Singular Value Decomposition, Hessian Errors, and Linear Algebra of Non-parametric Extraction of Partons from DIS

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By singular value decomposition (SVD) of a numerically singular Hessian matrix and a numerically singular system of linear equations for the experimental data (accumulated in the respective $\chi^2$ function) and constraints, least square solutions and their propagated errors for the non-parametric extraction of Partons from $F_2$ are obtained.

SVD and its physical application is phenomenologically described in the two cases. Among the subjects covered are: identification and properties of the boundary between the two subsets of ordered eigenvalues corresponding to range and null space, and the eigenvalue structure of the null space of the singular matrix, including a second boundary separating the smallest eigenvalues of essentially no information, in a particular case. The eigenvector-eigenvalue structure of ”redundancy and smallness” of the errors of two pdf sets, in our simplified Hessian model, is described by a secondary manifestation of deeper null space, in the context of SVD.

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INTRODUCTION

Singularity is widespread in nature and intriguing. Linearization is a major tool of human mathematical mind. In many areas of science and engineering, in particular, in physics, the enchanting theorem of linear algebra and the algorithm embodying it, the Singular Value Decomposition (SVD) of singular matrices is finding its applications. We encountered singularity in the way of extraction of parton distributions directly from the structure function $F_2$, in deep inelastic scattering (DIS), first in [1].

Based on one of the possibly typical examples, we are having an exposition of the physics of SVD, of how it diagnoses singularity in a linear system without ambiguity, and how it allows for its removal, bringing out desirable physical answer from its hiding place in the singular linear system. The example is a numerically singular Hessian matrix ($H$) error analysis in the context of a numerically singular system of linear equations for the least square (LS) estimates, of non-parametric extraction of parton distribution functions (pdfs), from experimental data, $F_2$, accumulated in a $\chi^2$, plus the equations of constraints, [2].

Notation and definitions

\[ e_k = h_k/\sqrt{\lambda_k}, \text{ defined under } h_k; \]
\[ f = n_F + 1 \leq 6: \text{ no. of pdf sets, ordered as in TABLE [1];} \]
\[ H(\text{or}A): \text{ numerically singular Hessian (or coefficient) matrix;} \]
\[ h_k, k = 1, ..., R: \text{ set of orthonormal eigenvector basis of the range of } H, \text{ corresponding to the eigenvalues } \lambda_k, k = 1, ..., R; e_k = h_k/\sqrt{\lambda_k}; \]
\[ \lambda_k, \text{ defined under } h_k; \]
\[ m = f \times n - 1 \leq 65: \text{ dimension of pdf-variables space; } \]
\[ n = 11: \text{ no. of } x-\text{bins of BCDS [3], TABLE [1];} \]
\[ \chi^2 \]

Generally, for a set of $n$ analyzed experimental $F_2^{exp}$ data points with a given covariance matrix $V_{exp}$,

\[ \chi^2 = \sum_{i,j=1}^{n} \Delta F_2^i (V_{exp}^{-1})^{ij} \Delta F_2^j, \]  
(1)
where $\Delta F^2_{ij} = F^2_{ij}^{exp} - F^2_{ij}^{th}$, and the theory (th) is LO PQCD here containing our pdfs directly as would be parameters, whose least square (LS) estimates minimizes $\chi^2$. In (1), obviously, constraint equations are left out of $\chi^2$.

There is a formal similarity between (1) and a Taylor expansion of $\chi^2$ about its LS minimum:

$$\Delta \chi^2 = \chi^2 - \chi^2_{min} \approx \sum_{i,j=1}^{m} \Delta u_i (V^{-1})_{ij} \Delta u_j. \quad (2)$$

In (2), $V_{ij}$ is the covariance matrix of the unknown pdf-variables $u_i, i = 1, ..., m$, here replacing the "pseudo-inverse" of the numerically singular Hessian matrix of second partial derivatives of $\chi^2$ (2): $\Delta u_i$ is the respective variation of $u_i$ about its least square (LS) estimate, $u_i^0$ - or the error.

We may always compute the Hessian $H$, (3), and use (2); however, we are not often given a full experimental covariance matrix. It is diagonal, having independent measurements, and (1) reduces to the following form that we use:

$$\chi^2 = \chi^2_p + \chi^2_d = \sum_{i,j=1}^{n_p} \left( \frac{\Delta F^2_{ij}^{p}}{\sigma^p_i} \right)^2 + \sum_{i,j=1}^{n_d} \left( \frac{\Delta F^2_{ij}^{d}}{\sigma^d_i} \right)^2. \quad (3)$$

In our present analysis, in (3), $n_{p,d} = 153,146$ are the total number of $p,d$ data left for analysis of BCDMS [3] after application of typical LO PQCD cuts for higher twists in the invariant mass squared, or squared boson-nucleon center-of-mass energy $W^2 = 20 GeV^2$ [4].

