Uncertainties of predictions from parton distribution functions. II. The Hessian method

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We develop a general method to quantify the uncertainties of parton distribution functions and their physical predictions, with emphasis on incorporating all relevant experimental constraints. The method uses the Hessian formalism to study an effective chi-squared function that quantifies the fit between theory and experiment. Key ingredients are a recently developed iterative procedure to calculate the Hessian matrix in the difficult global analysis environment, and the use of parameters defined as components along appropriately normalized eigenvectors. The result is a set of 2D eigenvector basis parton distributions (where $d \approx 16$ is the number of parton parameters) from which the uncertainty on any physical quantity due to the uncertainty in parton distributions can be calculated. We illustrate the method by applying it to calculate uncertainties of gluon and quark distribution functions, *W* boson rapidity distributions, and the correlation between *W* and *Z* production cross sections.

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I. INTRODUCTION

The partonic structure of hadrons plays a fundamental role in elementary particle physics. Interpreting experimental data according to the standard model (SM), precision measurement of SM parameters, and searches for signals of physics beyond the SM, all rely on the parton picture of hadronic beam particles that follows from the factorization theorem of quantum chromodynamics (QCD). The parton distribution functions (PDF's) are nonperturbative—and hence at present uncalculable—functions of momentum fraction x at a low-momentum transfer scale Q_0 . They are determined phenomenologically by a global analysis of experimental data from a wide range of hard-scattering processes, using perturbative QCD to calculate the hard scattering and to determine the dependence of the PDF's on Q by the renormalization-group based evolution equations.

Considerable progress has been made in several parallel efforts to improve our knowledge of the PDF's [1–3], but many problems remain open. In the conventional approach, specific PDF sets are constructed to represent the "best estimate" under various input assumptions, including selective variations of some of the parameters [4–6]. From these results, however, it is impossible to reliably assess the uncertainties of the PDF's or, more importantly, of the physics predictions based on PDF's. The need to quantify those uncertainties for precision SM studies and new physics searches in the next generation of collider experiments has stimulated much interest in developing new approaches to this problem [7,8]. Several attempts to quantify the uncer-

tainties of PDF's in a systematic manner have been made recently [9–13].

The task is difficult because of the diverse sources of experimental and theoretical uncertainty in the global QCD analysis. In principle, the natural framework for studying uncertainties is that of the likelihood function [12,14,15]. If all experimental measurements were available in the form of mutually compatible probability functions for candidate theory models, then the combined likelihood function would provide the probability distribution for the possible PDF's that enter the theory. From this, all physical predictions and their uncertainties would follow. Unfortunately, such ideal likelihood functions are rarely available from real experiments. To begin with, most published data sets used in global analysis provide only effective errors in uncorrelated form, along with a single overall normalization uncertainty. Secondly, published errors for some well-established experiments appear to fail standard statistical tests, e.g., the χ^2 per degree of freedom, may deviate significantly from 1.0, making the data set quite "improbable." In addition, when the few experiments that are individually amenable to likelihood analysis are examined together, they appear to demand mutually incompatible PDF parameters. A related problem is that the theoretical errors are surely highly correlated and by definition poorly known. All these facts of life make the idealistic approach impractical for a real-world global QCD analysis.

The problems that arise in combining a large number of diverse experimental and theoretical inputs with uncertain or inconsistent errors are similar to the problems routinely faced in analyzing systematic errors within experiments, and in averaging data from measurements that are marginally compatible [16]. Imperfections of data sets in the form of unknown systematic errors or unusual fluctuations—or both—are a common occurrence. They need not necessarily destroy the value of those data sets in a global analysis; but we must adapt and expand the statistical tools we use to analyze the data, guided by reasonable physics judgement.

In this paper we develop a systematic procedure to study the uncertainties of PDF's and their physics predictions, while incorporating all the experimental constraints used in the previous CTEQ analysis [1]. An effective χ^2 function, called χ^2_{global} , is used not only to extract the "best fit," but also to explore the neighborhood of the global minimum in order to quantify the uncertainties, as is done in the classic error matrix approach. Two key ingredients make this possible: (i) a recently established iterative procedure [17] that yields a reliable calculation of the Hessian matrix in the complex global analysis environment, and (ii) the use of appropriately normalized eigenvectors to greatly improve the accuracy and utility of the analysis.

The Hessian approach is based on a quadratic approximation to χ^2_{global} in the neighborhood of the minimum that defines the best fit. It yields a set of PDF's associated with the eigenvectors of the Hessian, which characterize the PDF parameter space in the neighborhood of the global minimum in a process-independent way. In a companion paper, referred to here as "the preceding paper" [18], we present a complementary process-dependent method that studies χ^2_{global} as a function of whatever specific physical variable is of interest. That approach is based on the Lagrange multiplier (LM) method [17], which does not require a quadratic approximation to χ^2_{global} , and hence is more robust; but, being focused on a single variable (or a few variables in a generalized formulation) it does not provide complete information about the neighborhood of the minimum. We use the LM method to verify the reliability of the Hessian calculations, as discussed in Sec. V. Further tests of the quadratic approximation are described in Appendix B.

The outline of the paper is as follows. In Sec. II we summarize the global analysis that underlies the study, and define the function χ^2_{global} that plays the leading role. In Sec. III we explore the quality of fit in the neighborhood of the minimum. We derive the eigenvector basis sets, and show how they can be used to calculate the uncertainty on any quantity that depends on the parton distributions. In Sec. IV we apply the formalism to derive uncertainties of the PDF parameters and of the PDF's themselves. In Sec. V we illustrate the method further by finding the uncertainties on predictions for the rapidity distribution of W production, and the correlation between W and Z production cross sections. We summarize our results in Sec. VI. Two appendixes provide details on the estimate of overall tolerance for the effective χ^2_{global} function, and on the validity of the quadratic approximation inherent in the Hessian method. Two further appendixes supply explicit tables of the coefficients that define the best fit and the eigenvector basis sets. The mathematical methods used here have been described in detail in [17]. Some preliminary results have also appeared in [7,8].

II. GLOBAL QCD ANALYSIS AND EFFECTIVE CHI SQUARED

Global χ^2 analysis is a practical and effective way to combine a large number of experimental constraints. In this section, we describe the main features of the global QCD analysis, and explain how we quantify its uncertainties through the behavior of χ^2_{elobal} .

A. Experimental and theoretical inputs

We use the same experimental input as the CTEQ5 analysis [1]: 15 data sets on neutral-current and charged-current deep inelastic scattering (DIS), Drell-Yan lepton pair production, forward backward lepton asymmetry from W production, and high p_T inclusive jets, as listed in Table I of Appendix A. The total number of data points is 1295, after cuts such as Q>2 GeV and W>4 GeV in DIS designed to reduce the influence of power-law suppressed corrections and other sources of theoretical error. The experimental precision and the information available on systematic errors vary widely among the experiments, which presents difficulties for the effort to quantify the uncertainties of the results.

The theory input is next-to-leading-order (NLO) perturbative QCD. The theory has systematic uncertainties due to uncalculated higher-order QCD corrections, including possible resummation effects near kinematic boundaries, powersuppressed corrections, and nuclear effects in neutrino data on heavy targets. These uncertainties—even more than the experimental ones—are difficult to quantify.

The theory contains free parameters $\{a_i\}=\{a_1,\ldots,a_d\}$ defined below that characterize the nonperturbative input to the analysis. Fitting theory to experiment determines these $\{a_i\}$ and thereby the PDF's. The uncertainty of the result due to experimental and theoretical errors is assessed in our analysis by an assumption on the permissible range of $\Delta \chi^2$ for the fit, which is discussed in Sec. II D.

B. Parametrization of PDF's

The PDF's are specified in a parametrized form at a fixed low-energy scale Q_0 , which we choose to be 1 GeV. The PDF's at all higher Q are determined from these by the NLO perturbative QCD evolution equations. The functional forms we use are

$$f(x,Q_0) = A_0 x^{A_1} (1-x)^{A_2} (1+A_3 x^{A_4})$$
(1)

with independent parameters for parton flavor combinations $u_v \equiv u - \overline{u}$, $d_v \equiv d - \overline{d}$, g, and $\overline{u} + \overline{d}$. We assume $s = \overline{s} = 0.2(\overline{u} + \overline{d})$ at Q_0 . A somewhat different parametrization for the $\overline{d}/\overline{u}$ ratio is adopted to better fit the current data:

$$\overline{d}(x,Q_0)/\overline{u}(x,Q_0) = B_0 x^{B_1} (1-x)^{B_2} + (1+B_3 x)(1-x)^{B_4}.$$
(2)

The specific functional forms are not crucial, as long as the parametrization is flexible enough to include the behavior of the true—but unknown—PDF's at the level of accuracy to which they can currently be determined. The parametrization should also provide additional flexibility to model the degrees of freedom that remain indeterminate. On the other hand, too much flexibility in the parametrization leaves some parameters poorly determined at the minimum of χ^2 . To avoid that problem, some parameters in the present study were frozen at particular values.

