# Uncertainty-Aware Multidimensional Ensemble Data Visualization and Exploration

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**Abstract**—This paper presents an efficient visualization and exploration approach for modeling and characterizing the relationships and uncertainties in the context of a multidimensional ensemble dataset. Its core is a novel dissimilarity-preserving projection technique that characterizes not only the relationships among the mean values of the ensemble data objects but also the relationships among the distributions of ensemble members. This uncertainty-aware projection scheme leads to an improved understanding of the intrinsic structure in an ensemble dataset. The analysis of the ensemble dataset is further augmented by a suite of visual encoding and exploration tools. Experimental results on both artificial and real-world datasets demonstrate the effectiveness of our approach.

Index Terms—Ensemble visualization, uncertainty quantification, uncertainty visualization, multidimensional data visualization

#### 1 Introduction

R ECENT developments in scientific simulation research have resulted in an increased role for scientific simulations and analysis. One common way to study uncertainty is the ensemble simulation, which employs stochastic initial conditions or multiple parameterizations to produce an ensemble of simulation outcome. For example, in the numerical weather simulation, both initial conditions and simulation parameters (e.g., cumulus schemes and microphysics schemes) can be perturbed in an ensemble forecast simulation. The number of ensemble runs ranges from dozens to thousands. In this paper, all ensemble members of a single data entry are called an ensemble data object, e.g., the ensemble numerical weather forecast at a geospatial location. The ensemble members of an ensemble data object may be averaged to obtain the ensemble mean, which is regarded as a representative of the ensemble members. Unfortunately, important information is lost in the averaging process. It is highly beneficial to characterize

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the uncertainty of an ensemble dataset during the entire analysis process.

In most scientific applications, the resulting simulation output is a multivariate field. For instance, the output of a typical weather research and forecasting (WRF) simulation [1] includes more than 100 variables such as temperature, pressure, and wind direction. Therefore, challenges for analyzing an ensemble dataset include both the *uncertainty* with respect to each output variable and the *high dimensionality* of the dataset.

There have been a large number of uncertainty visualization and analysis methods [2] that go beyond traditional summary statistics [3] and depict ensemble data in more detail. Conventional visualization solutions exploit glyphs [4], [5], [6] and visual variables [7], [8] to encode uncertainties. However, most of them are only designed for 1D or 2D datasets and have limited capabilities to reveal the intrinsic structures in the ensemble dataset. Recent methods can effectively characterize and analyze the uncertainty structures [9], [10] and forms [11], [12] but are designed for data objects with one variable.

To address the second challenge, high dimensionality, multidimensional projection techniques [13], [14] are widely used to build a low-dimensional layout that respects the distances among data objects in the high-dimensional space. In this low-dimensional layout, closely positioned points indicate similar data objects in the high-dimensional space. However, most conventional projection methods are developed for datasets in which a data object has only one instance. Simply using the ensemble mean as an instance to represent a data object for projection suffers from heavy information loss, because the shape of distribution for each ensemble data object is lost during the averaging process. A conceptual example is shown in Figs. 1a and 1b, where four ensemble data objects with similar ensemble means but different ensemble distributions are lumped together by a conventional multidimensional projection method to the ensemble means. This can cause misleading perceptions from users. For example, users might advocate that these

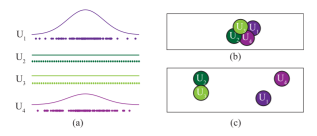


Fig. 1. Projecting an example 1D ensemble dataset (a) which consists of four ensemble data objects with similar mean values but different distributions. (b) The result of a conventional multidimensional projection algorithm to the ensemble means. (c) The result of our method.

four data objects are almost the same as they are located close to each other.

It remains a challenging task to visually explore the intrinsic structures of a high dimensional ensemble dataset as well as the uncertainty of each individual ensemble data object. The key contribution of this paper is a novel uncertainty-aware multidimensional projection approach that generalizes the conventional projection methods for ensemble datasets. Its core is a new dissimilarity measure of the ensemble data objects and an enhanced Laplacian-based projection scheme. Compared with the conventional projection methods that can solely respect the distances among ensemble means, our dissimilarity measure preserves the relationships among the ensemble distributions as well as the ensemble means. By setting the distributional difference weight to zero (see Equation (7)), our approach regresses to a conventional projection scheme. The enhanced Laplacianbased projection scheme achieves a balance between the local and global point layout by imposing global constraints in constructing the Laplacian system.

Fig. 1c illustrates the effectiveness of our uncertainty-aware projection approach, with which four data objects are positioned in the 2D visual plane. In this projection,  $U_2$  and  $U_3$  are located close to each other due to their similar ensemble means and distributions. As  $U_1$  and  $U_4$  follow different distributions from  $U_2$  and  $U_3$ , they are positioned far away from  $U_2$  and  $U_3$  even though all of them have similar ensemble means.

We note that as with any projection method, our method will not preserve the high-dimensional structures with 100 percent accuracy. Our goal is to provide users with a projection method to visualize not only the differences between the means of ensemble data objects, but also the differences between the shapes of the ensemble distributions. As demonstrated in Section 6 (see Fig. 6 and Fig. 11), a better separation of data objects with different ensemble distributions can be achieved with our method, but not with previous projection methods using ensemble means.

In addition, we augment users' abilities to visually study an ensemble dataset with a suite of visual exploration widgets: 1) an uncertainty histogram that allows for selecting the interesting range of uncertainty values in the visualized data; 2) an ensemble bar that embodies key information in an ensemble data object; 3) a parallel coordinates view that shows the details of the ensemble distribution for a particular ensemble data object; and 4) an optional geo-location view that depicts the uncertainty patterns identified in the projection view. Experimental results on a synthetic dataset

and numerical weather simulations alike show that ensemble data objects with similar ensemble means but different ensemble distributions can be clearly distinguished with our approach.

In summary, the main contributions of this work include:

- A novel uncertainty-aware multidimensional projection technique for ensemble datasets.
- A multi-view interactive visualization system for effective exploration of intrinsic structures and uncertainties in ensemble datasets.

# 2 RELATED WORK

Our work relates to multiple topics of visualization including uncertainty visualization, ensemble data visualization, and multidimensional/multivariate data visualization.

Uncertainty spreads throughout the entire data analysis pipeline including acquisition, transformation, and visualization [15], [16]. Depicting uncertainty can significantly help analysts make better decisions [17]. Thomson et al. [18] proposed a typology for uncertainty visualization in intelligence analysis. Potter et al. [19] presented a comprehensive survey for uncertainty quantification and visualization of scientific data. In the past decade, a large number of uncertainty visualization techniques have been developed. They can be roughly classified into four categories: glyph based, visual variable based, geometry based, and animation based. Glyph based methods encode uncertainties into well-designed glyphs (e.g., the flow radar glyph [4], the circular glyph [20], the summary plot [5], and the uncertain ODF glyph [21]) and place them into the original data field. Similarly, visual variables such as color [10], [22], brightness [8], [23], blurriness [24], and texture [7], [25] can also be employed to encode uncertainty. The geometry based approaches are a family of techniques that adapt the basic geometry to represent uncertainty, including point [22], line [26], cube [27], and surrounding volume [10], [28]. Coninx et al. [29] and Lundstrom et al. [30] demonstrate that animation can be used to express uncertainty as well. In general, most of these techniques focus on 1D or 2D datasets and cannot be readily extended to multidimensional datasets. Wu et al. [31] introduced the standard error ellipsoid to characterize uncertainty arising in any stage of a visual analytic process for multidimensional datasets. In addition, evaluations of different uncertainty visualization techniques have attracted much attention recently. Deitrick and Edsall [17] conducted an empirical evaluation to explore the influence of uncertainty visualization on decision making. Sanyal et al. [32] compared four commonly used techniques for 1D and 2D datasets. They identified that the glyph representations are good options generally.