SINGULARITY AND SVD OF ZM VFN LO HESSIAN AND OTHER MATRICES FROM $\chi^2$ FOR $F^2_{p,d}$

Singular Value Decomposition of a matrix decomposes it into three: the middle one contains the eigenvalues, the others contain the eigenvector bases of the subspaces of interest of the singular matrix for separating and managing the singularity, namely null space - on the right, and range - on the left [5]. The magic of SVD is to pin-point numerically too small, ignorable, eigenvalues, corresponding to a ignorable set of eigenvectors of the numerical null space within domain. Remaining eigenvalues correspond to the remainder of domain which maps to the range, of the same dimension, in short, they correspond to range. Null space of $H$ is the subspace corresponding to the set of largest eigenvalues of its pseudoinverse, the covariance matrix $V$, which is side stepped or ultimately deleted, as the site of singularity. $\chi^2$, carrying the physical information of both our theory and the experimental data ($F^2_{exp}$), justifiably assumed to be decisive in forming the numerical range, is decisive in determining the boundary between the two subsets of eigenvalues corresponding to range and null space, here called "r-n border", for all the singular matrices obtained via its derivatives.

Operationally, i.e. in the process of trial motion up or down the scale of eigenvalues in the simulation program (determining $R$ in (8)), there is a single physical criterion for uncovering the r-n border of $H$: appearance of well patterned set of errors for the LS estimate of every pdf set. For $H$, or other singular matrices of the theory, r-n borderline acts as a "physical ordering lens", suddenly focusing the desirable physical properties of the solutions. We'll be using this concept as the first characterization or "qualitative, intuitive, physical definition" of the r-n border of a singular matrix. The direction of motion from r-n border towards null space has a more immediate and dramatic effect on the mal-formation of the error groups compared to the other way. However, even the direction of motion towards the range has clear observable indications favoring optimal uncovering of the r-n border.

Matrices of the theory are constructed from $\chi^2$, or $\chi^2_p, i = p,d, (4)$. For each, the most notable are two numerically singular matrices to begin with. Soon we will see that they may be made one and the same. One of them is the matrix of coefficients $A$ of the linear system of equations of 1st derivatives and the VFN constraints, to obtain the LS estimates of the pdf-variables, (5,2):

$$H_{ij} = \frac{1}{2} \frac{\partial^2 \chi^2}{\partial u_i \partial u_j}, i,j = 1, ..., m. \quad (5)$$

For BCDMS $\chi^2$, or $\chi^2_p$, (3), $H$, (5), is a square matrix on an $m = f \times n - 1 = 65$ dimensional pdf-variable space. For $\chi^2_d$, $q^3$ is absent, thus $f = 5$, and $m = 54$. Subtraction of the number 1 in $m$ is due to the fact that BCDMS data has no information on $q^3(x_{11}) = .07$, or on the quark $b(x_{11})$, as there is no data point with $Q^2 \geq m_b^2$ at $x_{11} = .07$. (This statement is true in our simple "zero mass" VFN scheme.) Thus, $q^3(x_{11})$ is dropped from the set of variables and the set of constraint equations in (3) and (5). Dropping the VFN constraint equation with the missing $q^3(x_{11})$ leaves $(2n-1)$ equations of VFN constraint, at $m_c^2$ and $m_b^2$, thus resulting in an $[m + (2n - 1)] \times m$ rectangular A matrix.

For $H$ (of $\chi^2$), the r-n border, as a "physical ordering lens", takes its place at $R = 41$, corresponding to eigenvalues of range $\lambda_k, k = 1, ..., R$ in the interval $[11.92, 1.77 \times 10^6]$, and a 24-dimensional "null space", half (12) corresponding to relatively larger eigenvalues in the interval $[10^{-7}, .08]$ and the other half (12) to eigenvalues numerically zero (within the accuracy of computation), figure [7]. There is considerable information in the 1st half of the null space, and very little in the second half, discussed in details later.
The characteristic break seen in figure 1 at the r-n border of all the Hessians is a main lesson of nature here, being translated to the act of a "physical ordering lens", we had noted prior to seeing the curves. We use it as a second, and "quantitative definition of the r-n border".

