A Method for Merging the Results of Separate Least-Squares Fits and Testing for Systematic Errors

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A method is presented for merging the results of separate least-squares fits to obtain the most precise, single values for each of the molecular constants of a spectroscopic system. The output molecular constants and accompanying variance-covariance matrices from each of the separate fits are taken together as input to a correlated least-squares fit, which then yields the desired merged single values. The method is applied to a $\Sigma^2-\Sigma^2$ diatomic model band system using well-defined synthetic data as an example. Other currently used, multistate methods of obtaining molecular constants for a spectroscopic system are special cases of the method described here. For the nonlinear least-squares fits required for multiplet states, the merge method is particularly advantageous. The properties of the merged values of the molecular constants and their standard errors are examined with regard to forming confidence limits for these estimates and it is shown that, in most spectroscopic applications, the confidence limits have their usual meaning. Furthermore, it is similarly shown that the confidence limits for the variance of the merged fit provide a simple and convenient test for the presence of relative systematic errors between the separate least-squares results being merged. Although the method is illustrated here by merged separate band-by-band results, it also can be applied to any type of spectroscopic results.

INTRODUCTION

The band plays a central role in the analysis of molecular spectra. For example, the recognition of the band structure of a spectrum and the branch patterns of each band are the first steps in the assignment of vibrational and rotational quantum numbers. Furthermore, once the assignments are complete, the measured line positions of a band system are commonly grouped and tabulated by bands. Consequently, it is not surprising that, among the several approaches of reducing a band system to molecular constants, one popular way has been to make a least-squares fit of the calculated line

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1 Other approaches to reducing a system of bands to molecular constants involve two steps: first, the construction of intermediate “data,” such as combination differences or term values, that pertain to the vibrational levels of each electronic state separately, and second, the independent fitting of a rotational Hamiltonian to the intermediate “data” of each vibrational level to obtain the molecular constants of that level. These indirect, or one-state, approaches therefore differ distinctly from the direct, or multistate, approach employed in the band-by-band reduction discussed here. The former approach fits
positions derived from upper- and lower-state rotational Hamiltonians to the measured line positions of each band separately. Such a band-by-band reduction of a system of bands has the appeal of treating the data set conveniently in multiple "bite sizes," in contrast to a huge simultaneous multiband fit involving the Hamiltonians of all upper and lower vibrational levels and all of the measured line positions of all of the bands.

However, in almost all instances, many of the bands of a molecular band system will share common upper or lower vibrational levels. For such redundant systems, a band-by-band reduction will produce multiple values for each of the molecular constants of the common vibrational levels. Clearly, to halt the analysis of the band system at this point is undesirable, since the presence of multiple values for a constant indicates a missed opportunity, namely, the opportunity to treat the system as a whole and obtain a "best" single value for the molecular constant that, ideally, is more precise than that obtained from any one band separately.

When using the band-by-band reduction method, many approximate ways have been used to treat the system as a whole. One way would be to simply average all of the multiple values for each constant. However, this would ignore, among other things, the generally different precision of each value. Even the more preferable weighted average would ignore the often strong correlations (4, Section D-2-c) that generally exist between the errors of many pairs of molecular constants. Another approximate method of combining results of substantially different precision has been to simply hold certain constants in a least-squares fit fixed to separately obtained, high-precision values and then to determine values and errors for the remainder of the constants. Familiar examples of this are fixing to microwave results in fits to electronic bands, or fixing to combination-difference results for the zeroth vibrational level in a set of absorption bands. Clearly, this procedure assumes that the errors associated with the fixed values are negligible in comparison to the others. While this can be a good assumption in some cases, there are many cases where it is not. If the relative precision is not manyfold, then the assumption of no error in the fixed values usually means that the error estimates obtained for the other constants are too small. Rather than these approximate approaches, which require special cases to be valid, what is needed is a general technique.

We present here a method of "merging" the results of band-by-band fits to obtain the "best" single value for each of the molecular constants and show that this method (a) can yield the identical results as the huge simultaneous fit mentioned above, (b) is much more practical for nonlinear applications, and (c), as a bonus, provides a convenient quantitative indication of possible relative systematic errors between bands. Thus, the convenient "bite sizes" of the band-by-band fit and the enhanced precision available from treating a redundant band system as a whole are both achieved. Although the

with one Hamiltonian at a time; the latter approach fits with more than one Hamiltonian at a time. Both approaches have their merits and demerits. For example, the direct, multistate approach has the capability of using all of the measured lines in its reduction, whereas, the indirect, one-state approach cannot, in general. On the other hand, the direct, two-state approach cannot be applied if one of the rotational Hamiltonians in a band cannot be specified adequately (e.g., perturbations); whereas, the indirect, one-state approach can still be applied to obtain the constants of the state whose Hamiltonian can be specified. The details of these two different approaches have been examined recently by Allbritton et al. (1), Åslund (2), and Pliva and Telfair (3). Although the present paper applies the merge method to only the direct, or multistate, approach, it can be applied equally well in all of these approaches.
ability to test for relative systematic errors between bands arises somewhat as a dividend of this method, it may in time prove to be its most valuable feature, since such errors among the far-flung bands of an extensive band system are almost unavoidable.

We illustrate the merge method by applying it to a $\Pi - \Sigma$ diatomic band system. However, the method is not limited to this simple example, but applies equally well to electronic, infrared, Raman, and/or microwave data for transitions between or within molecular states of any multiplicity of diatomic and polyatomic molecules to determine the minimum-variance estimates of term values, combination differences, molecular constants, and/or Dunham-type coefficients, giving proper statistical regard to different measurement precision and correlation.

MERGED BAND-BY-BAND FITS

The present method has been applied to the bands of the $\text{HCl}^+ A^2\Sigma^+ - X^2\Pi$ system (5) and to the $\text{O}_2^+ b^3\Sigma_g^+ - a^3\Pi_u$ system in an accompanying paper (6), but here we apply it to a simple, but nevertheless realistic, band system that is designed to most clearly illustrate the features of the method.

Model Band System

Figure 1 schematically shows the three bands of a model $\Sigma - \Sigma$ system. Two electronic transitions are included: a $(0, 0)$ band having the first 50 $R$ lines and the first 50 $P$ lines and a $(0, 1)$ band having the first 25 $R$ lines and the first 25 $P$ lines. In addition, the model system contains one infrared transition: the $(1, 0)$ fundamental vibration-rotation band having the first 50 $R$ lines and the first 50 $P$ lines. It is clear that the model system has considerable redundancy. A band-by-band reduction will produce two values for each of the rotational constants $B$, and $D$, and will yield values for three band origins, one of which is related to the other two. Finally, each of the bands has at least one vibrational level in common with at least one other band, which is a necessary condition for treating a system as a whole. For example, a $(1, 2)$ band could not be combined with these bands.