The number of free parameters has increased over the years, as the accuracy and diversity of the global data set has gradually improved. A useful feature of the Hessian approach is the feedback it provides to aid in refining the parametrization, as we discuss in Sec. IV A. The current analysis uses a total of d=16 independent parameters, referred to generically as $\{a_i\}$. Their best-fit values, together with the fixed ones, are listed in Table III of Appendix C. (Some of the fit parameters a_i are defined by simple functions of their related PDF shape parameters A_i or B_i , as indicated in the table, to keep their relevant magnitudes in a similar range, or to enforce positivity of the input PDF's, etc.) The set of fit parameters $\{a_i\}$ could also include parameters associated with correlated experimental errors, such as an unknown normalization error that is common to all of the data points in a particular experiment; however, such parameters were kept fixed for simplicity in this initial study. The QCD coupling was similarly fixed at $\alpha_s(M_Z) = 0.118$.

C. Effective chi-squared function

Our analysis is based on an effective global chi-squared function that measures the quality of the fit between theory and experiment:

$$\chi^2_{\text{global}} = \sum_n w_n \chi^2_n, \qquad (3)$$

where n labels the 15 different data sets.

The weight factors w_n in Eq. (3), with default value 1, are a generalization of the selection process that must begin any global analysis, where one decides which data sets to include (w=1) or exclude (w=0). For instance, we include neutrino DIS data (because it contains crucial constraints on the PDF's, although it requires model-dependent nuclear target corrections) but we exclude direct photon data (which would help to constrain the gluon distribution, but suffers from delicate sensitivity to k_{\perp} effects from multiple soft gluon emission). The w_n can be used to emphasize particular experiments that provide unique physical information, or to deemphasize experiments when there are reasons to suspect unquantified theoretical or experimental systematic errors (e.g., in comparison to similar experiments). Subjectivity such as this choice of weights is not covered by Gaussian statistics, but is a part of the more general Bayesian approach, and is, in spirit, a familiar aspect of estimating systematic errors within an experiment, or in averaging experimental results that are marginally consistent.

The generic form for the individual contributions in Eq. (3) is

$$\chi_n^2 = \sum_I \left(\frac{D_{nI} - T_{nI}}{\sigma_{nI}} \right)^2, \tag{4}$$

where T_{nI} , D_{nI} , and σ_{nI} are the theory value, data value, and uncertainty for data point *I* of data set (or "experiment") *n*. In practice, Eq. (4) is generalized to include correlated errors such as overall normalization factors, or even the full experimental error correlation matrix if it is available [18].

The value of χ^2_{global} depends on the PDF set, which we denote by *S*. We stress that $\chi^2_{global}(S)$ is an "effective χ^2 ," whose purpose is to measure how well the data are fit by the theory when the PDF's are defined by the parameter set $\{a_i(S)\}$. We use $\chi^2_{global}(S)$ to study how the quality of fit varies with the PDF parameters, but we do not assign *a priori* statistical significance to specific values of it—e.g., in the manner that would be appropriate to an ideal chi-squared distribution—since the experimental and theoretical inputs are often far from being ideal, as discussed earlier.

D. Global minimum and its neighborhood

Having specified the effective χ^2 function, we find the parameter set that minimizes it to obtain a "best estimate" of the true PDF's. This PDF set is denoted by S_0 .¹ The parameter values that characterize S_0 are listed in Table III of Appendix C.

To study the uncertainties, we must explore the variation of χ^2_{global} in the neighborhood of its minimum, rather than focusing only on S_0 as has been done in the past. Moving the parameters away from the minimum increases χ^2_{global} by an amount $\Delta \chi^2_{global}$. It is natural to define the relevant neighborhood of the global minimum as

$$\Delta \chi^2_{\text{global}} \leq T^2, \tag{5}$$

where *T* is a *tolerance parameter*. The Hessian formalism developed in Sec. III provides a reliable and efficient method of calculating the variation of all predictions of PDF's in this neighborhood, as long as *T* is within the range where a quadratic expansion of χ^2_{global} , in terms of the PDF parameters, is adequate.

In order to quantify the uncertainties of physical predictions that depend on PDF's, one must choose the tolerance parameter *T* to correspond to the region of "acceptable fits." Broadly speaking, the order of magnitude of *T* for our choice of χ^2_{global} is already suggested by self-consistency considerations: Our fundamental assumption—that the 15 data sets used in the global analysis are individually *acceptable* and mutually *compatible*, in spite of departures from ideal statistical expectations exhibited by some of the individual data

¹It is very similar to the CTEQ5M1 set [1], with minor differences arising from the improved parametrization (2) for $\overline{d}/\overline{u}$.



FIG. 1. Illustration of the basic ideas of our implementation of the Hessian method. An iterative procedure [17] diagonalizes the Hessian matrix and rescales the eigenvectors to adapt the step sizes to their natural scale. The solid points represent the resulting eigenvector basis PDF's described in Sec. III C. Point $\mathbf{p}(\mathbf{i})$ is explained in Sec. IV A.

sets, as well as signs of incompatibility between some of them if the errors are interpreted according to strict statistical rules [12]—must, in this effective χ^2 approach, imply a value of *T* substantially larger than that of ideal expectations. More quantitatively, estimates of *T* have been carried out in the preceding paper [18], based on the comparison of $\Delta \chi^2_{\text{global}}$ with detailed studies of experimental constraints on specific physical quantities. The estimates of *T* will be discussed more extensively in Sec. V, where applications are presented, and in Appendix A. For the development of the formalism in the next section, it suffices to know that (i) the order of magnitude of these estimates is

$$T \approx 10$$
 to 15, (6)

and (ii) the master formulas, given in Sec. III D, imply that all uncertainties are proportional to T, hence the results are fully scalable, according to the best available estimate of T.

III. THE HESSIAN FORMALISM

The most efficient approach to studying uncertainties in a global analysis of data is through a quadratic expansion of the χ^2 function about its global minimum.² This is the well-known error matrix or Hessian method. Although the method is standard, its application to PDF analysis has, so far, been hindered by technical problems created by the complexity of the theoretical and experimental inputs. Those technical problems have recently been overcome [17].

The Hessian matrix is the matrix of second derivatives of χ^2 at the minimum. In our implementation, the eigenvectors of the Hessian matrix play a central role. They are used both for an accurate evaluation of the Hessian itself, via the iterative method of [17], and to produce an eigenvector basis set of PDF's from which uncertainties of all physical observables can be calculated. The basis PDF's provide an optimized representation of the parameter space in the neighborhood of the minimum χ^2 .

The general idea of our approach is illustrated conceptually in Fig. 1. Every PDF set *S* corresponds to a point in the *d*-dimensional PDF parameter space. It can be specified by the original parton shape parameters $\{a_i(S)\}$ defined in Sec. II B, as illustrated in Fig. 1(a); or by the eigenvector basis coordinates $\{z_k(S)\}$, which specify the components of *S* along the eigenvector basis PDF's that will be introduced in Sec. III C, as illustrated in Fig. 1(b). The solid points in both Figs. 1(a) and 1(b) represent the basis PDF sets.

A. Quadratic approximation and the Hessian matrix

The standard error matrix approach begins with a Taylor series expansion of $\chi^2_{\text{global}}(S)$ around its minimum S_0 , keeping only the leading terms. This produces a quadratic form in the displacements from the minimum:

$$\Delta \chi^2 = \chi^2 - \chi_0^2 = \frac{1}{2} \sum_{i=1}^d \sum_{j=1}^d H_{ij}(a_i - a_i^0)(a_j - a_j^0), \quad (7)$$

where $\chi_0^2 = \chi^2(S_0)$ is the value at the minimum, $\{a_j^0\} = \{a_j(S_0)\}$ is its location, and $\{a_j\} = \{a_j(S)\}$. We have dropped the subscript "global" on χ^2 for simplicity. We also suppress the PDF argument (*S*) in χ^2 and $\{a_i\}$ here and elsewhere when it is not needed to make the discussion clear.

The Hessian matrix H_{ij} has a complete set of orthonormal eigenvectors v_{ik} defined by

$$\sum_{j=1}^{d} H_{ij} v_{jk} = \epsilon_k v_{ik}, \qquad (8)$$

$$\sum_{i=1}^{d} v_{il} v_{ik} = \delta_{lk}, \qquad (9)$$

where $\{\epsilon_k\}$ are the eigenvalues and δ_{lk} is the unit matrix. Displacements from the minimum are conveniently expressed in terms of the eigenvectors by

$$a_i - a_i^0 = \sum_{k=1}^d v_{ik} s_k z_k, \qquad (10)$$

²The Lagrange multiplier method [17,18] is a complementary approach that avoids the quadratic approximation.



FIG. 2. Distribution of eigenvalues of the Hessian matrix for fits using d=13, 16 (standard), and 18 free PDF parameters.

where scale factors s_k are introduced to normalize the new parameters z_k such that

$$\Delta \chi^2 = \sum_{k=1}^d z_k^2. \tag{11}$$

With this normalization, the relevant neighborhood (5) of the global minimum corresponds to the interior of a hypersphere of radius *T*:

$$\sum_{k=1}^{d} z_k^2 \leqslant T^2.$$
(12)

The scale factors s_k are approximately equal to $\sqrt{2/\epsilon_k}$, as is explained in Appendix B.

The transformation (10) is illustrated conceptually in Fig. 1, where $\{k, l\}$ label two of the eigenvector directions. One of the eigenvectors \mathbf{v}_l is shown both in the original parameter basis and in the normalized eigenvector basis.