Ensemble data is common in many scientific fields such as meteorology, hydrology, and chemistry. Ensembles effectively represent multiple simulations of a model with varied initial or boundary conditions, yielding slightly different results. They can be regarded as a type of uncertainty data. A convenient visualization method for ensemble data is the small-multiple method [33], [34] associated with linking and brushing operations. Spaghetti plots [35] is another approach to visualize ensemble dataset. Potter et al. [34]

introduced a framework, Ensemble-Vis, to visualize and explore ensemble dataset by leveraging multiple coordinated views. Sanyal et al. [6] developed a tool named Noodles to visualize ensemble uncertainty modeled with standard deviation. Unfortunately, Noodles is only applicable for single variable ensemble data. Recent work in this area has progressed from simple visualization to formal analysis of the results. Hollt et al. [36] presented an interactive system to study off-shore structures in an ensemble ocean forecasting dataset. Gosink et al. [11] proposed a method to characterize different types of predictive uncertainty in ensemble datasets based on the Bayesian model averaging. Mathias et al. [12] developed a Lagrangian framework for ensemble flow field analysis.

The differences between the terms multidimensional and multivariate are subtle. The term multidimensional refers to independent dimensions, while the term multivariate refers to that of dependent variables [37]. Nevertheless, multivariate data can be treated as multidimensional because the relationships among dimensions are typically unknown in advance. Conventional approaches to visualizing multidimensional data employ scatter plot matrix, parallel coordinates, star coordinates, etc. Among these approaches, multidimensional projection has gained much attention due to its ability to characterize similarities among data points in a visual space (2D or 3D), and has been proven to be a useful tool in many applications. Daniels et al. [38] employed a LSP-like projection technique for interactive vector field analysis. Chen et al. [39] proposed to embed high-dimensional DTI fibers to a 2D space with MDS for interactive exploration. Joia et al. [14] introduced an advanced projection technique called local affine multidimensional projection (LAMP) to interactively correlate similar data instances. Anand et al. [40] used random projection to find interesting substructures in a high-dimensional dataset. Although these approaches are capable of visualizing data by their low-dimensional layout, they are designed for data without uncertainty. Our approach advances the multidimensional projection scheme by taking uncertainty into account.

# 3 UNCERTAINTY-AWARE MULTIDIMENSIONAL PROJECTION

Multidimensional projection is emerging as an effective visual exploration technique for high-dimensional datasets. It projects a set of data points in the d-dimensional space into an l-dimensional visual space, typically  $l \in \{2,3\}$ . Conventional techniques assume that the value of each data object is deterministic. Many recent scientific simulations produce a collection of values at each data object due to the perturbed initial condition or parameterization. This type of data is called ensemble data.

The conceptual model of our data consists of n ensemble data objects  $U = \{U_1, U_2, \ldots, U_n\}$ . Each object has m d-dimensional ensemble members,  $U_i = \{U_i^1, U_i^2, \ldots, U_i^m \mid U_i^t \in \mathbb{R}^d\}$ . The goal of our uncertainty-aware projection scheme is to build an l-dimensional representation  $V = \{V_1, V_2, \ldots, V_n \mid V_i \in \mathbb{R}^l\}$  to preserve the relationships among data objects in terms of both the ensemble means and the ensemble distributions.

# 3.1 Approach Overview

To strike a balance between projection efficiency and accuracy, we employ a two-step multidimensional projection: first, a small set of control points are selected from U based on the ensemble means and projected to a 2D space with the conventional multidimensional scaling method [13]; then all other objects in U are projected to the 2D space with an enhanced Laplacian system that combines the influences from both control points and other data points.

One distinctive feature of our approach is that we take both the ensemble mean differences and the ensemble distribution differences into account to measure the dissimilarity  $D(U_i,U_j)$  of any pair of ensemble data objects  $U_i$  and  $U_j$ . The former is described by the euclidean distance between the ensemble means  $E(\overline{U}_i,\overline{U}_j)$ , and the latter is captured by a difference measure between ensemble distributions  $J(U_i||U_j)$  based on relative entropy [41]. To reconstruct the continuous ensemble distribution for each data object, a multidimensional kernel density estimation (KDE) method that considers the dimensional correlations is employed.

# 3.2 Ensemble Data Object and Probability Distribution

For simplicity, we suppose that the data objects are independent. Currently, many uncertainty modeling and visualization methods [9], [10] assume that the ensemble members for a given data object are drawn from a known parametric distribution such as Gaussian. In practice, the ensemble distribution is often not Gaussian and can vary from data object to data object.

KDE is a non-parametric approach to approximate the underlying continuous distribution by using a sum of kernels centered at each sample. Specifically, the multidimensional KDE [42] for data object  $U_i$  is defined as

$$U_i(x) = \sum_{t=1}^m w_t K_{\mathbf{H}} \left( x - U_i^t \right), \tag{1}$$

where **H** is a bandwidth matrix,  $w_t$  is a weight factor which can be determined by prior knowledge, and  $\sum_{t=1}^m w_t = 1$ .  $K_{\mathbf{H}}$  is the multidimensional kernel function satisfying that  $K_{\mathbf{H}}(x) \geq 0$  in the entire domain and  $\int K_{\mathbf{H}}(x) dx = 1$ . As such, the estimated density function can be interpreted as the probability density function.

A range of kernels can be adopted for KDE including uniform, Gaussian, and Epanechnikov kernels. We employ the widely used normal kernel [43], [44], because the kernel type is less important than the bandwidth parameter in terms of influences on the estimation [45].

In general, there are three choices for  $\mathbf{H}$ : the scaled identity matrix  $\mathbf{H} = h^2\mathbf{I}$ , the diagonal matrix  $\mathbf{H} = diag(h_1^2, h_2^2, \ldots, h_d^2)$ , and the general symmetric positive definite matrix. The scaled identity matrix implies that the variance in all dimensions are identical. The diagonal matrix implies that each dimension has its own bandwidth and no correlation exists between any two dimensions. In practice, a variety of automatic data-driven bandwidth selection methods can be used to estimate the bandwidth

 $h_k$  on the kth dimension, e.g., the Silverman's rule of thumb [45]

$$h_k = \left(\frac{4\sigma_k^5}{3m}\right)^{1/5} \approx 1.06\sigma_k m^{-1/5},$$
 (2)

where  $\sigma_k$  is the standard deviation of samples on the kth dimension, m is the number of samples. More sophisticated approaches like cross-validation exist in [46]. The general symmetric positive definite matrix encodes the individual bandwidth for each dimension and the linear relationship between any two dimensions.

Each type of bandwidth matrix has its own advantages and limitations. The scaled identity matrix is prone to under-fitting and has a large fitting error. The general matrix, which requires that **H** is symmetric and positive definite, is the most accurate one but needs more parameters and is prone to over-fitting. The diagonal matrix strikes a balance between them.