![Figure 1. The structure of the model band system. $T'\Sigma'' = 0$.](image-url)
Table I. "True" Molecular Constants Used to Construct the Synthetic Lines\textsuperscript{a}

<table>
<thead>
<tr>
<th></th>
<th>Band 1 (0,0)</th>
<th>Band 2 (0,1)</th>
<th>Band 3 fundamental</th>
</tr>
</thead>
<tbody>
<tr>
<td>(f_0)</td>
<td>12000</td>
<td>10000</td>
<td>2000</td>
</tr>
<tr>
<td>(B_0)</td>
<td>1.5</td>
<td>1.5</td>
<td></td>
</tr>
<tr>
<td>(B''_0)</td>
<td>6.0 x 10^{-6}</td>
<td>6.0 x 10^{-6}</td>
<td></td>
</tr>
<tr>
<td>(B''_1)</td>
<td>0.98</td>
<td>0.98</td>
<td></td>
</tr>
<tr>
<td>(B''_2)</td>
<td>5.5 x 10^{-6}</td>
<td>5.5 x 10^{-6}</td>
<td></td>
</tr>
<tr>
<td>(B'')</td>
<td>1.0</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>(B''_0)</td>
<td>5.0 x 10^{-6}</td>
<td>5.0 x 10^{-6}</td>
<td></td>
</tr>
<tr>
<td>(\sigma)</td>
<td>0.05</td>
<td>0.05</td>
<td>0.005</td>
</tr>
</tbody>
</table>

\textsuperscript{a} All units are reciprocal centimeters.

Rather than use laboratory data for three bands of these types for some particular molecule from the literature, it is much more informative here to use "synthetic data," which offer far more flexibility in probing individual features of the method. (For examples of previous similar uses of such synthetic data, see Refs. (3, 4, 7).) In particular, a set of synthetic data for the three bands of Fig. 1 is created by (a) calculating "exact" line positions from the "true" molecular constants in Table I using the well-known energy level expressions for diatomic \(\Sigma\) states (8), and (b), to these "exact" line positions, adding random, normally distributed "errors" with specified standard deviations to form the "measured" line positions. Representative values for standard deviations were used here: \(\sigma = 0.05 \text{ cm}^{-1}\) for the two electronic bands and \(\sigma = 0.005 \text{ cm}^{-1}\) for the infrared band. It is to be emphasized that the values used for the molecular constants and standard deviations are completely arbitrary; the values used here are given only to be specific.

\textbf{Band-by-Band Least-Squares Fits}

A band-by-band reduction of the model system in Fig. 1 requires three separate least-squares fits. Corresponding to the commonly made assumptions (4, Section C-2) that the measurement errors of each band \(k\) \((k = 1, 2, \text{ and } 3)\) are randomly scattered with variance \(\sigma_k^2\) and zero covariance, then the fit to each band is the usual unweighted, uncorrelated least-squares solution (4, Section D) of the overdetermined matrix equation

\[ y_k = X_k \beta_k + \varepsilon_k, \quad k = 1, 2, \text{ and } 3, \]

where \(y_k, \beta_k, \text{ and } \varepsilon_k\) are the column vectors containing the \(n_k\) known measured line positions, the five desired molecular constants \(\nu_0, B', D', B'', \text{ and } D''\), and the \(n_k\) un-
Table II. A Set of Estimated Molecular Constants From a Band-by-Band Reduction\textsuperscript{a}

<table>
<thead>
<tr>
<th></th>
<th>Band 1</th>
<th>Band 2</th>
<th>Band 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>( v_0 )</td>
<td>12000.0007(10)</td>
<td>10000.0099(15)</td>
<td>20000.00027(92)</td>
</tr>
<tr>
<td>( b_0^{(0)} )</td>
<td>1.49959(22)</td>
<td>1.49934(56)</td>
<td></td>
</tr>
<tr>
<td>( a_0^{(0)} )</td>
<td>5.879(68) ( \times 10^{-6} )</td>
<td>5.30(67) ( \times 10^{-6} )</td>
<td></td>
</tr>
<tr>
<td>( b_1^{(0)} )</td>
<td>0.07939(56)</td>
<td></td>
<td>0.079985(21)</td>
</tr>
<tr>
<td>( d_1^{(0)} )</td>
<td>4.82(67) ( \times 10^{-6} )</td>
<td></td>
<td>5.4932(62) ( \times 10^{-6} )</td>
</tr>
<tr>
<td>( b_0^{(1)} )</td>
<td>0.99961(22)</td>
<td></td>
<td>0.999886(21)</td>
</tr>
<tr>
<td>( d_0^{(1)} )</td>
<td>4.887(68) ( \times 10^{-6} )</td>
<td></td>
<td>4.9933(62) ( \times 10^{-6} )</td>
</tr>
<tr>
<td>( a_k )</td>
<td>0.0529</td>
<td>0.0471</td>
<td>0.00482</td>
</tr>
</tbody>
</table>

Degrees of freedom: 95 45 95

\textsuperscript{a} The numbers in parentheses are the uncertainty in the last digits that corresponds to one standard error, i.e. \( 5.879(68) \times 10^{-6} = (5.879 \pm 0.068) \times 10^{-6} \).

Known measurement errors, respectively, and \( X_k \) is the known \( n_k \times 5 \) coefficient matrix (see Ref. (4, Section D-1), for an example), all for the \( k \)th band. The resulting least-squares values \( \hat{\beta}_k \) of the molecular constants are

\[
\hat{\beta}_k = (X_k^T X_k)^{-1} X_k^T y_k, \quad k = 1, 2, \text{ and } 3
\]

and the accompanying variance-covariance matrix is

\[
\hat{\Theta}_k = \sigma_k^2 (X_k^T X_k)^{-1}, \quad k = 1, 2, \text{ and } 3,
\]

where the estimated variance is

\[
\sigma_k^2 = (y_k - X_k \hat{\beta}_k)^T (y_k - X_k \hat{\beta}_k)/(n_k - 5), \quad k = 1, 2, \text{ and } 3.
\]

The band-by-band reduction yields three sets of estimated molecular constants \( \hat{\beta}_k \), \( k = 1, 2, \text{ and } 3 \), with five values per set, and three accompanying \( 5 \times 5 \) variance-covariance matrices \( \hat{\Theta}_k \), \( k = 1, 2, \text{ and } 3 \). A set of estimated molecular constants that arise from one particular set of random errors is listed in Table II. The different precision of the estimates from bands 1 and 2 caused by the different number of lines in each band is readily apparent, particularly for the molecular constant \( D_0' \). Moreover, the different precision of the estimates from bands 1 and 3 due to the different precision of the measurement errors is also readily apparent, particularly for the molecular constant \( D_0'' \). While the multiple values are very similar, as expected, they are not identical, as also expected, due to the random "measurement" errors. Considering the three-band system...