B. Eigenvalues of the Hessian matrix

The square of the distance in parameter space from the minimum of χ^2 is

$$\sum_{i=1}^{d} (a_i - a_i^0)^2 = \sum_{k=1}^{d} (s_k z_k)^2$$
(13)

by Eqs. (9), (10). Because $s_k \approx \sqrt{2/\epsilon_k}$, an eigenvector with large eigenvalue ϵ_k , therefore, corresponds to a "steep direction" in $\{a_i\}$ space, i.e., a direction in which χ^2 rises rapidly, making the parameters tightly constrained by the data. The opposite is an eigenvector with small ϵ_k , which corresponds to a "shallow direction," for which the criterion $\Delta \chi^2 \leq T^2$ permits considerable motion—as is the case for \mathbf{v}_l illustrated in Fig. 1. The distribution of eigenvalues, ordered from largest to smallest, is shown in Fig. 2. Interestingly, the distribution is approximately linear in $\log \epsilon$. The eigenvalues span an enormous range, which is understandable because the large global data set includes powerful constraints—particularly on combinations of parameters that control the quark distributions at moderate *x*—leading to steep directions; while free parameters have purposely been added to Eqs. (1) and (2) to the point where some of them are at the edge of being unconstrained by the data, leading to shallow directions.

Figure 2 also shows how the range of eigenvalues expands or contracts if the number of adjustable parameters is changed; the 16-parameter fit is the standard one used in most of this paper; the 18-parameter fit is defined by allowing $A_1^{u_v} \neq A_1^{d_v}$ and $A_3^g \neq 0$ with $A_4^g = 6$; the 13-parameter fit is defined by $A_4^{u_v} = A_4^{d_v} = 1$ and $A_3^{\overline{d}+\overline{u}} = 0$. The range spanned by the eigenvalues increases with the dimension *d* of the parameter space [roughly as $\log(\epsilon_1/\epsilon_d) \propto d^{0.4}$].

The large $(10^{\circ}:1)$ range spanned by the eigenvalues makes the smaller eigenvalues and their eigenvectors very sensitive to fine details of the Hessian matrix, making it difficult to compute H_{ij} with sufficient accuracy. This technical problem hindered the use of Hessian or error matrix methods in global QCD analysis in the past. The problem has been tamed by an iterative method introduced in [17], which computes the eigenvalues and eigenvectors by successive approximations that converge even in the presence of numerical noise and nonquadratic contributions to³ χ^2 .

The Hessian method relies on the quadratic approximation (7) to the effective χ^2 function. We have extensively tested this approximation in the neighborhood of interest by comparing it with the exacet χ^2_{global} . The results are satisfactory, as shown in Appendix B, which also explains how the approximation is improved by adjusting the scale factors s_k for the shallow directions.

C. PDF eigenvector basis sets S_l^{\pm}

The *k*th eigenvector of the Hessian matrix has component v_{ik} along the *i*th direction in the original parameter space, according to Eq. (8). Thus v_{ik} is the orthogonal matrix that transforms between the original parameter basis and the eigenvector basis. For our application, it is more convenient to work with coordinates $\{z_k\}$ that are normalized by the scale factors $\{s_k\}$ of Eq. (10), rather than the "raw" coordinates of the the eigenvector basis. Thus we use the matrix

$$M_{ik} \equiv v_{ik} s_k \tag{14}$$

rather than v_{ik} itself. M_{ik} defines the transformation between the two descriptions that are depicted conceptually in Fig. 1:

$$a_i - a_i^0 = \sum_{k=1}^d M_{ik} z_k.$$
 (15)

³The iterative algorithm is implemented as an extension to the widely used CERNLIB program MINUIT [30]. The code is available at http://www.pa.msu.edu/~pumplin/iterate/

It contains information about the physics in the global fit, together with information related to the choice of parametrization, and is a good object to study for insight into how the parametrization might be improved, as we discuss in Sec. IV A.

The eigenvectors provide an optimized orthonormal basis in the PDF parameter space, which leads to a simple parametrization of the parton distributions in the neighborhood of the global minimum S_0 . In the remainder of this section, we show how to construct these *eigenvector basis* PDF's $\{S_l^{\pm}, l=1, \ldots, d\}$, and in the following section, we show how they can be used to calculate the uncertainty of any chosen variable X(S).

The *eigenvector basis* sets S_l^{\pm} are defined by displacements of a standard magnitude *t* "up" or "down" along each of the *d* eigenvector directions. Their coordinates in the *z* basis are thus

$$z_k(S_l^{\pm}) = \pm t \,\delta_{kl} \,. \tag{16}$$

More explicitly, S_1^+ is defined by $(z_1, \ldots, z_d) = (t, 0, \ldots, 0)$, etc. We make displacements in both directions along each eigenvector to improve accuracy; which direction is called "up" is totally arbitrary. As a practical matter, we choose t=5 for the displacement distance. This choice improves the accuracy of the quadratic approximation by working with displacements that have about the same size as those needed in applications.⁴

The $\{a_i\}$ parameters that specify the eigenvector basis sets S_i^{\pm} are given by

$$a_i(S_l^{\pm}) - a_i^0 = \pm t M_{il}, \qquad (17)$$

by Eqs. (15), (16). Hence

$$a_i(S_l^+) - a_i(S_l^-) = 2tM_{il}.$$
 (18)

Interpreted as a difference equation, this shows directly that the element M_{il} of the transformation matrix is equal to the gradient of parameter a_i along the direction of ${}^5 z_l$.

Basis PDF sets along two of the eigenvector directions are illustrated conceptually in Figs. 1(a) and 1(b) as solid points displaced from the global minimum set S_0 . The coefficients that specify all of the sets S_l^{\pm} are listed in Table IV of Appendix D.

D. Master equations for calculating uncertainties using the eigenvector basis sets $\{S_L^{\pm}\}$

Let X(S) be any variable that depends on the PDF's. It can be a physical quantity such as the W production cross section; or a particular PDF such as the gluon distribution at specific x and Q values; or even one of the PDF parameters a_i . All of these cases will be used as examples in Secs. IV and V.

The best-fit estimate for *X* is $X^0 = X(S_0)$. To find the uncertainty, it is only necessary to evaluate *X* for each of the 2*d* sets $\{S_l^{\pm}\}$. The gradient of *X* in the *z* representation can then be calculated, using a linear approximation that is essential to the Hessian method, by

$$\frac{\partial X}{\partial z_k} = \frac{X(S_k^+) - X(S_k^-)}{2t},\tag{19}$$

where t is the scale used to define $\{S_l^{\pm}\}$ in Eq. (16). It is useful to define

$$D_k(X) = X(S_k^+) - X(S_k^-), \qquad (20)$$

$$D(X) = \left(\sum_{k=1}^{d} \left[D_k(X)\right]^2\right)^{1/2},$$
 (21)

$$\hat{D}_k(X) = D_k(X) / D(X), \qquad (22)$$

so that $D_k(X)$ is a vector in the gradient direction and $\hat{D}_k(X)$ is the unit vector in that direction.

The gradient direction is the direction in which X varies most rapidly, so the largest variations in X permitted by Eq. (12) are obtained by displacement from the global minimum S_0 in the gradient direction by a distance $\pm T$. Hence

$$\Delta X = \sum_{k=1}^{d} (T\hat{D}_k) \frac{\partial X}{\partial z_k}.$$
 (23)

From this, using Eqs. (19)–(22), we obtain the *master equation* for calculating uncertainties,

$$\Delta X = \frac{T}{2t} D(X). \tag{24}$$

This equation is applied to obtain numerical results in Secs. IV and V.

For applications, it is often important to also construct the PDF sets S_X^+ and S_X^- that achieve the extreme values $X = X^0 \pm \Delta X$. Their *z* coordinates are

$$z_k(S_X^{\pm}) = \pm T\hat{D}_k(X), \qquad (25)$$

which follows from the derivation of Eq. (24). Their physical parameters $\{a_i\}$ then follow from Eqs. (15) and (18):

$$a_i(S_X^{\pm}) - a_i^0 = \frac{\pm T}{2t} \sum_{k=1}^d \hat{D}_k(X) [a_i(S_k^+) - a_i(S_k^-)]. \quad (26)$$

⁴The value chosen for *t* is somewhat smaller than the typical *T* given in Eq. (6) because in applications, the component of displacement along a given eigenvector direction will generally be smaller than the total displacement.

⁵Technically, we calculate the orthogonal matrix v_{ij} using displacements that give $\Delta \chi^2 \approx 5$, where the iterative procedure [17] converges well. The eigenvectors are then scaled up by an amount that is adjusted to make $\Delta \chi^2 = 25$ exactly for each S_k^{\pm} to improve the quadratic approximation.



In practice, we calculate the parameters for S_X^+ and S_X^- by applying Eq. (26) directly to the parton shape parameters $\ln A_0^{u_v}, A_1^{u_v}, \ldots$ listed in Table IV, except that the normalization factors $A_0^{u_v}, A_0^{d_v}$, and A_0^g are computed from the momentum sum rule and quark number sum rules

$$\sum_{i} \int_{0}^{1} x f_{i}(x) dx = 1,$$
(27)

$$\int_{0}^{1} u_{v}(x) dx = 2, \quad \int_{0}^{1} d_{v}(x) dx = 1$$
(28)

to ensure that those sum rules are satisfied exactly.