To take the advantage of simplicity by the diagonal matrix while preserving correlations among dimensions, we choose to perform KDE in a space defined by the *principal component transformation* [42]. Specifically, a mean centering process is first applied to each data object before estimation so that the distributional differences (to be detailed in Section 3.3) are not influenced by ensemble means

$$\widehat{U}_i^t = U_i^t - \overline{U}_i. \tag{3}$$

We then apply principal component analysis to  $\widehat{U}_i = \{\widehat{U}_i^t \mid t=1,2,\ldots,m\}$  that yields a transformation matrix  $\Phi$ .  $\Phi$  is composed by the eigenvectors of the covariance matrix of  $\widehat{U}_i$ . Lastly, we transform each ensemble member  $U_i^t$  into a new set  $U_i^* = \{U_i^{t*} \mid t=1,2,\ldots,m\}$  by

$$U_i^{t*} = \Phi^T \widehat{U}_i^t. \tag{4}$$

Consequently, the probability  $U_i(x)$  at location x is computed by  $U_i^*(x^*)$  on  $U_i^*$  instead, where  $x^* = \Phi^T(x - \overline{U}_i)$ . Because the bases of the new space are eigenvectors that are orthogonal to each other, the diagonal bandwidth matrix  $\mathbf{H}^*$  for estimating  $U_i^*(x^*)$  can be quickly estimated with Equation (2) on  $U_i^*$ .

# 3.3 Dissimilarity Estimation

The dissimilarity measure between pairs of data objects plays an essential role in most multidimensional projection techniques. Geometric distances such as the euclidean distance, the cosine distance, and the geodesic distance have prevailed in the multidimensional projection literature for years. However, these measures cannot be directly applied to ensemble datasets.

A naive way to deal with ensemble datasets is to use the summary statistics like the ensemble means to represent the data objects and calculate the dissimilarities among them. Unfortunately, this simple scheme results in a large amount of information loss. The Anscombe's quartet dataset [47] shows that four data objects with the same means and variances have very different ensemble distributions. Clearly, it

is impossible to distinguish them from each other with the ensemble means.

Therefore, it is natural to extend the definition of dissimilarities among ensemble data objects by considering their probability distributions. *Jensen-Shannon divergence* (JSD) [41] or relative entropy is a dissimilarity measure of distributions based on the *Kullback-Leibler divergence*. JSD is symmetric, non-negative, and bounded. It is widely used in the community of uncertain data mining [48]. The distribution difference  $J(U_i||U_j)$  between the data object  $U_i$  and  $U_j$  is defined as

$$J(U_i||U_j) = \frac{1}{2} \int_{\mathbb{D}} U_i(x) \log\left(\frac{2U_i(x)}{U_i(x) + U_j(x)}\right) dx + \frac{1}{2} \int_{\mathbb{D}} U_j(x) \log\left(\frac{2U_j(x)}{U_i(x) + U_j(x)}\right) dx,$$
 (5)

where  $U_i(x)$  and  $U_j(x)$  are the reconstructed probability density functions for  $U_i$  and  $U_j$  respectively,  $\mathbb D$  denotes the high-dimensional space where all the ensemble members  $U_i^t$  reside. Typically, we set the base of the log function as 2 so that  $J(U_i||U_j)$  is bounded by 1. Only when  $U_i$  and  $U_j$  have the same distributions does  $J(U_i||U_j)$  reach 0. By convention,  $0 \log 0$  is defined as 0.

In most cases, it is impossible to obtain the analytical solution of Equation (5). Therefore we compute its numerical solution with sampling. In our implementation, the Metropolis-Hastings (MH) [49] algorithm, a Markov Chain Monte Carlo method, is employed, because it can generate a sequence of samples from a function proportional to the target probability density distribution when direct sampling is difficult. As more and more samples are produced, the distribution of samples more closely approximates the target distribution. Consequently, by the law of large numbers, we can rewrite Equation (5) as

$$J(U_{i}||U_{j}) \approx \frac{1}{2S} \sum_{t=1}^{S} \log \left( \frac{2U_{i}(x_{i}^{t})}{U_{i}(x_{i}^{t}) + U_{j}(x_{i}^{t})} \right) + \frac{1}{2S} \sum_{t=1}^{S} \log \left( \frac{2U_{j}(x_{j}^{t})}{U_{i}(x_{j}^{t}) + U_{j}(x_{j}^{t})} \right),$$
(6)

where  $\{x_i^t | t = 1, 2, ..., S\} \sim U_i(x)$  and  $\{x_j^t | t = 1, 2, ..., S\} \sim U_j(x)$  are S samples generated by MH.

Therefore, we define the dissimilarity between  $U_i$  and  $U_j$  as a weighted sum of the euclidian distance between their ensemble means and the distributional difference between their probability density functions

$$D(U_i, U_j) = \alpha E(\overline{U}_i, \overline{U}_j) + \beta J(U_i \parallel U_j), \tag{7}$$

where  $\alpha \in [0,1]$ ,  $\beta \in [0,1]$  are two parameters adjustable by users, and  $\alpha + \beta > 0$ . When  $\alpha \neq 0$ ,  $\beta = 0$ , the distributional differences are missed by our dissimilarity measure. In such a case, our approach will regress to the conventional projection methods because only the dissimilarities among ensemble means are captured.

In summary, the proposed dissimilarity measure is endowed with the following properties:

• Symmetry  $(D(U_i, U_j) = D(U_j, U_i))$ . The dissimilarities between any pair of data objects are symmetric.

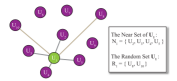


Fig. 2. The neighborhood graph for  $U_1$  with a near set of size 4 and a random set of size 2.

Therefore, it can be seamlessly incorporated into many state-of-the-art multidimensional projection techniques.

 Uncertainty-awareness. The dissimilarity considers the differences in ensemble distributions therefore differentiating data objects indistinguishable by summary statistics.

# 3.4 The Enhanced Laplacian-Based Projection

Our scheme of projecting ensemble data objects to the visual space is inspired by the least square projection (LSP) technique [50]. Generally, it is a two-step local technique. In the first step, a subset of data objects are projected to the visual space. In the second step, the rest of the data objects are interpolated according to the K-nearest neighborhood graph. However, this method inherits the drawbacks of local methods. It may bias the data objects projected in the second step in favor of control points projected in the first step [51] (see Fig. 3a). Thus, instead of directly using the Laplacian system originally proposed in [50], we propose to add global constraints into the equations. In other words, our method associates each data object with two small sets: a near set and a random set, instead of only one K-nearest set. Both sets are used for constructing the neighborhood graph for interpolation. The random set plays the role of the global constraints. This simple scheme strikes a balance between the locality maintained by the near set and the global layout preserved by the random set.

A large number of samples from the ensemble distributions may be needed to obtain an accurate estimation of the relative entropy (Equation (6)), especially when the ensemble distribution is complicated. To avoid pair-wise distributional difference calculations, we select control points and construct the neighborhood graph only based on the ensemble means  $\overline{U} = {\{\overline{U}_1, \overline{U}_2, \dots, \overline{U}_n\}}$ . Control points  $C = \{C_i \mid C_i \in U, i = 1, 2, ..., K\}$  are selected with a K-center algorithm [52]. If we do not have any priori knowledge about the dataset, the number of control points is set as  $K = \sqrt{n}$  [53]. The near set  $N_i$  for each ensemble object  $U_i$  is then defined as the ensemble objects whose means are the K-nearest neighbors of  $\overline{U}_i$ . Apparently,  $N_i$  might not be the true K-nearest neighbors of  $U_i$ because only the ensemble mean information is utilized. However, in practice the K-nearest neighbors in  $N_i$  can be guaranteed by enlarging the near set. In such a case, all other objects in  $N_i$  that are not true K-nearest neighbors of  $U_i$  can be deemed as objects in the random set  $R_i$ . Fig. 2 shows an example to build the neighborhood graph for the ensemble data object  $U_1$ .