In figure 1, a similar behavior of $H_p$ and $H_d$ can be seen, except for $R_p = 29$ and $R_d = 30$, reflecting having less information, loosely speaking, than $H$ with $R = 41$.

Having a quadratic $\chi^2$ for $U$ everywhere in the $m = 65$ dimensional pdf-variable space, twice the matrix of second derivatives, $H$, coincides with the data part of the matrix of coefficients $A$ of the linear system of equations, constructed from the 1st derivatives of $\chi^2$.

Thus, twice the eigenvalues of $H$ in figure 1 are the same as the eigenvalues of the data part of $A$, figure 2. The extra 13 (or 14 for $A_p$) eigenvalue tail, of the range of $A$, is due to the 21 constraint equations in $A$, not in $H$.

A particular property of the r-n borderline is that its eigenvalues are of the order of $r = 1$ to 10 for all matrices of the model for all the subsets of the BCDMS data we encounter. It is interesting that the order of the eigenvalues in our range is similar to that of figure 2 of [6], $(r, 10^6)$, in spite of our highly different contexts.

The non-singular matrices of this data analysis arise from $F_3 = F_2^{p-n}$ for direct determination of the triplet pdf $q^3(x_i, Q^2), i = 1, ..., n$.

**SIMPLE LINEAR ALGEBRA AND ERRORS**

Our simple linear algebra everywhere in the $m$ dimensional pdf-variables space, a quadratic $\chi^2$ everywhere, exact expansion about its minimum [2], constant Hessian, has the advantage that renders our method simpler than the methods of parametric determination of pdfs [4, 6]. Our choice of direct pdf-variables instead of their parameters, permitted through the linear algebraic matrix method of evolution - method of solution of DGLAP [1, 2] - allows a simple linear decomposition of $F_2$ in terms of $m$ pdf-variables - instead of their highly non-linear parameter dependence. The foundation of having a simple linear algebra everywhere can be seen to be:

1. having $m$ direct pdf-variables, instead of intermediate parameters,
2. having our matrix method for their evolution,
3. having SVD for dealing with coefficient ($A$) and Hessian ($H$) matrices, and obtaining physically acceptable solutions.

However, there is a limitation in our simple linear system which can be compared with the methods of parametric determination of pdfs for dealing with a single data set, BCDMS. More specifically, in the error analysis of our $m$-dimensional singular Hessian, for the errors $\sigma_i, i = 1, ..., m$, it is the $R$-dimensional range which contains the main information. Thus there is some lack of information on the remaining $(m - R)$ errors! We can extract some of the information of the null space by a procedure in the lines that follow.

In line with [6] and [7] modified by SVD, Let $h_k, k = 1, ..., R$ be the set of orthonormal eigenvector basis of the range of $H$, corresponding to the eigenvalues $\lambda_k, k = 1, ..., R$, each having $m$ elements. $m = 65$ and $R = 41$ for $H$ of figure 1. Scale the set as: $e_k =$
\[ h_k / \sqrt{\lambda_k}, k = 1, \ldots, R. \] Expand the variation \( \sigma_i \) about the LS estimate, \( u_i^0 \), in terms of the scaled basis,

\[ \sigma_i = \Delta u_i = u_i - u_i^0 = \sum_{k=1}^{R} e_{ik} z_k, i = 1, \ldots, m, \quad (6) \]

where \( e_{ik} \) denotes the \( i \)th component of the vector \( e_k \). Thus (2) implies

\[
\Delta \chi^2 = \chi^2 - \chi^2_{\text{min}} = \chi^2(U) - \chi^2(U^0) = \sum_{i,j=1}^{m} \Delta u_i (V_n^{-1})_{ij} \Delta u_j = \sum_{k,l=1}^{R} z_k e_{ik} H_{ij} e_{jl} z_l \]

\[ = \sum_{k,l=1}^{R} z_k z_l (\lambda_l / \sqrt{\lambda_k}) \sum_{i=1}^{m} h_i^t H_{jl} h_l = \sum_{k=1}^{R} z_k^2. \quad (7) \]

\[ z_i \text{ are } R \text{ appropriately normalized linear combination of } \Delta u_i, \text{ defining orthogonal directions in the space of deviations of pdf-variables. By } \chi^2, \text{ a surface of constant } \chi^2 \text{ is a hyper-sphere of given radius in } z\text{-space. The error of the quantity } u_i \text{ in } z\text{-space is, equation (7) of } \chi^2, \]

\[ \sigma_i = \sqrt{\Delta \chi^2} = \sqrt{\sum_{k=1}^{R} (\partial u_i / \partial z_k)^2} = \sqrt{\sum_{k=1}^{R} e_{ik}^2}, i = 1, \ldots, m. \quad (8) \]

where \( \Delta \chi^2 = 1 \) was used, corresponding to one standard deviation errors, or 68 percent confidence level.