\textsuperscript{a} The notation follows that given in Ref. (4), Section D.
as a whole, these 15 values are not a desirable stopping point for the reduction. There are only eight basic molecular constants of this model system: two rotationless energies3 and six rotational constants, as shown on the right-hand side of Fig. 1. Thus, the problem at hand is to merge the 15 values from the three band-by-band fits into the “best” (i.e., minimum variance, linear, and unbiased, denoted MVLU (4, Section C-1)) values for these eight molecular constants, using a method that accounts for the different precision of the values and the nonzero correlation between them. The present method is a rigorous solution to that problem.

Merged Least-Squares Fit

The output of the band-by-band fits, $\hat{\mathbf{g}}_k$ and $\hat{\mathbf{\Theta}}_k$, $k = 1, 2, 3$, is taken to be the input of another least-squares fit that “merges” the 15 redundant values into eight nonredundant values, or specifically, a least squares solution to the overdetermined 15 equations:

$$y = X\mathbf{g} + \delta,$$  \hspace{1cm} (5)

the details of which are defined in Table III. The input “data” from the three band-

Table III. Details of the Equations of the "Merge" Model (Eq. 5).

<table>
<thead>
<tr>
<th>$\hat{\mathbf{g}}_{0}^{(1)}$</th>
<th>$\hat{\mathbf{g}}_{0}^{(2)}$</th>
<th>$\hat{\mathbf{g}}_{0}^{(3)}$</th>
<th>$\hat{\mathbf{g}}_{1}^{(1)}$</th>
<th>$\hat{\mathbf{g}}_{1}^{(2)}$</th>
<th>$\hat{\mathbf{g}}_{1}^{(3)}$</th>
<th>$\hat{\mathbf{g}}_{2}^{(1)}$</th>
<th>$\hat{\mathbf{g}}_{2}^{(2)}$</th>
<th>$\hat{\mathbf{g}}_{2}^{(3)}$</th>
<th>$\hat{\mathbf{g}}_{3}^{(1)}$</th>
<th>$\hat{\mathbf{g}}_{3}^{(2)}$</th>
<th>$\hat{\mathbf{g}}_{3}^{(3)}$</th>
<th>$\hat{\mathbf{g}}_{4}^{(1)}$</th>
<th>$\hat{\mathbf{g}}_{4}^{(2)}$</th>
<th>$\hat{\mathbf{g}}_{4}^{(3)}$</th>
<th>$\hat{\mathbf{g}}_{5}^{(1)}$</th>
<th>$\hat{\mathbf{g}}_{5}^{(2)}$</th>
<th>$\hat{\mathbf{g}}_{5}^{(3)}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 1 0 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 0 0 0 0 0 0 1 0 0 0 0 0 0 0</td>
<td>0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>1 0 0 -1 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 1 0 0 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 1 0 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 0 1 0 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 0 0 1 0 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 0 0 0 1 0 0 0 0 0 0 0 0 0 0</td>
<td>0 0 0 0 0 0 1 0 0 0 0 0 0 0 0 0</td>
<td>0 0 0 0 0 0 0 1 0 0 0 0 0 0 0 0</td>
<td>0 0 0 0 0 0 0 0 0 1 0 0 0 0 0 0</td>
<td>0 0 0 0 0 0 0 0 0 0 1 0 0 0 0 0</td>
<td>0 0 0 0 0 0 0 0 0 0 0 1 0 0 0 0</td>
<td>0 0 0 0 0 0 0 0 0 0 0 0 1 0 0 0</td>
<td>0 0 0 0 0 0 0 0 0 0 0 0 0 1 0 0</td>
</tr>
</tbody>
</table>

3 $T'_0'$ and $T''_0'$ are actually energy differences, $T'_0' - T''_0'$ and $T'_1' - T''_1'$, respectively (since this is all that can be determined physically); however, we have defined the rotationless energy $T''_0'$ of the lowest vibrational level to be zero. The choice of which level and what value to be the absolute energy reference is, of course, arbitrary and the present choice is based only on aesthetics.
by-band fits are stacked in the 15 elements of \( y \). The eight nonredundant unknown constants are represented by \( \mathbf{g} \). The coefficient matrix \( X \) relates the redundant values of \( y \) to the corresponding members of \( \mathbf{g} \). For example, the second and seventh individual equations in the matrix Eq. (5) are

\[
\begin{align*}
\hat{B}_0'(1) &= B_0' + \delta_{2y}, \\
\hat{B}_0'(2) &= B_0' + \delta_{7y}.
\end{align*}
\]

Equation (5) also makes similar statements about the other redundant values. Of course, because of random measurement errors, the redundant values are almost never equal. Therefore, the least-squares solution of Eq. (5) finds values for the molecular constants \( \mathbf{g} \) that minimize the sum of the squares of the unknown \( \delta \), subject to known interrelations among the \( \delta \). In fact, recognizing that there are indeed interrelations among the unknown errors of the input values, \( \delta_k, k = 1, 2, \) and 3, is at the heart of the present method.

These interrelations arise because the variance–covariance matrices \( \hat{\Theta}_k \) associated with the \( \delta_k, k = 1, 2, \) and 3, generally do not have equal diagonal elements and zero off-diagonal elements, i.e., \( \hat{\Theta}_k \neq \hat{\delta}I \), where \( \hat{\delta} \) is a constant and \( I \) is the identity matrix. To account for these unequal variances and nonzero covariances, the solution of Eq. (5) must employ the weighted, correlated least-squares formalism (4, Section F). Using this, the molecular constants that minimize \( \delta^T \delta \) subject to the interrelations contained in \( \hat{\Theta}_k, k = 1, 2, \) and 3, are given by

\[
\hat{\mathbf{g}}^M = (X^T \hat{\Phi}^{-1} X)^{-1} X^T \hat{\Phi}^{-1} y,
\]

where \( \hat{\Phi} \) is the non-diagonal 15 \( \times \) 15 matrix composed of the individual \( \hat{\Theta}_k \),

\[
\hat{\Phi} = \begin{bmatrix}
\hat{\Theta}_1 & 0 & 0 \\
0 & \hat{\Theta}_2 & 0 \\
0 & 0 & \hat{\Theta}_3
\end{bmatrix}.
\]

In addition to the circumflex \( \hat{\cdot} \) that denotes that the \( \hat{\mathbf{g}}^M \) values are MVLU estimates, we have added the superscript \( M \) to denote that they are estimates of the merged method, a distinction that will be useful below.