In our implementation, an iterative majorization algorithm called scaling by majorizing a convex function

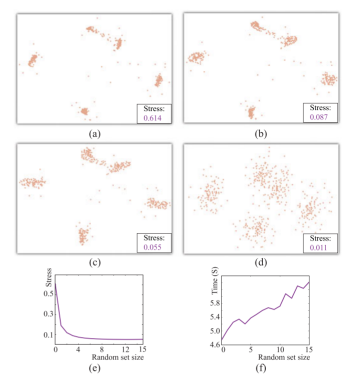


Fig. 3. The influence of different sizes of the random set in our approach. (a)  $|R_i|=0$ , which is the LSP technique. (b) Our method with  $|R_i|=4$ . (c) Our method with  $|R_i|=8$ . (d) MDS projection. (e) The stress plot. (f) The time plot.

(SMACOF) [13] is employed to build a low-dimensional representation  $\{V_i^c \mid V_i^c \in \mathbb{R}^2, i=1,2,\ldots,K\}$  for control points.  $V_i^c$  is the 2D projection of the control point  $C_i$ .

Technically, the Laplacian-based projection scheme rests on the theory of convex combination. That is, the low dimensional representation for each high-dimensional data object can be regarded as a linear combination of its neighborhoods in the visual space. Mathematically, let  $V_i \in \mathbb{R}^2$  be the projection of ensemble data object  $U_i$ , according to the convex combination theory,  $V_i$  can be written as

$$V_i = \sum_{U_j \in \{N_i \cup R_i\}} \tau_{ij} V_j, \tag{8}$$

where  $\tau_{ij} > 0$ ,  $\sum \tau_{ij} = 1$ . Typically, the inverse of dissimilarity  $D_{inv}(U_i, U_j) = 1/D(U_i, U_j)$  between the ensemble data object  $U_i$  and  $U_j$  is used to define the weight

$$\tau_{ij} = \frac{D_{inv}(U_i, U_j)}{\sum_{U_j \in \{N_i \cup R_i\}} D_{inv}(U_i, U_j)}.$$
 (9)

By reorganizing Equation (8) for all ensemble data objects into a matrix representation, a sparse linear system constrained by the control points can be derived [50] which can then be solved in a least square sense. The solution of this system is the low-dimensional representation of the ensemble dataset U.

Fig. 3 demonstrates the effect of using a random set. A synthetic dataset consisting of 500 data objects in five clusters is employed. Each ensemble data object has 80 members. Fig. 3a displays the projection result without the

random set. In this case, our method regresses to the Least Square Projection technique. Figs. 3b and 3c show the results with the size of the random set tested at 4 and 8 respectively. It is reasonable that a large random set preserves more global relationships among points, leading to low stress. This is empirically verified by the plot of standard normalized stress over the size of the random set (see Fig. 3e). For comparison, the projection result of MDS implemented with SMACOF [13], which optimizes the global relationships among points, is shown in Fig. 3d. MDS provides the smallest stress but requires the pairwise dissimilarity matrix that it is usually not feasible in projecting large datasets. Fig. 3f shows the time of our method with different sizes of the random set. From this plot, we can see that the consuming time is closely linear to the sum of  $|N_i|$  and  $|R_i|$ ,  $|N_i| = 20$  in this example. This is because the distributional difference estimation is much more time-consuming compared to building and solving the linear system. In summary, the addition of a random set provides a balance between the fast but inaccurate local methods (e.g., Laplacian-based projection methods) and the accurate but slow global methods (e.g., MDS).

# 4 UNCERTAINTY QUANTIFICATION AND VISUALIZATION FOR INDIVIDUAL ENSEMBLE DATA OBJECT

The uncertainty-aware 2D projection of the multidimensional ensemble data objects helps users understand the relationships among data objects in a simple 2D layout. It is equally important to allow users to further examine the uncertainty of each ensemble data object. This section describes our solutions for quantifying and visualizing uncertainty of the ith ensemble data object  $U_i$ .

#### 4.1 Uncertainty Quantification

Quantifying uncertainty plays an important role in the pipeline of uncertainty visualization [15]. It offers the variables for visual encoding. In this section, we describe two measures for quantifying the overall uncertainty of an ensemble data object and the detailed deviation for each ensemble member. We also discuss the limitations and application scenarios of each measure.

In the 1D case, the standard deviation is a widely used metric to quantify uncertainty of a random variable. Inspired by this rule, our first measure models the *overall uncertainty*  $O_i$  of the ensemble data object  $U_i$  as a sum of the standard deviations in all dimensions

$$O_i = \sum_{k=1}^d \sigma_i^k,\tag{10}$$

here  $\sigma_i^k$  represents the standard deviation on the kth dimension. The deviation  $\delta_i^t$  of the tth ensemble member of  $U_i$  is defined as its euclidean distance to the ensemble mean

$$\delta_i^t = \parallel U_i^t - \overline{U}_i \parallel . \tag{11}$$

The first measure is simple and can be used for most applications. However, it does not take the correlations

among dimensions into account. The second measure employs the covariance matrix to quantify uncertainty for each ensemble data object. The covariance matrix can be considered as a generalization of variance to multiple dimensions. It measures the dispersion of an ensemble data object with respect to the ensemble mean. A geometrical representation of a covariance matrix is the hyper ellipsoid [31], [54], whose axes correspond to the eigenvectors and square roots of eigenvalues of the covariance matrix. Similar to [31], we use the volume of the hyper ellipsoid to represent the overall uncertainty of an ensemble data object

$$O_i = \frac{\pi^{\frac{d}{2}}}{\Gamma(\frac{d}{2}+1)} \prod_{k=1}^d \sqrt{\lambda_k},\tag{12}$$

where  $\Gamma(\cdot)$  indicates the Gamma function,  $\lambda_k$  represents the eigenvalues of the covariance matrix  $\sum_i$ . Once established, the deviation of an ensemble member is defined as the Mahalanobis distance [55] to the ensemble mean

$$\delta_i^t = \sqrt{\left(U_i^t - \overline{U}_i\right)^T \sum_i^{-1} \left(U_i^t - \overline{U}_i\right)}.$$
 (13)

The second measure requires more computation than the first one due to a matrix inversion. In addition, a sufficient number of ensemble members are demanded in the second measure to correctly represent correlations among dimensions. Thus, the second measure is not a good option when the number of ensemble members is much smaller than the dimensionality of the dataset.

# 4.2 From Quantification to Visualization

Following the basic principles identified by Sanyal et al. [32] that the glyph size and color are two effective variables to display uncertainty in a 2D space, we design a color bar based representation called *ensemble bar* to depict the overall uncertainty and the distribution of ensemble members. Our goal is to provide users a quick preview of uncertainty patterns in an ensemble data object rather than complete details, which will be displayed in the parallel coordinates view. The height of the ensemble bar encodes the overall uncertainty: a higher bar implies an ensemble data object of larger overall uncertainty. The pattern shown in the bar depicts the distribution of all ensemble member deviations. To help users easily explore the distribution differences of different ensemble data objects, all ensemble bars have an identical width.

Let  $\delta_i = \{\delta_i^1, \delta_i^2, \dots, \delta_i^m\}$  be the deviations of the m ensemble members from the ensemble mean  $\overline{U}_i$ . The height of the ensemble bar is given by

$$H_i = (1 - O_i)H_{min} + O_iH_{max},$$
 (14)

where  $H_{min}$  and  $H_{max}$  represent the minimum and maximum heights of all ensemble bars, respectively.  $O_i \in [0, 1]$  is the normalized overall uncertainty of  $U_i$ .