IV. UNCERTAINTIES OF PARTON DISTRIBUTIONS

A. Uncertainties of the PDF parameters $\{a_i\}$

As a useful as well as illustrative application of the general formalism, let us find the uncertainties on the physical PDF parameters a_i . We only need to follow the steps of Sec. III D. Letting $X = a_i$ for a particular *i*, Eqs. (20) and (18) give

$$D_k(a_i) = a_i(S_k^+) - a_i(S_k^-) = 2tM_{ik}.$$
 (29)

The uncertainty on a_i in the global analysis follows from the master equation (24):

$$\Delta a_i = T \left(\sum_k M_{ik}^2\right)^{1/2}.$$
(30)

The parameter sets $\{a_j(a_i^+)\}$ and $\{a_j(a_i^-)\}$ that produce the extreme values of a_i can be found using Eq. (26). In the conceptual Fig. 1, the parton distribution set with the largest value of a_i for $\Delta \chi^2_{\text{global}} = T^2$ is depicted as point $\mathbf{p}(\mathbf{i})$. The uncertainties $\{\Delta a_i\}$ of the standard parameter set, cal-

The uncertainties $\{\Delta a_i\}$ of the standard parameter set, calculated from Eq. (30) with T=5 are listed along with the central values $\{a_i^0\}$ in Table III. To test the quadratic approximation, asymmetric errors are also listed. These are defined by displacements in the gradient direction (29) that are adjusted to make $\Delta \chi^2$ exactly equal to $T^2=25$. They agree quite well with the errors calculated using Eq. (30), which shows that the quadratic approximation is adequate for our purposes. FIG. 3. Two extreme gluon distributions (left) and *u*-quark distributions (right) for Q=2 GeV (long dash) and Q=100 GeV (short dash) with T=10. Each curve is calculated to give the minimum or maximum value for some particular *x*. The entire allowed region, which is the envelope of all such curves, is shaded.

Table III also lists the components of the displacement vectors $z_k(a_i^+)$ of Eq. (25) which have been renormalized to $\Sigma z_k^2 = 1$. These reveal which features of the PDF's are governed most strongly by specific eigenvector directions. The table is divided into sections according to the various flavor combinations that are parametrized. One can see, for example, that the flattest direction z_{16} is strongly related to the gluon parameters A_1^g and A_2^g , confirming that the gluon distribution at $Q_0=1$ GeV is a highly uncertain aspect of the PDF's. The second-flattest direction z_{15} relates mainly to the $\overline{d}/\overline{u}$ ratio, as seen by the large components along z_{15} for $B_0^{\overline{d}/\overline{u}}$ and $B_1^{\overline{d}/\overline{u}}$. Meanwhile, the steepest direction z_1 mainly influences the valence quark distribution via $A_1^{u_v}$.

All of the parametrized aspects of the PDF's at Q_0 , namely, u_v , d_v , g, $\overline{d} + \overline{u}$, and $\overline{d}/\overline{u}$ receive substantial contributions from the four-flattest directions 13–16, which shows that the current global data set could not support the extraction of much finer detail in the PDF's. This can be confirmed by noting that the error ranges of the individual parameters a_i are not small.

B. Uncertainties of the PDF's

The uncertainty range of the PDF's themselves can also be explored using the eigenvector method. For example, letting the gluon distribution g(x,Q) at some specific values of (x,Q) be the variable X that is extremized by the method of Sec. III D leads to the extreme gluon distributions shown in the left-hand side of Fig. 3. The envelope of such curves, obtained by extremizing at a variety of x values at fixed Q, is shown by the shaded region, which is defined by T=10, i.e., by allowing χ^2_{global} up to 100 above its minimum value.

The right-hand side of Fig. 3 similarly shows the allowed region and two specific cases for the u-quark distribution. The uncertainty is much smaller than for the gluon, reflecting the large amount of experimental data included in the global analysis that is sensitive to the u quark through the square of its electric charge.

The dependence on x in these figures is plotted as a function of $x^{1/3}$ to better display the region of current experimental interest. The values are weighted by a factor $x^{5/3}$, which makes the area under each curve proportional to its contribution to the momentum sum rule. Note that the uncertainty decreases markedly with increasing Q as a result of evolu-



tion. Also note that the gluon distribution is large and fairly mo

well determined at smaller x values and large Q—the region that will be vital for physics at the CERN Large Hadron Collider (LHC).

Figure 4 displays similar information for Q=10 GeV, expressed as the fractional uncertainty as a function of log *x*. It shows that the gluon distribution becomes very uncertain at large *x*, e.g., x>0.25. (At x>0.6, where the distribution is extremely small, the lower envelope of fractional uncertainty begins to rise. This is an artifact of the parametrization with $A_3^g=0$; making the parametrization more flexible by freeing A_3^g with $A_4^g=6$ leads to a broader allowed range indicated by the dotted curves.)

The boundaries of the allowed regions for the PDF's are not themselves possible shapes for the PDF's, since if a particular distribution is extremely high at one value of x, it will be low at other values. This can be seen most clearly in the gluon distributions of Figs. 3 and 4, where the extreme PDF's shown push the envelope on the high side in one region of x, and on the low side in another.

V. UNCERTAINTIES OF PHYSICAL PREDICTIONS

In applying the Hessian method to study the uncertainties of physical observables due to PDF's, it is important to understand how the predictions depend on the tolerance parameter *T*, and how well *T* can be determined. We discuss these issues first, and then proceed to illustrate the utility of this method by examining the predictions for the rapidity distribution of *W* and *Z* boson production as well as the correlation of *W* and *Z* cross sections in $p\overline{p}$ collisions.

First, note that the uncertainties of all predictions are linearly dependent on the tolerance parameter *T* in the Hessian approach, by the master formula (24); hence they are easily scalable. The appropriate value of *T* is determined, in principle, by the region of "acceptable fits" or "reasonable agreement with the global data sets" in the PDF parameter space. Physical quantities calculated from PDF sets within this region will range over the values that can be considered "likely" predictions. As discussed in the introductory sections, the complexity of the experimental and theoretical input to the function χ^2_{global} in the global analysis, in particular, the unknown systematic errors reflected in apparent abnormalities of some reported experimental errors as well as seeming incompatibilities between some data sets, makes it difficult to assign an unambiguous value to *T*. However, as



mentioned in Sec. II D, self-consistency considerations inherent to our basic assumption that the 15 data sets used in the global analysis are acceptable and compatible, in conjunction with the detailed comparison to experiment conducted in the preceding paper [18] using the same χ^2_{global} function, yield a best estimate of $T \approx 10$ to 15 [(Eq. 6)]. Details of these considerations are discussed in Appendix A.

Of the estimates of *T* described there, the most quantitative one is based on the algorithm of the preceding paper [18] to combine 90% confidence level error bands from the 15 individual data sets for any specific physical variable such as the total production cross section of *W* or *Z* at the Fermilab Tevatron or LHC (see Appendix A for a summary, and the preceding paper [18] for the detailed analysis). For the case of σ_W^{TeV} , the uncertainty according to the specific algorithm is ±4%, corresponding to $T \sim 13$. With our working hypothesis $T \approx 10-15$, the range of the uncertainty of σ_W^{TeV} will be ±3.3% to ±4.9%.

The numerical results on applications presented in the following sections are obtained with the same choice of T as in Sec. IV, i.e., T = 10. Bearing in the mind the linear dependence of the uncertainties on T, one can easily scale these up by the appropriate factor if a more conservative estimate for the uncertainty of any of the physical quantities is desired. We should also note that the experimental data sets used in this analysis are continuously evolving. Some data sets (cf., Table I in Appendix A) will soon be updated (H1, ZEUS) or replaced (CCFR).⁶ In addition, theoretical uncertainties have yet to be systematically studied and incorporated. Therefore, the specific results presented in this paper should be considered more as a demonstration of the method rather than definitive predictions. The latter will be refined as new and better inputs become available.

A. Rapidity distribution for W production

Figure 5 shows the predicted rapidity distribution $d\sigma/dy$ for W^+ production in $p\bar{p}$ collisions at $\sqrt{s} = 1.8$ TeV. The cross section is not symmetric in y because of the strong contribution from the valence u quark in the proton—indeed, the forward backward asymmetry produces an observable

⁶Compare talks presented by these collaborations at DIS2000 *Workshop on Deep Inelastic Scattering and Related Topics*, Liverpool, England, 2000.



asymmetry in the distribution of leptons from W decay, which provides an important handle on flavor ratios in the current global analysis.

The left-hand side of Fig. 5 shows the six rapidity distributions that give the extremes (up or down) of the integrated cross section $\sigma = \int (d\sigma/dy) dy$, the first moment $\langle y \rangle$, or the second moment $\langle y^2 \rangle$, as calculated using the Hessian formalism for T=10. To show the differences more clearly, the right-hand side shows the difference between each of these rapidity distributions and the best-fit distribution.

Figure 6 shows three of the same difference curves as in Fig. 5 along with results obtained using the Lagrange multiplier method of the preceding paper [18]. The good agreement shows that the Hessian formalism, with its quadratic approximation (7), works well at least for this application.