The precision of the estimates \( \hat{\mathbf{g}}^M \) is, of course, indicated by their standard errors, which are the square roots of the diagonal elements of the variance–covariance matrix associated with \( \hat{\mathbf{g}}^M \):

\[
\hat{\Sigma}^M = \hat{\vartheta}_M \hat{\Sigma}^M,
\]

where the merged dispersion matrix is given by

\[
\hat{\Sigma}^M = (X^T \hat{\Phi}^{-1} X)^{-1}.
\]

The estimated variance of the merged fit \( \vartheta_M^2 \) is given by

\[
\vartheta_M^2 = (y - X\hat{\mathbf{g}}^M)^T \hat{\Phi}^{-1} (y - X\hat{\mathbf{g}}^M)/f_M,
\]

where the degrees of freedom of the merged fit are denoted by \( f_M \). For the example considered here, the degrees of freedom are \( f_M = 15 - 8 = 7 \).
Equations (1)–(10) define the merged band-by-band method. It is a two-step approach. The first step (Eqs. (1)–(4)) is a set of band-by-band fits\(^4\) to determine $\hat{\beta}_k$ and $\hat{\delta}_k$ for each of the bands. The second step (Eqs. (5)–(10)) is a correlated fit to these output values to determine the “best” single values for the molecular constants of each vibrational level. It is clear that the merged estimates $\hat{\beta}^M$, $\hat{\delta}^M$, and $\hat{\sigma}_M^2$ depend not only on the redundant values $\hat{\beta}_k$, $k = 1, 2, 3$, from the band by band fits, but also on the accompanying variance–covariance matrices $\hat{\Theta}_k$, $k = 1, 2, 3$, composing $\hat{\Theta}$.

The pervasive appearance of $\Phi$ in Eqs. (6)–(10) properly accounts for two features of the input $\hat{\beta}_k$ making up $y$. First, the unequal diagonal elements of $\Phi$ account for the fact that the input values $y$ are generally of different precision. For example, the variances of the two values $\Bar{\beta}_o'(1)$ and $\Bar{\beta}_o'(2)$ from the separate fits to bands 1 and 2, respectively, differ by about a factor of six. The elements $\Phi_{22}$ and $\Phi_{77}$ will account for this by weighting the merged value for $\Bar{\beta}_o'$ toward $\Bar{\beta}_o'(1)$ by this factor.\(^5\) Secondly, the band-by-band fits are not limited to unweighted, uncorrelated least-squares formulations. If the lines of a band are of different estimatable precision, then a weighted fit can be made. The correlations among line position measurements are much more difficult to estimate reasonably and always have been assumed to be zero.

\(^4\) This does not mean that the merged value for $\Bar{\beta}_o'$ will be a factor-of-six weighted average of $\Bar{\beta}_o'(1)$ and $\Bar{\beta}_o'(2)$, since there are many other interrelations involving $\Bar{\beta}_o'$ for which the merged fit will also simultaneously account.
nonzero off-diagonal elements of $\Phi$ account for the fact that some of the input values are generally correlated (4, Section D-2-c). For example, the errors in $B_0'(1)$ of the upper state and $B_o''(1)$ of the lower state are correlated with a coefficient of 0.9941. This means that the errors in $B_0'(1)$ and $B_o''(1)$ are very likely to be of the same magnitude and sign. The nonzero off-diagonal element $\Phi_{24}$ will account for this by weighting the difference of merged values of $B_0'$ and $B_o''$ toward the difference of $B_0'(1)$ and $B_o''(1)$. Thus, $\Phi$ is a “generalized weight” matrix that simultaneously balances all of the diagonal and off-diagonal interrelations among the errors of $\gamma$. Fundamentally, since $\Phi$ is made up of the elements of $\delta X_k^T X_k^{-1}$, $k = 1, 2, \text{and} 3$ (cf. Eqs. (3) and (7)), it contains not only the information about the energy level expressions, but also the differing rotational development and measurement precision among the bands, all of which go into creating the interrelations.

Table IV gives the values that result from merging the band-by-band results in Table II. Even though these are results that correspond to a particular set of random “measurement” errors, Table IV does show some of the general features of merging the band-by-band fits. Compared to Table II, there have been changes in both the precision and the values of the molecular constants.

First, the precision (as indicated by the standard errors) of the molecular constants of the $\tau' = 0$ level of the upper electronic state has improved considerably. For example, the standard error associated with the merged value for $B_0'$ is a factor of 10 smaller than that associated with the value for $B_0'$ obtained from the individual fit to band 1, the most favorable of the two electronic bands (see Table II). This improvement is mainly due to the fact that the high precision of the infrared band, which is associated only with the lower electronic state, has been “propagated” into the upper electronic state by the high correlation of the upper- and lower-state rotational constants of the two electronic bands. It is interesting to note that this correlation has often been considered a handicap in obtaining precise molecular constants; here, it obviously is an advantage! Such dramatic changes in the precision do not occur for the molecular constants of the two levels of the lower electronic state because these levels are already dominated by the more precise infrared band.

Secondly, the merged values for the molecular constants in Table IV differ, of course, in varying degrees from the corresponding band-by-band results in Table II. Generally, the merged results lie close to the more precise values from the band-by-band fit, namely, those of the more rotationally developed electronic band and particularly the infrared band. However, not all of the differences are what one might guess from such simple considerations. Using the upper-state constant $D_0'$ as an example, the merged value, $\bar{D}_0' = 5.9815 \times 10^{-6}$ cm$^{-1}$, does not lie between the values obtained from the separate fits to the two electronic bands, $\bar{D}_0'(1) = 5.879 \times 10^{-6}$ and $\bar{D}_0'(2) = 5.30 \times 10^{-6}$ cm$^{-1}$. Neither a simple average nor a weighted average based on the individual variances could have predicted this, thereby indicating the role of the covariances, which place strong interrelations on the values that highly correlated rotational constants can assume simultaneously. Other properties of these merged estimates $\bar{\delta}^M$, $\bar{\delta}^M$, and $\delta M^2$ will be discussed below.
COMPARISON TO SIMULTANEOUS MULTIBAND LEAST-SQUARES FIT

The increase in precision gained by merging the results of the band-by-band fits is, of course, just the familiar statistical gain that occurs when a data set is enlarged. For a molecular band system, this enlargement has, in the past, usually taken the form of combining all of the bands into a single, simultaneous multiband fit to obtain the best single values for the molecular constants of each vibrational level. For example, Pliva and Telfair (3) have explored the systematics of this gain for a simultaneous multiband fit to an absorption band system with varying number of bands.