In order to convey the distribution of ensemble member deviations in the bar, we first set the left and right edges of the bar to correspond to the minimum deviation  $min(\delta_i)$  and the maximum deviation  $max(\delta_i)$ , respectively. Then, we equally discretize the entire bar into a set of bins and

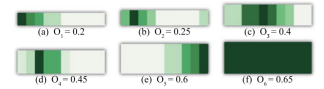


Fig. 4. Various types of ensemble bars show different types of ensemble data objects, each of which consists of 100 members. The overall uncertainty increases from (a) to (f). In each case, a bar in dark green indicates the peak of the distribution of ensemble member deviations. The ensemble member deviations approximately follow: (a-b) the positively skewed distribution; (c-d) the normal distribution; (e) the negatively skewed distribution; (f) the uniform distribution.

count the number of ensemble members in each bin. Each bin is drawn as a rectangle with a linearly varied sequential color map (e.g., from white to green in this paper).

The number of bins has a great impact on the pattern shown in an ensemble bar. In practice, various useful guidelines and rules of thumb can be employed. In our implementation, we take the square root of the number of ensemble members as the number of bins. This scheme is simple and has been widely used in many applications such as Excel. Users are allowed to interactively adjust the number of bins to discretize the ensemble bar.

In general, the ensemble bar allows users to compare overall uncertainties of different ensemble data objects through the heights of the ensemble bars (see Fig. 4). In addition, the patterns of bins enable users to identify different types of distributions. Taking the uniform distribution as an example, an approximately monochromatic ensemble bar is produced with our method (see Fig. 4f). This indicates that all bins in the ensemble bar almost have the same number of ensemble members. Similarly, for positively skewed distribution of which more ensemble members are close to the ensemble mean, an ensemble bar with the dark-green bin on the left is generated (see Figs. 4a and 4b).

#### 5 VISUAL EXPLORATION AND INTERACTIONS

This section describes an integrated system with a suite of visualization and interaction tools for visually exploring a multidimensional ensemble dataset.

#### 5.1 The Exploration Workflow

The exploration process starts with an uncertainty histogram, on which users can select target uncertainty intervals (e.g., intervals with high levels of uncertainty). Then, users can interactively investigate the low-dimensional layout of the ensemble data objects in the 2D projection view equipped with a set of interactions such as zoom in/out, and pointer/lasso selection. Once points of interest are specified, the geospatial locations associated with each data object will be highlighted in the Geo-Location view. Meanwhile, the ensemble bar view shows all selected data objects. Users can further drill down to a specific ensemble bar to explore the high-level distribution and the overall uncertainty of a data object. Afterwards, users can select bins of interest in the ensemble bar to examine the selected ensemble members in an animated continuous parallel coordinates view. The parallel coordinates view allows users to study the distribution of ensemble members on

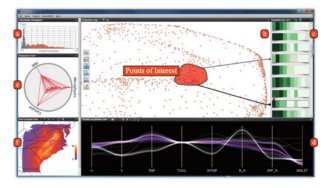


Fig. 5. The main interface of our visual exploration system. (a) The uncertainty histogram. (b) The 2D projection view. (c) The ensemble bar view. (d) The continuous parallel coordinates view. (e) The parameters view. (f) The Geo-Location view.

each dimension. To investigate the source of uncertainty (e.g., parameter perturbations in this paper), a radial plot is provided to display the parameter configurations that generate these ensemble members.

# 5.2 Exploration Tools

The main user interface is composed of a set of linked widgets and views to enhance interactivity. Fig. 5 shows an overview of our system.

*Uncertainty histogram.* At the top left corner, an uncertainty histogram widget allows users to select data points with overall uncertainty in a desired range.

Projection view. This view displays the intrinsic structures of an ensemble dataset in the 2D plane. The distances among 2D data points encode the similarities of the corresponding ensemble data objects with regard to both ensemble means and ensemble distributions. Users can either select a point or a group of points for further analysis. A set of operations are provided in this view to assist exploration including zooming, panning, hovering, pointer selection, lasso selection, and multi-selection with the control key pressed.

Ensemble bar view. To facilitate comparison of overall uncertainties for different data objects, the ensemble bar view is displayed at the right side of the main interface. The overall uncertainties are perceived by the heights of the bars, and the ensemble distributions are depicted by different types of patterns in the ensemble bars. Following interactions are provided in this view:

- Choosing a color map and the bin number;
- Dragging bars together for detailed comparison;
- Reordering bars according to the shown patterns;
- Selecting bins of interest to further examine ensemble members in the parallel coordinates view.

Parallel coordinates view. A continuous representation is employed to convey uncertainty at each dimension. Bright areas on each dimension indicate high certainty. Users can detect outliers with line dyeing, brushing, local zooming, and axis reordering. To ensure a smooth interaction, animated transitions are implemented in this view.

Parameters view. Studying the effect of parameters in an ensemble simulation (that is, ensemble members are generated by different parameterizations) is an important task. For this purpose, we employ a radial plot where each

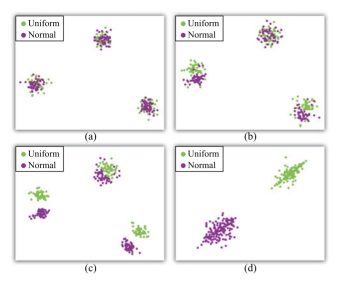


Fig. 6. The projections produced with different dissimilarity weights on the ensemble mean difference and the ensemble distribution difference in the dissimilarity measure. (a)  $\alpha=1.0$ ,  $\beta=0.0$ . (b)  $\alpha=0.5$ ,  $\beta=1.0$ . (c)  $\alpha=0.25$ ,  $\beta=1.0$ . (d)  $\alpha=0.0$ ,  $\beta=1.0$ .

parameter corresponds to one of the equiangular axes. Initially, all parameter configurations are shown as a context. Once users select a set of ensemble members, the corresponding parameter configurations are highlighted in this view.

*Geo-Location view*. For geolocation-related applications, a geo-location view is employed. Uncertainty features identified in the projection view may be easily understood and verified by the domain experts in this view.

#### 6 EXPERIMENTS AND DISCUSSION

We conducted three experiments to demonstrate the effectiveness and usefulness of our uncertainty-aware projection method and the exploration system. The system was implemented with the standard C++ and Qt 5.0. The experiments were performed on a PC equipped with an Intel Core 2 Duo 3.0 GHz CPU, 4 GB host memory and an NVIDIA Quadro 4000 video card with 1.5 GB video memory.

#### 6.1 The Synthetic Dataset

We generated a synthetic 5D dataset consisting of 300 ensemble data objects to demonstrate the effectiveness and the influence of dissimilarity weights. We first randomly chose three anchor points that constitute an equilateral triangle as cluster centers in a 5D space. Then, we generated 100 ensemble data objects in each cluster. Each data object had 250 ensemble members following either a uniform or a normal distribution. The ensemble mean of a data object in a cluster was set as the cluster center with a small random bias, and the correlation matrix was set as a fixed scaled identity matrix. This process generated a dataset with three clusters. The euclidean distances between any two cluster centers are equal. The geometrical differences of data objects in the same cluster are subtle but the distributional differences are significant. The equilateral triangle structure in this dataset is designed to demonstrate capability of our method to preserve the high-dimensional features in a visual space.

Fig. 6 shows the effect of the dissimilarity weights  $\alpha$  and  $\beta$ (see Equation (7)) in our uncertainty-aware multidimensional projection method. Without considering the distributional differences ( $\beta = 0$ ), all data objects with similar ensemble means are projected close to each other (see Fig. 6a) even though data objects within a cluster have distinct distributions. This is consistent with the result of the conventional multidimensional projection methods designed for certain datasets. When the distributional differences are incorporated ( $\beta > 0$ ), data objects within a cluster are gradually separated (see Figs. 6b and 6c). For demonstration, we color the uniformly distributed data objects with green and normally distributed data objects with purple. From Figs. 6b and 6c, we can observe that there are three clusters. In each cluster, we can further identify two subclusters, each of which corresponds to a particular type of distribution. However, these structures cannot be observed in Fig. 6a, as data objects within a cluster become distinguishable only when ensemble distributions are considered.