Figure 7 shows the same three curves from Fig. 5, together with 6 random choices of the PDF's with $\Delta \chi^2_{\text{global}}$ = 100. These random sets were obtained by choosing random directions in $\{z_i\}$ space and displacing the parameters from the minimum in those directions until χ^2_{global} has increased by 100. Note that none of this small number of ran-



FIG. 6. Comparison with the Lagrange multiplier method: three of the six curves from Fig. 5 (corresponding to maximum σ , $\langle y \rangle$, or $\langle y^2 \rangle$) are shown, together with the result of the exact LM method for $\Delta \chi^2_{\text{global}} = 100$.

FIG. 5. Left: Predicted rapidity distribution for $p\bar{p} \rightarrow W^+ + X$ at $\sqrt{s} = 1.8$ TeV. The curves are extreme predictions for the integrated cross section σ (solid), or the rapidity moments $\langle y \rangle$ (long dash), or $\langle y^2 \rangle$ (short dash). Right: same except the best-fit prediction is subtracted to show the details better.

dom sets give good approximations to the three extreme curves. This is not really surprising, since the extrema are produced by displacements in specific (gradient) directions, and in 16-dimensional space, the component of a random unit vector along any specific direction is likely to be small. But it indicates that producing large numbers of random sets would at best be an inefficient way to unearth the extreme behaviors.

B. Correlation between W and Z cross sections

One can ask what are the error limits on two quantities X and Y simultaneously, according to the $\Delta \chi^2_{global} < T$ criterion. In the Hessian approximation, the boundary of the allowed region is an ellipse, as shown in [17]. The ellipse can be expressed elegantly in a "Lissajous figure" form

$$X = X^0 + \Delta X \sin(\theta + \phi),$$

$$Y = Y^0 + \Delta Y \sin(\theta), \qquad (31)$$



FIG. 7. Comparison with random methods: three of the six curves from Fig. 5 (corresponding to maximum σ , $\langle y \rangle$, or $\langle y^2 \rangle$ for $\Delta \chi^2_{\text{global}} = 100$) are shown, together with results from PDF sets that are obtained by displacement in 6 random directions of $\{z_i\}$ space by $\Delta \chi^2_{\text{global}} = 100$.



FIG. 8. Error ellipse for predicted *W* and *Z* boson production in $p\bar{p}$ collisions at 1.8 TeV. The error limit (T=10) of the prediction is the interior of the ellipse. Error bars show data from DØ (dotted) and CDF (dashed). The dot-dash error bars show the result of reinterpreting the CDF data by using the same assumption for luminosity as DØ [31].

where $0 \le \theta \le 2\pi$ traces out the boundary. The shape of the ellipse is governed by the phase angle ϕ , which is given by the dot product between the gradient vectors for *X* and *Y* in $\{z_k\}$ space:

$$\cos\phi = \sum_{k=1}^{d} \hat{D}_{k}(X)\hat{D}_{k}(Y),$$
 (32)

where $\hat{D}_i(X)$ and $\hat{D}_i(Y)$ are defined by Eq. (22).

As an example of this, T=10 error limits for W^{\pm} and Z^0 production at the Tevatron are shown in Fig. 8. The error limits on the separate predictions for these cross sections are each about 3.3% for T=10. The predictions are strongly correlated ($\cos \phi = 0.60$), in part because the same quark distributions—in different combinations—are responsible for both W and Z production, and in part because the uncertainties of all the quark distributions are negatively correlated with the more uncertain gluon distribution, and hence positively correlated with each other.

The *W* and *Z* cross sections from Collider Detector at Fermilab (CDF) (dashed) and DØ (dotted) are also shown in Fig. 8 [31]. (The measured quantities $\sigma_W \cdot B_{W \to e\nu}$ and $\sigma_Z \cdot B_{Z \to e^+e^-}$ were converted to σ_W and σ_Z using world average values for the branching ratios [16]; the measured CDF and DØ branching ratios for *W* agree with the world average to within about 1%.) The data points are shown in the form of error bars defined by combining statistical and systematic errors (including the errors in decay branching ratios) in quadrature. The errors in these measurements are also highly correlated, in part through the uncertainty in overall luminosity, which both cross sections are proportional to—so the experimental points would also be better represented by ellipses. The two experiments in fact use different assumptions for the inelastic $p\bar{p}$ cross section which measures the luminosity; CDF uses its own measurement of it, while DØ uses the world average. The dot-dashed data point shows the result of reinterpreting the CDF point by scaling the luminosity down by a factor of 1.062 to correspond to the world average $p\bar{p}$ cross section [31].

VI. SUMMARY AND CONCLUDING REMARKS

Experience over the past two decades has shown that minimizing a suitably defined χ^2_{global} is an effective way to extract parton distribution functions consistent with experimental constraints in the PQCD framework. The goal of this paper has been to expand the scope of such analysis to make quantitative estimates of the uncertainties of PDF's and their predictions, by examining the behavior of χ^2_{global} in the neighborhood of the minimum. The techniques developed in Ref. [17] allow us to apply the traditional error matrix approach reliably in the complex global analysis environment. The eigenvectors of the Hessian (inverse of the error matrix) play a crucial role, both in the adaptive procedure to accurately calculate the Hessian itself, and in the derivation of the compact master formula for determining the uncertainties of parton distributions and their predictions, Eq. (24).

Our principal results are (i) the formalism developed in Sec. III D, leading to the master formulas, and (ii) the best-fit parton distribution set S_0 plus the 2*d* eigenvector basis sets S_k^{\pm} presented in Sec. III C, which are used in applications of the master formula (24). The uncertainties are proportional to *T*, the tolerance parameter for $\Delta \chi^2_{global}$. We present several estimates, based on current experimental and theoretical input, that suggest *T* is in the range 10–15. It is important to note, however, that this estimate can, and should, be refined in the near future. First, several important data sets used in the global analysis will soon be updated or replaced (cf. Sec. V, footnote 5). Second, there are other sources of uncertainties, which have yet to be studied and included in the analysis in a full evaluation of uncertainties. (The work of Botje [10] describes possible ways to incorporate some of these.)

This paper, focusing on the presentation of a new formalism and its utility, represents the first step in a long-term project to investigate the uncertainties of predictions dependent upon parton distributions. We plan to perform a series of studies on processes in precision SM measurements (such as the W mass) and in new physics searches (such as Higgs production cross section) which are sensitive to the parton distributions, at the Tevatron and LHC.

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TABLE I. Data sets used in the global analysis. If experiment *n* is omitted, Δ_n denotes the amount by which χ^2 for the remaining 14 experiments can be reduced by readjusting the fit parameters.

Expt n	Process	N_n	Name	Ref.	Δ_n
1	DIS $F_2(\mu p)$	168	BCDMS	[19]	19.7
2	DIS $F_2(\mu d)$	156	BCDMS	[19]	4.5
3	DIS $F_2(ep)$	172	H1	[20]	3.7
4	DIS $F_2(ep)$	186	ZEUS	[21]	9.7
5	DIS $F_2(\mu p)$	104	NMC	[22]	7.8
6	DIS $\mu p/\mu n$	123	NMC	[22]	1.9
7	DIS $\mu p/\mu n$	13	NMC	[22]	13.9
8	DIS $F_2(\nu p)$	87	CCFR	[23]	8.9
9	DIS $F_3(\nu p)$	87	CCFR	[23]	5.5
10	D-Y <i>pp</i>	119	E605	[24]	6.4
11	D-Y pd/pp	1	NA51	[25]	0.5
12	D-Y pd/pp	11	E866	[26]	0.6
13	W _{lept. asym.}	11	CDF	[27]	15.1
14	$\overline{p}p \rightarrow \text{jet}X$	24	DØ	[28]	3.4
15	$\overline{p}p \rightarrow \text{jet}X$	33	CDF	[29]	3.7

APPENDIX A: ESTIMATES OF THE TOLERANCE PARAMETER FOR $\Delta \chi^2_{\text{global}}$

This appendix provides details of the various approaches mentioned in Secs. II D and V to estimate the tolerance parameter *T* defined by Eq. (5). In our global analysis based on $\Delta \chi^2_{\text{global}}$, all uncertainties of predictions of the PDF's according to the master formula Eq. (24) are directly proportional to the value of *T*.

The first two estimates rest on considerations of selfconsistency, which are required by our basic assumption that the 15 data sets used in the global analysis (see Table I) are *acceptable* and mutually *compatible*—in spite of the departure from ideal statistical expectations exhibited within many of the individual data sets, as well as apparent incompatibility between experiments when the errors are interpreted according to strict statistical rules [12].

Tolerance required by acceptability of the experiments. One can examine how well the best fit S_0 agrees with the individual data sets, by comparing χ_n^2 in Eq. (3) with the range $N_n \pm \sqrt{2N_n}$ that would be the expected 1σ range if the errors were ideal. The largest deviations found lie well outside that range: $\chi_n^2 - N_n(\sqrt{2N_n}) = 65.5(17.7), -64.8(18.5),$ 65.1(19.3), -25.9(15.4), and 22.4(8.1), for experiments n =2,3,4,10, and 15, respectively. By attributing the "abnormal" χ_n^2 's to unknown systematic errors or unusual fluctuations (or both), and accepting them in the definition of $\chi^2_{
m global}$ for the global analysis, we must anticipate a tolerance for the latter which is larger than that for an "ideal" χ^2 function. (Compare Appendix A of the preceding paper [18] for a quantitative discussion of the increase in T due to neglected systematic errors.) Since the sources of the deviation of these real experimental errors from ideal expectations are not known, it is not possible to give specific values for the overall tolerance. However, the sizes of the above quoted deviations (which, in each case, imply a very improbable fit to any theory model, according to ideal statistics) suggest that the required tolerance value for the overall $\Delta \chi^2_{global}$ (involving 1300 data points) must be rather large.