There are only \( R \) degrees of freedom in terms of \( z_i \). However, we sought \( m \) solutions of the errors in (8). Thus there is a complication for \( (m - R) = 24 \) of the errors. This appears as 24 small errors, 22 of which are in 2 redundant error sets, discussed in the next section.

Having singularity and using SVD, the major difference with references [6] and [7] is limitation by \( R \), for r-n border, in [6] through [5], dictated by the criteria of the previous section which boils down to the numerical observation that the errors become highly unacceptable, as soon as any eigenvector of the null space is added.

Equation (5) constructs errors which are grouped, in Well formed, i.e. dominantly increasing, sets for each of the \( f \) pdf sets. It may be understood as follows. For the \( l \)th element of the \( J \)th pdf set, in the prescribed order of TABLES I and II, the error \( \sigma_j \), is defined in [5], with \( i = ((j-1) \times n + l) \), before \( q^{24}(x_{11}) \) is dropped, and \( i = ((j-1) \times n - 1 + l) \) afterwards. Each error \( \sigma_i \), being the \( i \)th component of the scaled eigenvectors \( e_k \) as summed in [5], is observed to be dominated by one, or more \( e_k \), bunched together within the range, at a distance, correlated with \( (n - l) \), roughly \( \geq (n - l) \), from the r-n border. It is so that, for each \( j \), increase of \( l = 1, \ldots, n \) \( n \rightarrow (n - 1) \) for \( q^{24} \) on the average decreases the distance of the dominant \( e_k \) bunch from the r-n border, creating the dominantly increasing trend of the errors, within the \( j \)th set, because of scaling of \( e_k \). This structure is imparted to the errors via \( \chi^2 \) through the Hessian and eventually, through the mentioned structure of the index \( i \) in [8].

Note that if 1/2 of the coefficients of the \((2n - 1)\) linear equations of VFN constraints were added to \( H \), making it \([m + (2n - 1)] \times m = 86 \times 65 \) dimensional, \( H \) would have exactly 1/2 of the whole spectrum of \( A \) of figure 2. However, such a simple addition of coefficients of constraints to \( H \) is not allowed, as it is the matrix of second derivatives of \( \chi^2 \). From \( H \) meaningful (well-formed) errors are calculated at the answer \( U^0 = (u_1^0, \ldots, u_m^0) \) at the chosen fixed \( Q^2 \) (here 37.5 GeV\(^2\)), having the effect of continuity of \( U^0 \) at matching points due to the constraints, but no continuity of their variations, the errors, which becomes an assumption in the determination of errors that follows.

Addition of the coefficients of constraint equations to \( A \) follows naturally. It leads to spread of the range at the r-n border as may be seen in figure 2 by some extra eigenvalues, all of the order of 1. Physically meaningful solutions of \( A \) are obtained only through such addition, and then, as expected from figure 2, if the r-n border is placed at \( R = 54 \) and \( R_d = R_p = 43 \), so that the range includes the constraints. Deleting from \( A \) the less critical n constraints at \( Q^2 = m^2 \), as explicitly shown in [2], leads to non-physical answers.

For \( A \), (constraints for) continuity of pdfs are essential to get any physical results. It is not so for \( H \), which allows doing without constraints, and having a simplifying assumption. However, there are problems, some faced in the next section, very likely beginning in this simplification.

**REdundancy and smallness of errors of Two pdf sets**

Using the Hessian ([5]), there are problems of redundancy and having too-small error bars, associated with redundancy. Within our method, redundancy has a straight forward definition, referring to sameness of the quantity of error, i.e. [9].

When the two separate SVD for matrices \( A \) and \( H \) are done, we are left with physically desirable LS estimate of the pdfs, for \( d \) and \( p + d \) data. And there are Well formed sets of errors for each of the \( f = 5 \) or \( 6 \) groups of pdfs, for \( d \) and \( p + d \) data respectively, except for a "group redundancy" in the following sets:

\[ \Delta q^{15}(x_i, Q^2) = \Delta q^8(x_i, Q^2), i = 1, \ldots, n, \quad (9) \]

as seen in TABLE I. The \( 2m = 22 \) redundant error bars, [9], like the others, are well-formed, and acceptable except for being too small to be one standard deviation, for all of the different analysis, \( p, d, \) and \( p + d \).