Weighted Simultaneous Multiband Least-Squares Fit

For the present example, a simultaneous multiband fit would be a single fit of the eight $\mathbf{g}$ molecular constants (Fig. 1, right-hand side) to the 250 line positions $y_1$, $y_2$, and $y_3$ of the three bands. Since electronic and infrared bands are generally expected to have different measurement precision, this simultaneous fit should be a weighted least-squares fit (4, Section E). In the absence of more specific information, the simplest weighting scheme would be to assume that the electronic measurements are of equal precision and that the infrared measurements are also of equal precision, but are more precise than the electronic. Thus, in the weighted fit, all of the electronic lines are assigned the same weight and all of the infrared lines could be assigned a greater weight, where the weights should be inversely proportional to some estimate of the generally unknown measurement precision. For these estimates, one could simply assume that, generally, infrared measurements are $R$ times more precise than electronic measurements and could use weights of 1 and $R^2$ for the electronic and infrared measurements, respectively. However, this somewhat overly general assumption can be replaced by quantitative estimates of the electronic and infrared measurement precision. These could come from fits to each band separately, examples of which are given as the $\delta_k$ values in Table II. Such a priori estimates are certainly generally preferable to a posteriori assumed ratios $R$. Thus, in the most general application, the weighted simultaneous multiband method has, as its first step, a set of band-by-band fits. This is the same as the first step of the merged band-by-band method, except that only the $\delta_k^2$ are of interest here, not $\mathbf{g}_k$ and $\mathbf{\phi}_k$.

The second step is an estimation of the eight molecular constants $\mathbf{g}$ by a weighted least-squares solution (4, Section E) of the 250 equations

$$\begin{align*}
y_1 &= Z_1 \mathbf{g} + \gamma_1, \\
y_2 &= Z_2 \mathbf{g} + \gamma_2, \\
y_3 &= Z_3 \mathbf{g} + \gamma_3,
\end{align*}$$

with the weight of the lines $y_k$ of the $k$th band being $1/\sigma_k^2$. The $250 \times 8$ matrix $Z$,

---

6 Because of this improvement in precision for the larger data set, it would appear wise to keep marginally statistically significant molecular constants in the band-by-band Hamiltonians and delay the final decision as to their significance until the increased precision of the enlarged data set permitted a more sensitive test.

7 In weighted least-squares fits, only the relative values of the weights influence the determination of the molecular constants and their standard errors (4, Section E-2).
where $Z^T = [Z_1^T Z_2^T Z_3^T]$, is unwieldy, but straightforward. (An example for a slightly different problem is given in Ref. (4), Eq. 19.) The weighted least-squares solution of Eq. (11) is

$$\hat{\beta}^W = \left[ \sum_{k=1}^{3} (Z_k^T \hat{\sigma}_k^2)^{-1} \right]^{-1} \left[ \sum_{k=1}^{3} (Z_k^T \hat{y}_k^b / \hat{\sigma}_k^2) \right].$$

(12)

Here, the superscript W is used to specifically identify estimates obtained from the weighted simultaneous multiband fit. The variance–covariance matrix that accompanies $\hat{\beta}^W$ is given by

$$\hat{\Theta}^W = \hat{\sigma}^W \hat{V}^W,$$

(13)

where the dispersion matrix is

$$\hat{V}^W = \left[ \sum_{k=1}^{3} (Z_k^T Z_k / \hat{\sigma}_k^2) \right]^{-1}.$$

(14)

The estimated variance of the weighted fit is

$$\hat{\sigma}^W = \left[ \sum_{k=1}^{3} (y_k^b - Z_k \hat{\beta}^W)^T (y_k^b - Z_k \hat{\beta}^W) / \hat{f}_W \right] / \hat{f}_W,$$

(15)

where $\hat{f}_W$ are the degrees of freedom. In the present example $\hat{f}_W = 250 - 8 = 242$.

The merged fit output $\hat{\beta}^M$, $\hat{\Theta}^M$, and $\hat{\sigma}^M$ and the simultaneous fit output $\hat{\beta}^W$, $\hat{\Theta}^W$, and $\hat{\sigma}^W$ are estimates of the same eight molecular constants $\beta$; therefore, it is interesting to compare the values obtained using the two methods. Table V shows the results of

<table>
<thead>
<tr>
<th>Table V. Weighted Simultaneous Molecular Constants That Correspond to the Band-by-Band Results in Table IIa</th>
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<tbody>
<tr>
<td>$\hat{\gamma}_0$</td>
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<tr>
<td>$\hat{b}_0$</td>
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<tr>
<td>$\hat{\beta}_0$</td>
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<tr>
<td>$\hat{r}_1$</td>
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<tr>
<td>$\hat{\gamma}_W$</td>
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<td>degrees of freedom</td>
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</table>

a See the footnote of Table II
applying the weighted simultaneous multiband fit to the same example used in Tables II and IV. The results are strikingly similar. In fact, starting with relations between the $X$ and $Z$ coefficient matrices, some tedious, but straightforward, matrix algebra shows that

$$\hat{\beta}^M = \hat{\beta}^W,$$  \hspace{1cm} (16)

$$\hat{V}^M = \hat{V}^W,$$  \hspace{1cm} (17)

and

$$\frac{(\delta^M - 1)}{(\delta^W - 1)} = f_w/f_M.$$  \hspace{1cm} (18)

These relations show that the *weighted simultaneous multiband fit and merged band-by-band fit are equivalent*. The results of one fit are either identical to, or can be derived from, the other. Since the first steps of these two methods are the same, the choice between the two depends on the features of the second step.