Tolerance required by mutual compatibility of the experiments. We can quantify the degree of compatibility among the 15 data sets by removing each one of them in turn from the analysis, and observing how much the total χ^2 for the remaining 14 sets can be lowered by readjusting the $\{a_i\}$. This is equivalent to minimizing χ^2 for each possible 14experiment subset of the data, and then asking how much increase in the χ^2 for those 14 experiments is necessary to accommodate the return of the removed set. These increases are listed as Δ_n in Table I. They range up to ≈ 20 . In other words, we have implicitly assumed that when a new experiment requires an increase of 20 in the χ^2_{global} of a plausible global data set, that new experiment is nevertheless sufficiently consistent with the global set that it can be included as an equal partner.⁷ Hence the value of T^2 must be substantially larger than 20.

A complementary, and more quantitative, estimate of the overall tolerance parameter *T* for our χ^2_{global} follows from the analyses of the preceding paper [18].

Tolerance calculated from confidence levels of individual experiments. In [18], we examine how the quality of fit to each of the 15 individual experiments varies as a function of the predicted value for various specific observable quantities such as σ_W or σ_Z . The fit parameters $\{a_i\}$ are continuously adjusted by the Lagrange multiplier method to yield the minimum possible value of χ^2_{global} for given values of the chosen observable. The constrainted fits obtained this way, and interpreted as "alternative hypotheses" in statistical analysis, are then compared to each of the 15 data sets to obtain a 90% confidence level error range for the individual experiments. Finally, these errors are combined with a definite algorithm to provide a quantifiable uncertainty measure for the cross section. In the case of the W production cross section at the Tevatron, $\sigma_W^{\rm TeV}$, this procedure yields an uncertainty of $\pm 4\%$, which translates into a value of ≈ 180 for $\Delta \chi^2_{\text{global}}$, or $T \approx 13$. This method is definite, but it is, in principle, process dependent. However, when the same analysis is applied to σ_Z^{TeV} , σ_W^{LHC} , and σ_Z^{LHC} (which probe different directions in the PDF parameter space) we find $\Delta \chi^2_{\text{global}}$ to be consistently in the same range as for σ_W^{TeV} , even though the percentage errors on the crossection vary from 4% at the Tevatron to 10% at LHC.

Based on all the above results, we adopt as our working hypothesis,

$$T \approx 10$$
 to 15,

which is quoted in Sec. II D [Eq. (6)], and used in Secs. IV and V for estimating the numerical results shown in the plots.

⁷Since 5 or 6 of the experiments require Δ_n in the range of 10–20, this level of inconsistency is not caused by problems with just one particular experiment—which would simply invite the removal of that experiment from the analysis.

TABLE II. Overall χ^2_{global} values and their increments above the best-fit value, for some current and historical parton distribution sets.

Current sets	χ^2	$\Delta\chi^2$	Historical sets	χ^2	$\Delta\chi^2$
CTEQ5M1	1188	-	CTEQ4M	1540	352
CTEQ5HJ	1272	84	MRSR2	1680	492
MRST99	1297	109	MRSR1	1758	570
MRST- $\alpha \downarrow$	1356	168	CTEQ3M	2254	1066
MRST- α^{\uparrow}	1531	343	MRSA'	3371	2183

Finally, it is of some interest to compare this tolerance estimate with the traditional—although by now generally recognized as questionable—gauge provided by differences between published PDF's.

Comparison of tolerance figures to differences between published PDF's. Table II lists the value obtained when our χ^2_{global} is computed using various current and historical PDF sets. The $\Delta \chi^2$ column lists the increase over the CTEQ5M1 set. Typical values for the modern sets are similar to the range 100–225 that corresponds to T=10-15. For previous generations of PDF sets, χ^2_{global} is much larger—not surprisingly, because the obsolete sets were extracted from much less accurate data, and without some of the physical processes such as W decay lepton asymmetry and inclusive jet production.

APPENDIX B: TESTS OF THE QUADRATIC APPROXIMATION

The Hessian method relies on a quadratic approximation (7) to the effective χ^2 function in the relevant neighborhood of its minimum. To test this approximation, Fig. 9 shows the dependence of χ^2 along a representative sample of the eigenvector directions. The steep directions 1 and 4 are indistinguishable from the ideal quadratic curve $\Delta \chi^2 = z^2$. The shallower directions 7, 10, 13, and 16, are represented fairly well by that parabola, although they exhibit noticeable cubic and higher-order effects. The agreement at small z is not perfect because we adjust the scale factors s_k in Eq. (10) (see footnote 5) to improve the average agreement over the important region $z \leq 5$, rather than defining the matrix H_{ii} in Eq. (7) strictly by the second derivatives at z=0. For this reason, the scale factors s_k in Eq. (17) are somewhat different from the $\sqrt{2}/\epsilon_k$ suggested by the Taylor series: the flattest directions are extremely flat only over very small intervals in z, so it would be misleading to represent them solely by their curvature at z=0.

Figure 10 shows the dependence of χ^2 along some random directions in $\{z_i\}$ space. The behavior is reasonably close to the ideal quadratic curve $\Delta \chi^2 = z^2$, implying that the quadratic approximation (7) is adequate. In particular, the approximation gives the range of z permitted by T=10 to an accuracy of $\approx 30\%$. Since the tolerance parameter T used to make the uncertainty estimates is known only to perhaps 50%, this level of accuracy is sufficient.



FIG. 9. Variation of χ^2 with distance along representative eigenvector directions 1, 4, 7, 10, 13, and 16. The first two, shown by solid curves, are nearly indistinguishable from each other and from the ideal $\Delta \chi^2 = z^2$. The remaining four, shown by dashed curves with increasing dash length denoting increasing eigenvector number, demonstrate that the quadratic approximation is adequate, though imperfect.

APPENDIX C: TABLE OF BEST FIT S₀

Table III lists the parameter values that define the "best fit" PDF set S_0 , which minimizes χ^2_{global} . It also lists the uncertainties (for T=5) in those parameters.

For each of the d=16 parameters, Table III also lists the components of a unit vector z_1, \ldots, z_d in the eigenvector



FIG. 10. Variation of χ^2 with distance along 10 randomly chosen directions in $\{z_i\}$ space. The dependence is represented acceptably well by the quadratic approximation $\Delta \chi^2 = z^2$, which is shown as the dotted curve.

TABLE III. Parameters of the global fit. Errors shown are for T=5, i.e., $\Delta \chi^2_{\text{global}}=25$. Fixed parameters: $A_4^{\vec{d}+\vec{u}}=1.0$, $B_2^{\vec{d}/\vec{u}}=15.0$, $B_3^{\vec{d}/\vec{u}}=10.0$, and $A_3^g=0.0 \Rightarrow A_4^g$ irrelevant. The z_k are proportional to the $z_k(a_i^{\pm})=\pm tM_{ik}$ of Eq. (29) and are normalized to $\Sigma_k z_k^2=1$.