Indeed, details of redundancy happen to be:

\[ e_{(i+p \times n)+k} = e_{(i+(p+1) \times n)+k}, i = 1, \ldots, n, k = 1, \ldots, R; \quad (10) \]
where \( p = 1 \) or \( 0 \), depending on whether proton data is included or not. Equations (10) and (8), not only imply (9), but also bring algebraic rigor to the term “information” and its shortage in terms of the properties of eigenvalues of \( H \).

Redundancy or duplication of errors, lack of information, being at the bottom of the null space are associated phenomena. As other aspects of the null space in our model, it may be eliminated

Exactly all of the eigenvectors of null space were forced out of counting in the sum of (8), to keep the formation of error sets well and acceptable. However, \( R \) eigenvectors of range, each have \( m \) components, associated with the \( m \) ordered eigenvalues, due to the symmetry of the \( m \times m \) matrix of eigenvectors. Thus, the relation of the last \( (m - R) \) components to the null space is to be taken into account now, leading to a completion of deletion of a bottom subspace of the null space.

A key observation, on the behavior of null space in the physical application of SVD to the Hessian, is that elimination of the two sets of redundant and smallest error bars completely eliminates the deepest part of the numerical null space, corresponding to 2\( n \) smallest eigenvalues. Elimination of a set of \( n \) redundant eigenvalues eliminates \( n \) smallest eigenvalues. Thus, along with redundancy, \( n \) dimensions are to be totally eliminated. However, the smallness of the remaining \( n \) error bars is essentially untouched, whose elimination completely drops the deepest \( n \) dimensions of the null space of \( H \).

Indeed, generally, a mapping of error sets into the ordered eigenvalue spectrum of the Hessian may be obtained by an ordered dropping, beginning with one set at a time. It takes the following order for \( F_{2,d} \) analysis: \( q^8, q^{24}, g, q^{8,15} \) (with an exception of 2 error points coming from above to below the \( g \) set, right below the \( r-n \) border). For \( F_{2,d} \) analysis, bulk of the extra \( q^8 \) set (10 out of \( n=11 \)) goes between \( q^8 \) and \( q^{24} \), in the above ordering.

The only by-product of dropping a set, is a slight extension or stretching of the eigenvalues towards the void the drop creates, which dies off with distance from the void! In figure 4, omission of the smallest 2\( n \) eigenvalues, after elimination of 2\( n \) redundant and small error bars of total \( H \) can be observed in \( H_{\text{filtered}} \). It results in a slight shrinkage of the eigenvalues (towards the created void), becoming pronounced for the smallest remaining eigenvalue(s), best observable for the last, or \((m - 2n)th = 43rd \), eigenvalue of \( H_{\text{filtered}} \).

Figure 5 shows a continuation of figure 4 for the eigenvalues of \( H \), obtained by raising the sensitivity of calculation. In it, the drop after \( r-n \) border is followed by a second drop before \( n \) approximately zero redundant eigenvalues, we may call “redundancy border”, beyond which there is essentially no information.

Beyond redundancy and its associated smallness of errors, belonging to the bottom of the null space, there are other relatively small, nevertheless, justifiable error bars observable in TABLE II and figure 4. The smallest such error bars are \( \Delta q_{p+d}^i(x_i, Q^2), i = 1, 2 \) and \( \Delta q_{p+d}^{24}(x_1, Q^2) \) (of which two are associated with the remaining highest two eigenvalues of null space, just below the \( r-n \) border), which, excluding inexact gluon, happen to be the largest relative errors, having the smallest LS central values; and the whole set \( \Delta q_{p+d}^i(x_i, Q^2), i = 1, ..., n \), which is acceptable on the grounds that \( SU(5) \) singlet is apparently the most exact parton distribution that our phenomenology (assumptions) gets out of this data, given the standard yardstick of LO MSTW.