A comparison certainly appears to favor the merged fit. Once each of the bands have been fitted to yield sets of $\hat{\beta}_k$ and $\hat{\Theta}_k$, there is no need to handle the input line positions $y_k$ again, as would be required by the weighted simultaneous multiband fit; the merge fit can get the *same* values for $\hat{\beta}$ by handling only the compact sets of $\hat{\beta}_k$ and $\hat{\Theta}_k$. This is much more convenient and should often require less computer storage. However, the most important factor is probably the fact that the second step of the merge method is always a *linear* fit. When the application involves the nonlinear Hamiltonians of higher-order multiplet electronic states, each band-by-band fit does, to be sure, involve a nonlinear fit to yield the molecular constants of the two vibrational levels. However, the second step of the merge method is *always* a linear least-squares fit, as given by Eq. (6). This is in contrast to the second step of the simultaneous multiband method, in which a nonlinear fit must be made involving the molecular constants of *all* of the vibrational levels. This generally expensive nonlinear second step clearly can be replaced by the less-expensive linear second step of the merged fit, since the results are equivalent. Discussion of other aspects of this comparison, particularly the consequences of Eq. (18), will be deferred until a later section below.

**Unweighted Simultaneous Multiband Least-Squares Fit**

Many times, it has been implicitly assumed that all of the recorded bands involved have the same measurement precision. An example would be the bands of one electronic system that have been recorded in a relatively restricted wavelength region by the same investigator on one instrument over a relatively short time. In such applications, the simultaneous multiband fit is often shortened to one step, an *unweighted* least-squares fit to the lines of all bands to determine the values $\hat{\beta}^U$, $\hat{\Theta}^U$, and $\hat{\phi}^U$, where $U$ has been used to identify estimates obtained with this common unweighted variation of the simultaneous multiband fit. The equations yielding these estimates would be Eqs. (11)–(15), with $\delta_i^2 = 1$, $k = 1, 2, 3$.

It is important to realize that the relations similar to Eqs. (16)–(18) do not rigorously apply now. Even though they seek estimates of $\beta$ from the same data, the two methods cannot be exactly equivalent because the least-squares models are different. As the price it must pay for its simplicity, the unweighted simultaneous multiband model involves the *additional* assumption that all of the lines of all bands have the same
measurement precision. If this assumption is a good one, i.e., if all of the \( \sigma_k \) are nearly equal, then

\[
\hat{\beta}_W \sim \hat{\beta}_U \tag{19}
\]

and

\[
\hat{\theta}_W \sim \hat{\theta}_U. \tag{20}
\]

In addition to requiring this extra assumption, the unweighted simultaneous fit has a rather serious disadvantage of a practical nature. Since it is a one-step method, relatively large least-squares fits must be made over and over during the inevitable cycles of discovering mispunched data cards, misassigned lines, inexplicably anomalous measurements, and model testing (e.g., checking to see if \( H_s' \) and \( H_s'' \) are needed). This disadvantage is greatly magnified if the application is nonlinear, for reasons similar to those mentioned above.

**PROPERTIES OF THE MERGED ESTIMATES**

Since the values \( \hat{\beta} \) obtained from a least-squares fit are only estimates, it is necessary to indicate their quality. This is done by stating confidence limits (4, Section D-3-b), or more commonly, stating standard errors, from which the confidence limits can be constructed. Such limits define the range about the estimated value \( \hat{\beta} \), in which the unknown \( \beta_{true} \) is believed, with a certain level of confidence, to lie. The establishment of confidence limits depends on the statistical properties of the estimates \( \hat{\beta} \) and \( \sigma^2 \), which arise from, among other things, the statistical properties of the measurement errors. The latter are, of course, unknown; hence, recourse commonly is made to a set of "reasonable assumptions."

In addition to assuming a correct physical model, the common least-squares assumptions (4, Sections C-2 and D-3) are that the measurement errors are (i) normally distributed with (ii) zero mean and (iii) have a variance–covariance matrix \( \sigma^2 \Phi \) that is known, except for the common factor \( \sigma^2 \). Given these assumptions, well-defined confidence limits for the estimates \( \hat{\beta} \) from the familiar single-step least-squares method can be constructed straightforwardly by following prescribed procedures. The merged band-by-band method, however, is a two-step fit in which the output estimates from the first fit are used as input "data" to the second fit. As we shall see below, this feature can alter the statistical properties of the merged estimates \( \hat{\beta}^M, \hat{\theta}^M, \) and \( \sigma^2_M \) in such a way that the confidence limits constructed from these estimates in the usual manner do not always have their normally expected, well-defined meaning. The origin of this loss of meaning of the merged confidence limits occurs between the first and second steps of the merged method. The output estimates from the first step must be examined with regard to the above three assumptions since they are the input to the correlated least-squares fit of the second step.

The assumption of normality causes no problem. If the measurement errors of the line positions \( y_k \) are normally distributed, as is commonly assumed,\(^a\) then it can be shown (9, p. 99) that the least-squares estimates from the first step \( \hat{\beta}_k \) are also normally distributed. The assumption of normally distributed measurement errors appears to be very realistic in most spectroscopic applications. There are simple tests with which its validity can be checked (see (4), Section D-3-a for an example).
distinguished. The assumption of zero mean requires that the systematic errors in both
the fitted model and the measurements of each band be much smaller than the random
errors. While this assumption is almost always implicitly evoked, it is clear that it is
often violated. For the present purposes, it is also evoked here, but the consequences
of its being violated are examined below.

The third assumption, that the variance-covariance matrix $\sigma^2 \Phi$ of the input $y$
(Eq. (5) and Table III) is known to within the common factor $\sigma_y^2$, is the root of the
problem. The matrix $\Phi$ is only an estimate of the dispersion matrix, because it contains
the estimates $\hat{\sigma}_k^2$, $k = 1, 2, 3$, of the measurement precision of the three bands. In
the following sections, the effects of this inadequate assumption on the confidence limits
of (i) the merged estimates of the molecular constants $\tilde{b}^M$ and (ii) the merged estimate
of the variance $\hat{\sigma}_v^2$ are examined.