In general, our method not only preserves the global structure of the three clusters but also distinguishes data objects between the two types of ensemble distributions within each cluster. The degree of separation of the data objects within a cluster depends on the relative value of  $\alpha$  and  $\beta$ . When the ensemble mean is not considered ( $\alpha = 0$ ), all data objects are separated into two clusters solely based on the two types of ensemble distributions (see Fig. 6d).

# 6.2 The NBA Players' Statistics Dataset

We applied our method to a dataset consisting of 933 NBA players' career statistics<sup>1</sup> since 1981. In this dataset, we treat each player as an ensemble data object, and the statistics in a season as an ensemble member. Each ensemble member is a 16-dimensional vector that describes the statistics in a season including: Games Played, Minutes Played, Field Goals, Field Goal Attempted, three-Point Field Goals, three-Point Field Goal Attempted, Free Throws, Free Throw Attempted, Offensive Rebounds, Defensive Rebounds, Assists, Steals, Blocks, Turnovers, Personal Fouls, and Points. Each ensemble data object has at least five ensemble members. Because the number of the ensemble members for most ensemble data objects is smaller than the dataset dimensionality of 16, we chose the first uncertainty quantification measure to evaluate the overall uncertainty (Equation (10)) and the ensemble member deviations from mean (Equation (11)). The overall uncertainty in this case reflects the fluctuation of a player's performance over years, or a player's inconsistency.

Fig. 7 shows the results of the uncertainty-aware projection with different parameters. The color encodes the position of a player. We can see a triangular structure in Figs. 7a and 7b. The points in the upper part mainly represent players who play center (C) and players who play both center and forward (C-F). The points in the lower part primarily represent players who play guard (G) and players who play

<sup>1.</sup> The statistics of a player who has career over five seasons are collected from a professional basketball statistics website (http://www.basketball-reference.com)

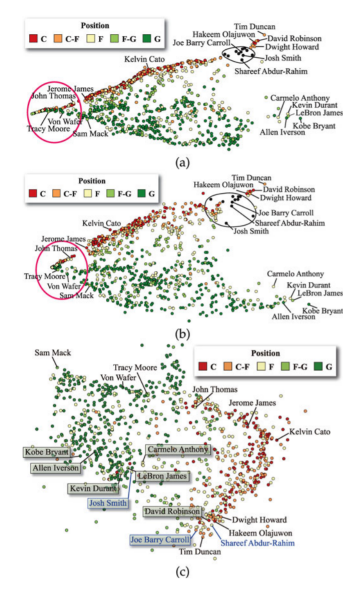


Fig. 7. Results for the NBA players' statistics dataset. (a) The result that considers only the geometrical differences,  $\alpha=1.0,~\beta=0.0.$  (b) The result of our method with  $\alpha=0.55,~\beta=0.45.$  (c) The result that considers only the distributional difference,  $\alpha=0.0,~\beta=1.0.$  In this example,  $|R_i|=8$  and  $|N_i|=2.$ 

both forward and guard (F-G). In addition, it is easy to find that yellow points are distributed in both the upper and the lower part. This is because players of forward (F) usually are versatile. For demonstration, we annotated a set of players with their names. The results in Figs. 7a and 7b show that points on the left side are role players and the points on the right side are key players. More specifically, the points in the upper right corner represent excellent players of C or C-F, and the points in the lower right corner represent excellent players of G or F-G.

Fig. 7c shows the result of projection using only distributional differences. This result indicates that the seasonal fluctuation patterns of the players in C and C-F positions are quite different from those of players in F-G and G positions. Furthermore, if we only use the geometrical differences among players to model their dissimilarities, the consistency of a player will be hidden. For example, without considering the distributional differences, the points in a region indicated

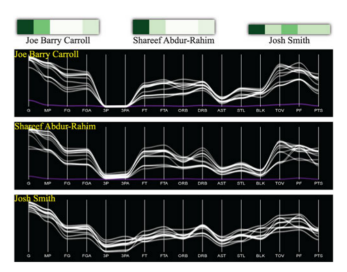


Fig. 8. The ensemble bars and parallel coordinates representation for the three selected players.

by the magenta circle in Fig. 7a are lumped together. This implies that their ensemble means are similar. However, Fig. 7c reveals that their ensemble distributions are distinct. This explains why points inside the magenta circle in Fig. 7b are much more dispersed. And this dispersion may help a general manager identify consistent players from a pool of players who are similar in mean performance.

To further inspect the difference between the results with and without considering distributional differences, we highlight a group of points in a region indicated by the black circle in Fig. 7a. The same group of points are also highlighted in black in Fig. 7b for comparison. Because Fig. 7a only considers the geometrical difference among ensemble means, the highlighted points close to each other have subtle differences among ensemble means. Take Josh Smith and Joe Barry Carroll as an example, they are located close to each other in Fig. 7a. However, they are dispersed when taking the distributional difference into account in Fig. 7b. This is because their distributional differences are significant, which can be clearly verified in Fig. 7c (notice the annotations in blue). The ensemble bars and the parallel coordinates views shown in Fig. 8 further verify this observation. On the other hand, Joe Barry Carroll and Shareef Abdur-Rahim are positioned closer to each other when considering the distributional difference because their ensemble distributions are quite similar, which is evident in their ensemble bars and the parallel coordinates representations in Fig. 8.

From the heights of the ensemble bars in Fig. 8, we can conclude that the consistency of Josh Smith is higher than those of Joe Barry Carroll and Shareef Abdur-Rahim. Further examining the details, we can see that the high overall uncertainties of Joe Barry Carroll and Shareef Abdur-Rahim are partly caused by some flat and close-to-zero outlier ensemble members (highlighted in purple). This was caused by seasons in which they may have injuries.

#### 6.3 The Numerical Weather Simulation Dataset

The dataset used in this experiment is an ensemble WRF simulation of the 1993 superstorm at 6:00 PM, Mar. 13th, 1993. We select a region with latitude from 30°N to 40°N,

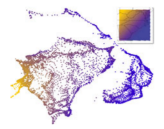


Fig. 9. The result of our projection method with  $\alpha=0.4$ ,  $\beta=0.6$ ,  $|R_i|=24$  and  $|N_i|=4$ . The color of a point is indexed from a 2D linear map (from violet to orange) with the geo-coordinates.

and longitude from 85°W to 75°W. The dataset consists of 7,050 geospatial grid points. Each grid point contains 8 variables, including wind speed in the zonal (E-W) and meridional (N-S) directions respectively, temperature, soil temperature, water temperature, relative humidity, specific humidity, and mean sea level pressure. Forty runs of the WRF simulation with different cumulative schemes and microphysics schemes generated the ensemble members. In other words, our simulation produced an 8-dimensional ensemble dataset with 7,050 ensemble data objects, each of which has 40 ensemble members. As the number of ensemble members is much larger than the dimensionality of the dataset, the second uncertainty quantification measure was employed in this case.

Fig. 9 presents our results. To help users easily associate the geospatial locations in the projection view, we use color to encode the geospatial locations. Specifically, we synthesize a 2D texture that covers the entire simulation region, i.e., latitude from 30°N to 40°N, and longitude from 85°W to 75°W. For simplicity, a linear gradient fill mode from violet to orange along the diagonal for this texture is adopted. In this way, the points sampled from the sea area will be encoded with color close to violet and the points sampled from the land area will be encoded with color close to orange (see Fig. 9). For clarity, the map is drawn as a background. From the projection result, we can easily find that points in the right part are closer to the sea area and points in the left part are closer to the land area.