Parameter	Value	Error	z_1	<i>z</i> ₂	<i>z</i> ₃	z_4	Z 5	z ₆	Z7	z_8
			Z9	z_{10}	z_{11}	z_{12}	z ₁₃	z_{14}	z_{15}	z_{16}
$A_{1}^{u_{v}} + 1$	0.466	0.094	0.04	-0.01	0.00	-0.08	-0.04	-0.09	0.05	0.07
1		+0.083 - 0.079	-0.06	-0.06	0.35	-0.74	-0.37	-0.39	-0.06	0.00
$A_2^{u_v}$	3.360	0.122	-0.01	-0.01	0.03	-0.14	-0.07	-0.37	-0.32	0.46
2		+.112 - 0.106	0.54	-0.13	0.06	-0.23	0.17	0.09	0.35	0.04
$\ln(1 + A_{2}^{u_{v}})$	2.553	0.51	0.00	0.00	0.00	0.00	0.06	-0.01	-0.06	-0.07
5		+0.44 - 0.45	0.10	0.02	-0.36	0.73	0.36	0.41	0.13	0.03
$A_{A}^{u_{v}}$	0.855	0.118	0.01	0.00	-0.04	0.19	-0.09	-0.13	-0.29	0.51
7		+0.110-0.118	0.48	-0.05	0.05	0.06	0.47	0.19	0.30	-0.01
$A_2^{d_v}$	4.230	0.45	0.00	0.00	0.00	0.00	0.06	-0.06	0.00	0.02
2		+0.45-0.40	-0.05	0.13	0.72	0.45	0.32	-0.23	-0.30	0.08
$\ln(1 + A_3^{d_v})$	2.388	0.64	0.00	0.00	0.00	0.00	-0.04	0.00	0.04	0.05
5		+0.59-0.62	-0.08	0.05	-0.04	0.88	0.38	0.26	-0.03	0.06
$A_4^{d_v}$	0.763	0.30	0.00	0.00	0.00	-0.01	-0.06	0.16	-0.13	-0.18
		+0.24-0.23	0.26	0.11	0.52	0.37	0.49	-0.11	-0.41	0.08
$\ln \tilde{f}_{g}$	-1.047	0.018	0.01	0.79	0.32	0.12	0.19	-0.25	-0.16	0.09
		+0.018 - 0.019	-0.19	0.26	0.00	-0.05	-0.03	0.00	-0.06	-0.11
$A_{1}^{g} + 1$	0.469	0.40	0.00	0.00	0.02	0.01	0.02	0.02	0.23	0.05
		+0.39-0.35	0.22	0.29	0.00	0.03	-0.09	0.04	0.12	0.89
A_2^g	5.574	2.65	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
		+2.98-2.30	0.00	-0.03	0.01	0.01	-0.03	0.03	0.11	0.99
$A_1^{d+u} + 1$	-0.009	0.148	0.00	-0.03	0.10	0.07	0.01	0.01	-0.24	-0.07
		+0.142 - 0.119	-0.15	-0.65	0.06	0.15	-0.13	-0.07	-0.09	-0.65
$A_2^{\overline{d}+\overline{u}}$	7.866	0.96	0.00	0.00	0.00	0.00	0.00	0.00	0.04	0.00
		+1.05 - 1.09	0.01	-0.21	0.03	-0.15	0.89	0.28	-0.10	-0.22
$\ln(1 + A_3^{\overline{d} + \overline{u}})$	2.031	0.89	0.00	0.00	0.00	0.00	0.00	0.00	-0.03	0.00
		+0.78-0.89	-0.04	0.29	-0.05	-0.28	0.66	0.20	0.04	0.59
$\ln B_0^{\overline{d}/\overline{u}}$	10.29	1.45	0.00	0.00	0.00	0.00	-0.01	-0.02	0.01	-0.04
0		+1.551 - 1.496	0.00	0.01	0.03	0.00	0.00	0.06	0.94	-0.32
$B_1^{\overline{d}/\overline{u}}$	5.379	0.85	0.00	0.00	0.00	-0.01	0.02	0.06	-0.02	0.10
1		+1.02-0.75	-0.02	-0.03	0.01	0.04	0.16	-0.21	0.93	-0.24
$B_{A}^{\overline{d}/\overline{u}}$	4.498	1.21	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.01
7		+1.20 - 1.12	0.00	0.01	0.06	-0.04	-0.27	0.94	-0.01	-0.20

basis. That unit vector gives the direction for which the parameter varies most rapidly with χ^2_{global} , i.e., the direction along which the parameter reaches its extreme values for a given increase in χ^2_{global} . For parameter a_i , the components z_k are proportional to M_{ik} according to Eq. (29).

APPENDIX D: TABLE AND GRAPHS OF THE EIGENVECTOR SETS S_{\perp}^{\pm}

Table IV and its continuation Table V completely specify the PDF eigenvector basis sets S_l^+ and S_l^- by listing all of their parameters at Q_0 . The notation and the best fit set S_0 are specified at the beginning of the table.

The coefficients listed provide all of the information that is needed for applications. For completeness, however, we state here explicitly the connections between these coefficients and the constructs that were used elsewhere in the paper to derive them. The fit parameters $\{a_i\}$ are related to the tabulated parameters by

$$a_{1} = A_{1}^{u_{v}} + 1, \quad a_{2} = A_{2}^{u_{v}}, \quad a_{3} = \ln(1 + A_{3}^{u_{v}}), \quad a_{4} = A_{4}^{u_{v}},$$

$$a_{5} = A_{2}^{d_{v}}, \quad a_{6} = \ln(1 + A_{3}^{d_{v}}), \quad a_{7} = A_{4}^{d_{v}}, \quad a_{8} = \ln \tilde{f}_{g},$$

$$a_{9} = A_{1}^{g} + 1, \quad a_{10} = A_{2}^{g}, \quad a_{11} = A_{1}^{\bar{d} + \bar{u}} + 1, \quad a_{12} = A_{2}^{\bar{d} + \bar{u}},$$

$$a_{13} = \ln(1 + A_{3}^{\bar{d} + \bar{u}}), \quad a_{14} = \ln B_{0}^{\bar{d}/\bar{u}}, \quad a_{15} = B_{1}^{\bar{d}/\bar{u}},$$

$$a_{16} = B_{4}^{\bar{d}/\bar{u}}.$$
(D1)

TABLE IV. Parameters of the best fit S_0 and their definitions, followed by eigenvector sets $S_1^+, S_1^-, S_2^+, S_2^-, \ldots$

-0.1941	-0.5337	3.3604	11.8404	0.8552	$\ln A_0^{u_v}$	$A_1^{u_v}$	$A_2^{u_v}$	$A_2^{u_v}$	$A_{A}^{u_{v}}$
-0.7175	-0.5337	4.2296	9.8950	0.7628	$\ln A_0^{d_v}$	$A_{1}^{d_{v}}$	$A_{2}^{\tilde{d}_{v}}$	$A_{2}^{d_{v}}$	$A_{A}^{\overset{4}{d}_{v}}$
1.8914	-0.5305	5.5737	0.0000	1.0000	$\ln A_0^g$	A_1^g	A_2^g	A_{3}^{g}	A_{4}^{g}
-1.1174	-1.0092	7.8658	6.6187	1.0000	$\ln A_{a}^{\overline{d}+\overline{u}}$	$A^{\overline{d}+\overline{u}}$	$A_{2}^{\overline{d}+\overline{u}}$	$A^{\overline{d}+\overline{u}}$	$A^{\overline{d}+\overline{u}}$
10.2940	5.3793	15.0000	10.0000	4.4980	$\ln R_0^{\overline{d}/\overline{u}}$	$B^{\overline{d}/\overline{u}}$	$\mathbf{R}^{\overline{d}/\overline{u}}$	$\mathbf{R}^{\overline{d}/\overline{u}}$	$R^{\overline{d}/\overline{u}}$
-0.1841	-0.5298	3 3596	11 8478	0.8558	-0.2042	-0.5376	33612	11 8331	0.8546
-0.7064	-0.5298	4 2295	9 8957	0.7629	-0.7285	-0.5376	4 2297	9 8942	0.0540
1 8919	-0.5304	5 5737	0.0000	1,0000	1 8910	-0.5376	5 5737	0.0000	1.0000
-1.1342	-1.0089	7 8658	6 6191	1,0000	-1.1010	-1.0095	7 8658	6.6182	1.0000
10 2940	5 3793	15 0000	10,0000	4 4980	10 2940	5 3793	15,0000	10,0000	4 4980
-0.1951	-0.5343	3 3597	11.8372	0.8557	-0.1931	-0.5331	3 3611	11 8437	0.8547
-0.7184	-0.5343	4 2291	9 8952	0.7636	-0.7165	-0.5331	4 2301	9 8947	0.7620
1 9025	-0.5321	5 5740	0.0000	1,0000	1 8802	-0.5288	5 5734	0.0000	1.0000
-1.1609	-1.0140	7 8662	6.6000	1,0000	-1.0751	-1.0043	7 8654	6.6256	1.0000
10 2940	5 3794	15,0000	10,0000	4 4980	10 2940	5 3792	15,0000	10,0000	4 4980
-0.1981	-0.5338	3 3645	11 8237	0.8501	-0.1903	-0.5336	3 3564	11.8567	0.8602
-0.7182	-0.5338	4 2287	9 9098	0.7633	-0.7167	-0.5336	4 2304	9 8806	0.7623
1 9095	-0.5244	5 5730	0.0000	1,0000	1 8738	-0.5364	5 5745	0.0000	1 0000
-1.0925	-0.9946	7 8651	6.6340	1.0000	-1.1422	-1.0234	7 8665	6.6037	1.0000
10 2939	5 3795	15 0000	10,0000	4 4980	10 2941	5 3791	15,0000	10,0000	4 4980
-0.1932	-0.5413	3 3436	11 8275	0.8774	-0.1960	-0.5260	3 3774	11.8536	0.8327
-0.7420	-0.5413	4 2312	9 8814	0.7592	-0.6928	-0.5260	4 2279	9 9087	0.0527
1 9005	-0.5271	5 5735	0.0000	1,0000	1 8822	-0.5200	5 5739	0.0000	1 0000
-1.0798	-0.9990	7 8655	6.6250	1.0000	-1.1544	-1.0195	7 8661	6.6122	1.0000
10 2964	5 3748	15,0000	10,0000	4 4970	10 2916	5 3839	15,0000	10,0000	4 4990
-0.2373	-0.5372	3 3513	12 2488	0.8440	-0.1528	-0.5303	3 3692	11 4555	0.8661
-0.7278	-0.5372	4 2550	9 5924	0.7435	-0.7078	-0.5303	4 2049	10 1974	0.0001
1 9071	-0.5244	5 5733	0.0000	1,0000	1 8761	-0.5364	5 5742	0.0000	1 0000
-1.1096	-1.0071	7 8668	6 6099	1.0000	-1.1246	-1.0112	7 8648	6.6272	1.0000
10 2848	5 3943	15,0000	10,0000	4 5012	10 3029	5 3647	15,0000	10,0000	1.0000
-0.2370	-0.5419	3 3147	11 7529	0.8399	-0.1529	-0.5258	3 4046	11 9257	0.8700
-0.6899	-0.5419	4 2039	9.8658	0.8113	-0.7483	-0.5258	4 2545	9 9232	0.7159
1 0021	-0.5228	4.2037 5.5734	0.0000	1 0000	1 8811	-0.5379	5 5741	0.0000	1 0000
-1.0010	-1.0074	7 8633	6 6099	1.0000	-1 1398	-1.0109	7 8682	6.6271	1.0000
10.2670	5 4325	15 0000	10,0000	4 5101	10 3201	5 3279	15,0000	10,0000	4 4863
-0.2078	-0.5282	3 3181	11 4087	0.8179	-0.1833	-0.5383	3 3960	12 2156	0.8866
-0.7700	-0.5282	1 2279	10 1816	0.7198	-0.6769	-0.5383	4 2310	9 6594	0.0000
2 0853	-0.4301	4.2277	0.0000	1,0000	1 7217	-0.6150	5 5770	0.0000	1.0000
-1.2033	-1.0484	7 9025	6 3721	1.0000	-1.0446	-0.9762	7 8349	6.8326	1.0000
10.3046	5 3626	15,0000	10,0000	1.0000	10 2850	5 3934	15 0000	10,0000	1.0000
-0.0901	-0.5260	3 /17/	11 2077	4.4717 0.0160	-0.2002	-0.5402	3 3060	10.0000	4.5051
-0.7805	-0.5209	J.4174 1 2276	10.22/1	0.9100	-0.6640	-0.5402	J. 2020	0 5 2 0 7	0.1911
1 0250	-0.5209	4.2370	0.0000	1 0000	1 8485	-0.5514	4.2220	7.3077 0.0000	1 0000
1.7337 - 1 1401	- 1 0105	7 9611	6 6229	1,0000	- 1 0929	-0.0004	J.J/00 7 8702	6.6147	1.0000
10.2407	5 4620	15 0000	10,0220	1.0000	1.0000	5 2005	15 0000	10,000	1.0000
10.2407	5.4029	10.0000	10.0000	4.0104	10.3449	5.2995	15.0000	10.0000	4.4014