Confidence Limits for $\tilde{b}^M$

In least-squares fits of $f$ degrees of freedom where the above three assumptions are
met, it can be shown (9, p. 99) that the difference between the estimate $\hat{\beta}_i$ and its
unknown "true" value $\beta_i^{true}$, divided by the standard error $\hat{\sigma}_v^{-1}$, has a "t" distribution
on $f$ degrees of freedom, i.e.,

$$(\hat{\beta}_i - \beta_i^{true})/\hat{\sigma}_v^{-1} \sim t(f).$$

(21)

This means that if the set of measurements could be repeated many times in an identical
fashion, except for random measurement error, then the set of $(\hat{\beta}_i - \beta_i^{true})/\hat{\sigma}_v^{-1}$ values
would be distributed symmetrically about zero in a fashion described by the $t(f)$ function,
which is a commonly tabulated function of statistics. Stated alternatively and
equivalently, the $t(f)$ function describes the probability of occurrence of a given
$(\hat{\beta}_i - \beta_i^{true})/\hat{\sigma}_v^{-1}$ value. This property leads straightway (9, p. 107) to the familiar
confidence interval (4, Section D-3-b)

$$[\hat{\beta}_i - t(f, 1 - \alpha/2)\hat{\sigma}_v^{-1} \leq \beta_i^{true} \leq \hat{\beta}_i + t(f, 1 - \alpha/2)\hat{\sigma}_v^{-1}],$$

(22a)

commonly abbreviated in the physical sciences as

$$\hat{\beta}_i \pm t(f, 1 - \alpha/2)\hat{\sigma}_v^{-1};$$

(22b)

within which one can be $100(1 - \alpha)\%$ confident that the unknown "true" value $\beta_i^{true}$
lies. Thus, in order to use Eq. (22b) to construct the same well-defined confidence
limits for the merged estimates $\tilde{b}^M$, it must be shown that the fraction $(\hat{\beta}_i^M - \beta_i^{true})/
(\hat{\sigma}_v^M)^{-1}$ is very nearly $t$ distributed for the merged fit.

In principle, the probability distribution of $(\hat{\beta}_i^M - \beta_i^{true})/(\hat{\sigma}_v^M)^{-1}$ can be derived
algebraically using the approach that produced Eq. (21). However, the resulting
algebra becomes exceedingly complicated, and since the end result is unlikely to be a
commonly tabulated function, it would not have much utility. We have taken an
alternate approach. The distribution of $(\hat{\beta}_i^M - \beta_i^{true})/(\hat{\sigma}_v^M)^{-1}$ is generated by fitting
1000 sets of synthetic data described earlier. Each set differs from the others only
because of the randomly generated, normally distributed "measurement" errors of
mean zero and specified variance. Thus, the generated $(\hat{\beta}_i^M - \beta_i^{true})/(\hat{\sigma}_v^M)^{-1}$ distribu-

$^*$ The $t$ distribution is compared graphically to the similar and more familiar normal distribution
in Ref. (10, p. 81).
Fig. 2. The histogram represents the distribution of \((\hat{B}_0''^M - B_0''^{\text{true}})/(\hat{\Theta}_{77}^M)^{1/2}\) values obtained from merged band-by-band fits to 1000 sets of synthetic data with randomly different, normally distributed errors for the model system described in Table I and Fig. 1. The solid curve is a \(t\) distribution with seven degrees of freedom.

tion can be compared to a true \(t(f)\) distribution for a variety of typical "experimental" conditions to find out when the fraction \((\hat{\beta}_i^M - \beta_i^{\text{true}})/(\hat{\Theta}_{ii}^M)^{1/2}\) is approximately \(t\) distributed. For those conditions, a standard error will have its usual meaning, and well-defined confidence limits for the merged estimates can be constructed via Eq. (22b).

For the model band system defined in Fig. 1, the scatter of 1000 values of \((\hat{B}_0''^M - B_0''^{\text{true}})/(\hat{\Theta}_{77}^M)^{1/2}\) is given by the histogram in Fig. 2. The solid curve is a \(t\) distribution with seven degrees of freedom, which are the degrees of freedom of the second step of the merged fit. It is clear that the histogram and the \(t(7)\) distribution match very well. This comparison can be made more quantitatively with a standard goodness-of-fit test (11, pp. 90–92), which yields a \(X^2\) value of 5.8. For the type of test applied, \(X^2\) values outside of the range 3.8 to 21.9 indicate that one would reject the observed histogram as being a 1000-member sample from a \(t(7)\) distribution (with only a 5% risk of erroneously rejecting a histogram that was indeed a sample from a \(t(7)\) distribution) and would accept the histogram as being such for values of \(X^2\) within that range. Consequently, for this model system, the merged molecular constant \(\hat{B}_0''^M\) and standard error \((\hat{\Theta}_{77}^M)^{1/2}\) have the usual familiar meaning. For example, 947 of the 1000 values in Fig. 2 fall within \(\pm 2.365\), which are 95% points for a \(t(7)\) distribution. Similar behavior was found for the merged estimates of the other molecular constants.

To explore the range in which this statement of validity can be made for the merged estimates, many experimental parameters in different model systems were varied: rotational development, number of bands, relative precision, degree of redundancy in the band system, and number and type of molecular constants in the Hamiltonian. Among all of these, it was found that the \textit{degrees of freedom in the separate fit to each band}
MERGING SEPARATE LEAST-SQUARES RESULTS

FIG. 3. The histogram represents the distribution of \( \frac{(B_o^{\text{true}} - B_o^{\text{true}})}{(\hat{\Theta}_{77})^\frac{1}{2}} \) values obtained from merged band-by-band fits to 1000 sets of synthetic data with randomly different, normally distributed errors for the model system described in Table I and Fig. 1, except that each band has only three \( R \) lines and three \( P \) lines. The solid curve is a \( t \) distribution with seven degrees of freedom.

(i.e., Eqs. (1)–(4), the first step of the merged fit) was the most sensitive. As an example of a very extreme case, the model band system of Fig. 1 was used, except that each band had only six lines (three \( R \) lines and three \( P \) lines). Thus, the degrees of freedom of the three band-by-band fits were \( f_k = 1, k = 1, 2, \) and 3. Figure 3 shows the scatter of the 1000 \( \frac{(B_o^{\text{true}} - B_o^{\text{true}})}{(\hat{\Theta}_{77})^\frac{1}{2}} \) values for this situation. The solid curve is a \( t(7) \) distribution. The match is clearly much poorer than that of Fig. 2. There are fewer values near zero and more values at large deviations. Approximately 100 values lie outside the range of this figure. The goodness-of-fit test gave a \( \chi^2 \) value of 884, indicating a poor match. Only 798 values fall within \pm 3.65, indicating that these are about 89% points for the non-\( t \) distribution followed by \( \frac{(B_o^{\text{true}} - B_o^{\text{true}})}{(\hat{\Theta}_{77})^\frac{1}{2}} \) for this example, in contrast to being 95% points for a \( t(7) \) distribution. To construct a 95% confidence limit for the case at hand, one would have to use about \( f_k \). The merged estimates of the other molecular constants showed a similar behavior.