Our system can help users explore the dataset from multiple perspectives and levels-of-detail. We present two exploration scenarios below.

Visual cluster analysis. Identifying highly correlated data objects is a crucial data exploration task to gain insights into an ensemble dataset because highly correlated data objects (i.e., the clusters) constitute the major structure of a dataset.

Fig. 10 shows the result of our uncertainty-aware projection method colored by the overall uncertainty. In this result, a color set from purple (low uncertainty) to yellow (high uncertainty) is employed. Because the overall uncertainty varies significantly, we employ a log transformation and scale the overall uncertainty to [0,1].

From the resulting 2D manifold, we can easily find that the entire dataset can be roughly grouped into three clusters C1, C2, and C3. With the lasso tool, we can further select the points in these clusters and show the geospatial locations associated with each data object in the Geo-Location view. The results show that C1 contains data objects located in the Chesapeake bay and Delaware bay area, C2 contains data objects located in the sea area, and C3 contains data objects

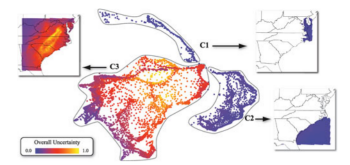


Fig. 10. Cluster identification and analysis based on our uncertainty-aware projection method. Three preliminary clusters are clearly shown with our method. The geospatial locations encoded with overall uncertainties are displayed on the map. In this example,  $\alpha=0.3,\ \beta=0.7,\ |R_i|=24,$  and  $|N_i|=4.$ 

located over the land. This separation of data points based on land types is particularly interesting given that the geospatial location is not part of the data used in projection. From the results in the Geo-Location view, we can further find that data objects over the sea have lower uncertainties than those over the land.

For comparison, we also projected this dataset by only considering the geometrical differences and the distributional differences respectively. The near set, the random set, the neighborhood graph, and the control points used in each projection were identical, while  $\alpha$  and  $\beta$  were different. Fig. 11a shows the result that only takes geometrical differences into account, i.e.,  $\alpha=1.0$  and  $\beta=0.0$ . Fig. 11c displays the result that only considers distributional differences, i.e.,  $\alpha=0.0$  and  $\beta=1.0$ . Fig. 11b presents the result that considers both geometrical and distributional differences with  $\alpha=0.3$  and  $\beta=0.7$ .

From Fig. 11a, we can see that the major structure shown in this result is similar to that in the result produced by our uncertainty-aware projection method in Fig. 11b. However, the difference between the data objects over the sea and the land is captured much more clearly by taking both the geometrical and distributional differences into account. Noticing the red points in C1 highlighted in Fig. 11a, many of them are lumped with points in C3. This is because the geometrical differences of ensemble means among these points are subtle. On the other hand, from Fig. 11c we know that the distributional differences between data objects in C1 and C3 are significant because C1 and C3 are clearly separated in this projection.

Furthermore, more points outside the three main clusters (e.g., points in the orange circle) are produced by the method without considering distributional differences in Fig. 11a. To investigate the reason why our method can avoid these outliers, three groups (G1, G2, and G3) of points are highlighted in different colors. For inspection, the points of G1, G2, and G3 are also highlighted in Figs. 11b and 11c. We calculate the average pairwise geometrical difference between two groups by

$$\overline{E}(P,Q) = \frac{\sum_{U_i \in P, U_j \in Q} E(\overline{U}_i, \overline{U}_j)}{|P||Q|}.$$
 (15)

Similarly, the average pairwise distributional difference between two groups is calculated by

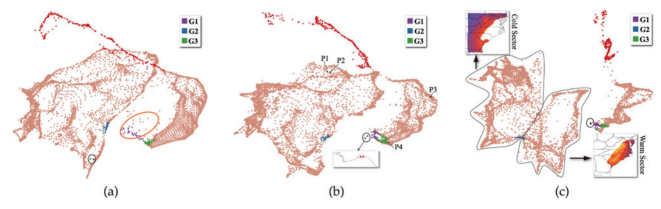


Fig. 11. (a) The result that considers only geometrical differences,  $\alpha=1.0$ ,  $\beta=0.0$ , i.e., the projection method applied to the ensemble means. (b) The result of our uncertainty-aware projection,  $\alpha=0.3$ ,  $\beta=0.7$ . (c) The result that considers only the distributional differences,  $\alpha=0.0$  and  $\beta=1.0$ . In this example,  $|R_i|=24$  and  $|N_i|=4$ .

$$\overline{J}(P,Q) = \frac{\sum_{U_i \in P, U_j \in Q} J(U_i, U_j)}{|P||Q|}.$$
(16)

In this case,  $\overline{E}(G1,G2)=0.29$ ,  $\overline{E}(G1,G3)=0.27$ . It means that the average geometrical dissimilarity of **G1** to **G2** and **G3** are almost the same. This faithfully explains why points in **G1** are located between **G2** and **G3** in Fig. 11a. However,  $\overline{J}(G1,G2)=0.82$ ,  $\overline{J}(G1,G3)=0.44$ . This means that the ensemble distributions of points in **G1** have higher dissimilarities to those points in **G2**. The resulting projection in Fig. 11c visually verifies this observation. From the result presented in Fig. 11b, we conclude that our method strikes a balance between methods that only consider geometrical differences or distributional differences.

Another interesting observation is that there are two points located between G1 and G2 in Fig. 11b (the region indicated by the black circle). We highlight their geospatial locations in the Geo-Location view. From the magnified map, we can see that they are sampled from the Florida shore of the Gulf of Mexico. For demonstration, we also highlight these two points both in Figs. 11a and 11c (the black points in the black circles). From Fig. 11a, we can infer that the ensemble means of these two points are similar to points over the land. From Fig. 11c, we can infer that their ensemble distributions are much more similar to points over the sea. These observations further prove the necessity of modeling the dissimilarities among ensembles with both geometrical and distributional differences.

Ensemble distribution investigation. Our system can also be employed to investigate the patterns of uncertainty in different areas and study the reason for the uncertainty of a single data object. For simplicity, we selected a set of representative points (P1, P2, P3, and P4 in Fig. 11b) in different areas for analysis. Fig. 12 shows the ensemble bars and parallel coordinate representations for the selected points. From the heights of the ensemble bars, we can see that P1 or P2 has a higher overall uncertainty than those of P3 and P4. From the patterns depicted in the ensemble bars of P1 and P2, we can infer that their ensemble member deviations follow an approximately uniform distribution. By examining their parallel coordinates representations, we can visually verify this observation. From the ensemble bars of P3 and P4, we may conclude that their ensemble member

deviations approximately follow a positively skewed distribution which is further confirmed by their parallel coordinates representations. Another interesting observation about the ensemble bars of P3 and P4 is that the color of the last bin varies greatly (see the magnified bins on the right side). This suggests that P4 has more outlier members. By selecting the last bin in the ensemble bars of P3 and P4, more ensemble members are highlighted in the parallel coordinates representation of P4.

#### 6.4 Discussion

Parameter configuration. Figs. 3e and 3f reveal that the near set  $|N_i|$  and the random set  $|R_i|$  have a great impact on projection efficiency and accuracy. A large near set and random set require more computation time and resource, while a high projection accuracy (i.e., low stress) can be obtained. Because the dissimilarity estimation process relates to the

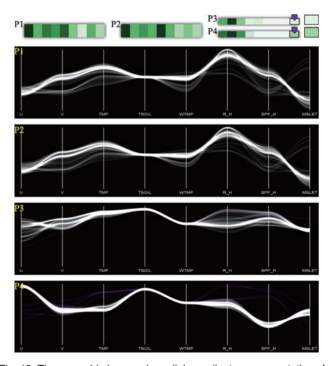


Fig. 12. The ensemble bars and parallel coordinates representations for four selected data objects. The selected bins are magnified on the right side. All selected ensemble members are highlighted in purple.

number of ensemble members and the dimensionality of the dataset, it is intractable to find an optimal size for the near set and the random set. In our implementation, a good balance between efficiency and accuracy is achieved by setting  $|N_i|$  close to  $\sqrt{n}/3$  and  $|R_i|$  no larger than 4.