Each of the a_i is thus related to a single PDF parameter, except for a_8 , which is related to \tilde{f}_g , the momentum fraction carried by gluons, and is thus determined by A_0^g, \ldots, A_4^g . The matrix elements of the transformation from the a_i coordinates to the eigenvector coordinates are given by

$$M_{il} = \frac{a_i(S_l^+) - a_i(S_l^-)}{2t}$$
 (D2) For

according to Eq. (18), where t=5 because that value was used to generate the S_l^{\pm} . Equations (9) and (14) imply

$$\sum_{i=1}^{d} M_{il} M_{ik} = s_l s_k \delta_{lk} \,. \tag{D3}$$

For $l \neq k$, this becomes $\sum_{i=1}^{d} M_{il} M_{ik} = 0$, which can serve as

TABLE V. Continuation of Table IV: parameters of the eigenvector sets $S_9^+, S_9^-, \ldots, S_{16}^+, S_{16}^-$.

-0.1659	-0.5391	3.4265	12.5012	0.9121	-0.2243	-0.5283	3.2943	11.2121	0.7983
-0.6220	-0.5391	4.2087	9.3472	0.8414	-0.8285	-0.5283	4.2505	10.4716	0.6842
2.0550	-0.4456	5.5690	0.0000	1.0000	1.7218	-0.6154	5.5785	0.0000	1.0000
-1.1277	-1.0312	7.8798	6.3693	1.0000	-1.0969	-0.9873	7.8518	6.8764	1.0000
10.2997	5.3659	15.0000	10.0000	4.4929	10.2883	5.3927	15.0000	10.0000	4.5031
-0.2209	-0.5385	3.3464	11.9679	0.8505	-0.1581	-0.5272	3.3793	11.6700	0.8616
-0.7087	-0.5385	4.2773	10.1973	0.7903	-0.7304	-0.5272	4.1650	9.4988	0.7256
2.0697	-0.4336	5.4996	0.0000	1.0000	1.6342	-0.6616	5.6741	0.0000	1.0000
-1.4530	-1.0911	7.6927	8.5071	1.0000	-0.6805	-0.8983	8.1000	4.6459	1.0000
10.3080	5.3568	15.0000	10.0000	4.5032	10.2750	5.4097	15.0000	10.0000	4.4910
0.0212	-0.4998	3.3685	9.5867	0.8619	-0.3963	-0.5645	3.3530	14.3021	0.8491
-0.3803	-0.4998	4.5680	9.5749	0.9267	-1.0637	-0.5645	3.9221	10.1942	0.6138
1.8986	-0.5295	5.5981	0.0000	1.0000	1.8849	-0.5314	5.5516	0.0000	1.0000
-1.0511	-0.9997	7.8935	6.2437	1.0000	-1.1519	-1.0178	7.8406	6.9762	1.0000
10.3431	5.3894	15.0000	10.0000	4.5718	10.2494	5.3702	15.0000	10.0000	4.4309
-0.6200	-0.6029	3.3322	17.6542	0.8617	0.2128	-0.4640	3.3889	7.8121	0.8487
- 1.0990	-0.6029	4.4299	18.0058	0.8742	-0.3418	-0.4640	4.0277	5.2176	0.6505
1.9215	-0.5171	5.5927	0.0000	1.0000	1.8610	-0.5440	5.5546	0.0000	1.0000
-0.9608	-0.9876	7.7202	4.9575	1.0000	-1.2706	-1.0310	8.0126	8.7623	1.0000
10.2885	5.4121	15.0000	10.0000	4.4446	10.2995	5.3462	15.0000	10.0000	4.5518
-0.3285	-0.5667	3.3800	14.2366	0.9075	-0.0441	-0.4965	3.3383	9.5883	0.7962
-0.7740	-0.5667	4.3664	12.6923	0.9007	-0.6624	-0.4965	4.0754	7.4209	0.6073
1.8124	-0.5621	5.5060	0.0000	1.0000	1.9800	-0.4949	5.6501	0.0000	1.0000
-1.3022	-1.0266	8.6732	12.2662	1.0000	-0.9707	-0.9896	6.9557	3.0774	1.0000
10.2946	5.5039	15.0000	10.0000	4.1962	10.2934	5.2389	15.0000	10.0000	4.8382
-0.4030	-0.5716	3.3718	14.9177	0.8788	-0.0012	-0.4980	3.3497	9.4869	0.8330
-0.9934	-0.5716	4.1229	11.9127	0.7280	-0.4696	-0.4980	4.3301	8.2828	0.7956
1.9461	-0.5128	5.6623	0.0000	1.0000	1.8398	-0.5471	5.4902	0.0000	1.0000
-1.1804	-1.0206	8.1430	8.1232	1.0000	-1.0571	-0.9985	7.6046	5.4285	1.0000
10.3763	5.1976	15.0000	10.0000	5.6698	10.2164	5.5506	15.0000	10.0000	3.3936
-0.2036	-0.5399	3.4062	12.7490	0.8931	-0.1860	-0.5282	3.3199	11.0884	0.8217
-0.9051	-0.5399	4.0867	9.6572	0.6340	-0.5709	-0.5282	4.3557	10.1093	0.8765
2.0530	-0.4819	5.8758	0.0000	1.0000	1.7480	-0.5734	5.3071	0.0000	1.0000
-1.1784	-1.0238	7.7670	6.9327	1.0000	-1.0517	-0.9963	7.9530	6.3518	1.0000
11.7435	6.2127	15.0000	10.0000	4.4913	9.0144	4.6435	15.0000	10.0000	4.5039
-0.2054	-0.5338	3.3661	12.0453	0.8534	-0.1853	-0.5336	3.3560	11.6819	0.8566
-0.7110	-0.5338	4.2688	10.4095	0.7879	-0.7226	-0.5336	4.1988	9.5075	0.7431
3.2798	-0.1380	8.5257	0.0000	1.0000	0.7313	-0.8385	3.2571	0.0000	1.0000
-1.7279	-1.1163	7.6289	12.7489	1.0000	-0.6889	-0.9252	8.0517	3.7937	1.0000
9.7697	5.1479	15.0000	10.0000	4.2287	10.7054	5.5609	15.0000	10.0000	4.7094

a check on numerical accuracy, while for l=k, it becomes $\sum_{i=1}^{d} M_{il}^2 = s_l^2$, which can be used to reconstruct s_1, \ldots, s_d .

Finally, for the benefit of the reader who is curious about them, graphs are shown in Fig. 11 of the differences described by each of the PDF eigenvector sets. One sees that the steeper directions (small values of l) mainly control aspects of the quark distribution, while the shallower directions (high values of *l*) control the gluon distribution, whose absolute uncertainty is larger. The variations in the gluon distribution show less variety than the quarks because the gluon distribution is described by only three parameters (including normalization) such that the most general variation for it is of the form $\Delta g/g = c_1 + c_2 \log x + c_3 \log(1-x)$.



FIG. 11. Displacements of the *u* quark and gluon distributions $f^+(x,Q)-f^-(x,Q)$ at Q=10GeV corresponding to $S_l^+ - S_l^-$. Length of dashes increases for l = 1, ..., 16.

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