### RESULTS AND DISCUSSION

One may remedy the relevant information limitation of the small error sets via a single enlargement (\(^toler-

### TABLE I. One standard deviation Hessian errors for proton plus deuteron \((p+d)\) data analysis, including redundancy and smallness of the errors of both \( p+d \) and \( d \) data \((\Delta q^8 = \Delta q^{15}\) columns).

| \( x_i \) | \( \Delta q_{p+d}^8 \) | \( \Delta q_{p+d}^{15} \) | \( \Delta q_{p+d}^{24} \) | \( \Delta q_{p+d}^g \) | \( \Delta q_{p+d}^d \) |
|-------|--------|--------|--------|--------|--------|
| 0.75  | 0.00280| 0.00334| 0.00234| 0.00245| 0.00290| 0.0844 |
| 0.65  | 0.00301| 0.00437| 0.00379| 0.0205 | 0.00511| 0.135  |
| 0.55  | 0.00344| 0.00764| 0.00742| 0.0449 | 0.0101 | 0.122  |
| 0.45  | 0.00388| 0.00785| 0.00590| 0.0341 | 0.00907| 0.112  |
| 0.35  | 0.00447| 0.00863| 0.00593| 0.0349 | 0.0105 | 0.105  |
| 0.275 | 0.00648| 0.0109 | 0.00728| 0.0473 | 0.0154 | 0.146  |
| 0.225 | 0.00712| 0.0115 | 0.00759| 0.0519 | 0.0176 | 0.188  |
| 0.18  | 0.00853| 0.0117 | 0.00816| 0.0561 | 0.0198 | 0.195  |
| 0.14  | 0.0106 | 0.0133 | 0.00929| 0.0628 | 0.0225 | 0.206  |
| 0.1   | 0.0134 | 0.0190 | 0.0130 | 0.0789 | 0.0238 | 0.173  |
| 0.07  | 0.0298 | 0.0135 | 0.0102 | 0.0433 | 0.330  |

### TABLE II. Least square estimated central values of the pdf solutions of equations (11) for proton plus deuteron \((p+d)\) BCDMS data analysis at \( Q_0^2 = 37.5 GeV^2 \), obtained at the \( x \)-points of the data.
FIG. 4. Points are ZM VFN LO parton momentum distributions at $Q^2 = 37.5 \text{ Gev}^2$, obtained at the $x$-points of the data. They have normal one standard deviation errors, except for $q^8$ and $q^{15}$, which arbitrarily have every other point missing (left and right show alternating $x$-points), to eliminate the redundancy of errors, and have the remaining error bars enlarged by a factor of 5, to remedy the smallness. Continuous curves are the corresponding GM global MSTW LO distributions with error margin. On the left, there are six sets (five quark flavors) obtained from 299 data points of BCDMS $F_2^{p,d}$ set. On the right, there are five sets obtained from 146 data points of BCDMS $F_2^d$ subset.
\section*{SUMMARY AND APPLICATIONS OF SVD}

We introduced a flexible linear algebraic direct method of determination of pdfo at the $x$-bins of the data, which has the advantage of simplicity. For our method, SVD is essential. The basics of SVD in terms of the properties of numerical null space and its border with range are abstracted and expounded. Our different cases of physical applications of SVD can be said to be essentially purging the null space in different ways, the most important being as the sight of singularity.

First, we begin with our criteria of determining the r-n border to get the physical answers, out of their hiding place due to singularity, for both matrices $A$ and $H$. For $A$, eigenvalue consideration is enough, and determination of $R$ brings us physical solution of the linear system of equations with coefficient matrix $A$. For $H$, eigenvectors are brought in. Those of the null space are purged out of calculations to arrive at physical answers. However, the $R$ eigenvectors of range, each have $m$ components, associated with the $m$ ordered eigenvalues, due to the symmetry of the $m \times m$ matrix of eigenvectors. Thus, the relation of the last $(m-R)$ components to the null space is analyzed. The last $n$ components are identified with the phenomenon of redundancy and with eigenvalues below redundancy border at the bottom of the null space, corresponding to the last 11 points of figure. Not showing redundant error bars in our resulting graphs is taken to mean a deleting of the last $n$ components.

SVD cut is not used to delete the remaining well formed, but too small error bar set, associated with redundancy. We try to keep whatever information which may be saved. There is an attempt to remedy the lack of information by a single degree of freedom corresponding to an enlargement factor $T$, equivalent to putting an arbitrary factor $T$ before a set of $n$ deleted eigenvectors of the null space.

Other experimental and phenomenological linear numerical modeling may run into singular matrices. How the spectrum can lend itself to interpretation depends on use value in the context. E.g. in such a linear modeling as signature of identification of molecules, a comparison of two singular spectrums with differences only in the bottom of the null space, of the type we have below the redundancy border in figure is a signal that two molecules can replace each other.

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