The experiment in Section 6.1 demonstrates the influence of the dissimilarity weights  $\alpha$  and  $\beta$ . Empirically, if users need to put more emphasis on geometrical differences, a large  $\alpha$  should be assigned. On the other hand, if users want to highlight the distributional differences among data objects, a large  $\beta$  needs to be assigned. Without any prior knowledge,  $\alpha$  and  $\beta$  are both set as 0.5 for a preliminary study of an ensemble data set.

*Projection performance.* The overall projection complexity of our method is determined by three steps: building the neighborhood graph, estimating dissimilarities, and solving the sparse linear system.

Building the neighborhood graph involves defining a near set and a random set for each data object. A standard procedure to find the nearest neighbors is prohibitive  $O(n^2)$ . By reorganizing the input data objects into  $\sqrt{n}$  clusters [50], the complexity can be reduced to  $O(n^{\frac{3}{2}})$ .

A core component of our method is to estimate the distributional differences among data objects in the neighborhood graph. The complexity is  $O(n(|N_i|+|R_i|)mdS)$ . Here  $|N_i|$  is the size of the near set,  $|R_i|$  is the size of the random set, m is the number of ensemble members, d is the dimensionality of the dataset, and S is the number of samples. In our implementation  $|N_i| = \sqrt{n}/3$ , both  $|R_i|$  and S are constants, thus the complexity becomes  $O(n^{\frac{3}{2}}md)$ .

As the derived linear system is sparse, symmetric, and positive definite, the conjugated gradient method is employed. In this case, the complexity is  $O(n\sqrt{k})$  [50], where k is the condition number of the matrix  $L^TL$ . (L represents the coefficient matrix of the linear system.)

In general, the complexity of our method is  $O(max(n^{\frac{3}{2}}md,n\sqrt{k}))$ . In practice, dissimilarity estimation dominates the projection time in our experiments. For example, several hours are required to compute the dissimilarities in the numerical weather simulation dataset on our computer. A parallel implementation of the entire system would alleviate this problem. The computation consumption can be further decreased by employing a fast KDE algorithm, e.g., the KD-tree based KDE. This is an avenue for future work. In addition, from Equation (7), we know that changing the value of  $\alpha$  and/or  $\beta$  for exploration does not need to re-estimate the geometrical and distributional differences. Accordingly, we can obtain a new projection in seconds by solving a new linear system.

Strengths and limitations. By considering both geometrical and distributional differences, data objects with similar summary statistics can be distinguished with our approach. Our enhanced Laplacian-based projection scheme strikes a balance between the computation time and accuracy, which can be easily verified in Figs. 3e and 3f. Currently, both the probability reconstruction and the dissimilarity estimation processes in our approach are limited to numerical data objects. In the future, we will seek to adapt our approach for categorical multivariate ensemble data visualization and exploration.

# 7 EVALUATION

We evaluated the exploration system with a meteorologist collaborator to analyze the ensemble simulation of the 1993 superstorm. We examined the ensemble data at the 44th hour of the simulation out of the entire 49-hour simulation. We intentionally chose a time point towards the end of the simulation when there are more diversities among the ensemble distributions. According to the findings in Section 6.3, the dissimilarity weights were specified as  $\alpha=0.3, \beta=0.7$  so that data objects from different land types were clearly separated. The meteorologist can interactively adjust these weights to compare projection patterns. The observations and feedback are summarized in this section.

## 7.1 The Uncertainty-Aware Projection Method

The meteorologist observed that, without considering the distributional differences among data objects, the Chesapeake bay/Delaware bay and the surrounding land are overlapped in the projection view (see Fig. 11a). When considering both geometrical and distributional differences, these two types of land covers are clearly separated (see Figs. 11b and 11c). The meteorologist hypothesized that this separation is caused by the more precise ensemble simulations over the sea than over the land. This hypothesis came from the past experience with WRF simulations and was validated by examining the parallel coordinates of data objects over the land and the sea.

The meteorologist also observed that the warm sector and the cold sector (see Fig. 11c) are much better separated in the projection view when considering distributional differences rather than considering geometrical differences only. This is because the uncertainty between the ensemble members significantly increased in the warm sector located closer to the center of storm at that time point. The reason for larger uncertainty is the uncertain timing of the cold front associated with the storm. The different ensemble members timed the cold frontal passage differently, which affects when each point in the warm sector leaves the warm sector and enters the cold sector.

Based on these observations, the meteorologist agreed that projecting the data objects according to both geometrical and distributional information provides a novel and useful perspective on the data.

# 7.2 The Uncertainty Visualization

The meteorologist stated that it is a routine but an effective means to use color for showing uncertainty on a geospatial map. The color map provided a useful overview of the uncertainty of data objects in regions of high or low uncertainty. The meteorologist further confirmed that the high uncertainty region (i.e., the yellow region) in Fig. 10 coincides with the path of the super storm center.

Besides color mapping, the ensemble bar representation provides a supplementary visualization for the ensemble uncertainty. The meteorologist asserted that this representation was intuitive and helped him study the consensus of ensemble members and compare different data objects. The meteorologist concluded that the ensemble bars can be used to characterize and identify different types of uncertainty. For example, the ensemble bars with dark green color to the

left indicate positive skew distributions where most ensemble members are grouped together with few outliers (e.g., data objects located in the sea surface). Ensemble bars with dark green color in the middle or right represent little to no agreement among ensemble members (e.g., data objects located in the warm sector).

## 7.3 The Linked Multi-View Interface

The meteorologist found the linked interaction mode helpful. A cluster of points in the projection view can be easily selected and their ensemble distributions can then be reviewed in the ensemble bar view immediately. The selection of bins in an ensemble bar allows a group of ensemble members to be quickly selected and reviewed in the parallel coordinates view and the parameter view. In particular, the last several bins in an ensemble bar usually contain the outlier ensemble members that require further analysis.

The parallel coordinates view helps the meteorologist examine the distribution of individual variables of a data object. The meteorologist prefers the continuous parallel coordinates plot for a more intuitive presentation of the distribution. The ability to reorder the axes in the parallel coordinates view helps organize the similarly distributed variables close to each other for comparison.

The meteorologist pointed out that the soil temperature variable was not simulated, but inherited from the input NARR dataset. Thus, it contributed no uncertainty in the ensembles. The meteorologist expressed the desire to interactively select a group of variables for analysis. Currently, this feature is not supported in our system due to the time-consuming dissimilarity computation. However, a parallel computing system can help achieve this goal in the future.

#### 8 Conclusion

We present a visualization and exploration system for a multidimensional ensemble dataset whose kernel is a novel uncertainty-aware multidimensional projection method. This new method considers not only the ensemble means but also the ensemble distributions. Experiments on an artificial dataset demonstrate the ability of the uncertainty-aware projection to distinguish between ensemble data objects with similar means but different ensemble distributions. Results on both real-world and simulation datasets verify that 1) differences in ensemble distributions are an important part and are crucial for proper ensemble analysis, and 2) our uncertainty-aware projection can distinguish ensemble distributions in real-world data.

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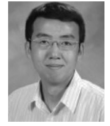
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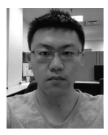
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