The example of one degree of freedom is obviously an extreme case. The situation rapidly gets better as the degrees of freedom in each of the band-by-band fits increase. For the model system in Fig. 1, for example, a close approximation to the \( t \) distribution occurs when \( f_k \gg 10 \). However, for a given \( f_k \), the comparison slowly deteriorates as the number of bands in the system increases. For an absorption band system with \( f_k = 15 \) for each band, the comparison is four times poorer (as indicated by \( \chi^2 \)) for six bands than three bands. Nevertheless, for \( f_k \gg 20 \), the comparisons are about the same. Thus, one can conclude from these and other tests that when the degrees of freedom in the fit to each band are typical for most spectroscopic applications, say \( f_k \gg 30 \), the esti-
Fig. 4. The histogram represents the distribution of 1000 $\delta_M^2$ values generated under the conditions described in Fig. 2. The solid curve is a $\chi^2/f$ distribution with seven degrees of freedom.

...mated molecular constants and standard errors have their usual statistical meaning; hence, confidence limits can be constructed in the standard way (4, Section D-3-b).

Confidence Limits for $\delta_M^2$

In addition to $\delta$, a least-squares fit produces an estimate $\delta^2$ of the variance of the measurement errors of the fitted data. This is an important estimate because it is probably the most valuable single indicator of the quality of the fit. In least-squares fits of $f$ degrees of freedom where the three assumptions mentioned above are met, it can be shown (9, pp. 99–100) that the estimate $\delta^2$ divided by its unknown "true" value $\sigma_{\text{true}}^2$ has a $\chi^2/f$ distribution, i.e.,

$$\frac{\delta^2}{\sigma_{\text{true}}^2} \sim \chi^2/f.$$  \hspace{1cm} (23)

This means that the probability of occurrence for a given $\delta^2/\sigma_{\text{true}}^2$ ratio is described by the $\chi^2/f$ function, which is again a common tabulated function of statistics. Unlike the symmetric $t$ function, which has a range of $-\infty < t < +\infty$ and a mean of zero, the asymmetric $\chi^2/f$ function has a range of $0 \leq \chi^2/f \leq \infty$ (since $\delta^2$ is positive definite) and a mean of unity. This property leads (10, pp. 102–103) to confidence limits in much the same way as for $\delta$, except that the confidence limits are asymmetric. For example, the $100(1 - \alpha)\%$ confidence limits for $\sigma_{\text{true}}^2$ are

$$\frac{\delta^2}{P(f, 1 - \alpha/2)} \leq \sigma_{\text{true}}^2 \leq \frac{\delta^2}{P(f, \alpha/2)},$$  \hspace{1cm} (24)

10 The $\chi^2/f$ distribution should not be confused with the $\chi^2$ values of the goodness-of-fit test. The $\chi^2/f$ distribution is graphed for $f = 1, 2,$ and 6 in Ref. (10, p. 102).
where $P(f, \alpha/2)$ and $P(f, 1 - \alpha/2)$ are the $100\alpha/2$ and $100(1 - \alpha/2)$ percentage points of a $X^2/f$ distribution. For example, the 95% confidence limits are $0.43782 < \sigma^2 < 4.15$ for $f = 7$. Thus, in order to use Eq. (24) to construct the same well-defined confidence limits for the merged estimate $\delta_M^2$, it must be shown that the ratio $\delta_M^2/\sigma^2_{true}$ is very nearly $X^2/f$ distributed for the merged fit.

For a least-squares fit that meets the above assumptions and for which the input data are weighted inversely by their true variances, $\sigma^2_{true}$ is effectively unity (4, Section E-4) and $\delta^2$ will scatter about unity as described by the $X^2/f$ distribution. This means that the limits in Eq. (24) would include unity $100(1 - \alpha)%$ of the time, or equivalently

$$P(f, \alpha/2) \leq \sigma^2 \leq P(f, 1 - \alpha/2).$$

For example, the 0.95 probability range for $\delta_M^2$ is $0.241 \leq \delta_M^2 \leq 2.29$ for $f = 7$. We can test whether Eqs. (24) and (25) hold for the merged fit, which weights by estimates of the true variances, by examining the distribution of $\delta_M^2$ from the tests described above. For the model band system defined in Fig. 1, the scatter of 1000 values of $\delta_M^2$ is given by the histogram in Fig. 4. The solid curve is a $X^2/7$ distribution. It is clear that the match is very good. The goodness-of-fit test yields a $X^2$ of 8.2, indicating that we can accept the histogram as a sample from a $X^2/7$ distribution. Consequently, for this model system, confidence limits for $\sigma_M^2$ are given by Eq. (24), or the percentage distribution of $\delta_M^2$ is given by Eq. (25). For example, 947 of the values in Fig. 4 fall within 0.241 and 2.29, which are the 2.5% and 97.5% points for a $X^2/7$ distribution.

However, just as was found above, the validity of these statements decreases when the degrees of freedom in the fit to each band are small. Figure 5 shows the extreme case.
of one degree of freedom in the fit to each of the three bands of the model system. The histogram of the $\delta M^2$ values only slightly resembles the $\chi^2/7$ distribution given as the solid curve and, understandably, the goodness-of-fit test yields a huge value for $\chi^2$. Values of $\delta M^2$ near unity are much less probable and large values are much more probable, in comparison to the $\chi^2/7$ distribution. Approximately 360 values lie outside of the right-hand range of this figure. Only 442 of the $\delta M^2$ values fall within 0.241 and 2.29, the 95% confidence limits of a $\chi^2/7$ distribution. To construct 95% limits for the $\delta M^2$ values in this example, the data of this histogram suggest that one would have to use about 0.22 and 150!

However, for larger degrees of freedom in the fit to each band, the comparison is much more favorable. For the model system, a close approximation ($\chi^2 \lesssim 20$) to the $\chi^2/f$ distribution occurs when $f_k \gtrsim 30$. Thus, this favorable match requires slightly higher $f_k$ than that for $\delta M$ examined above. It is also more sensitive to the number of bands in the band system. From numerous tests on a variety of model systems, it appears that Eqs. (24) and (25) apply reasonably well for the merged estimate $\delta M^2$ of the variance if the degrees of freedom in the band-by-band fits are greater than about 40.

The bands of light molecules, which generally have a small number of lines, often may not fall into this category. For those low-degree-of-freedom cases for which the proper establishment of confidence limits happens to be of crucial importance, it should be feasible to establish them using a simulation study much like the one done here for the model band system. Random, normally distributed errors of the appropriate variances can be added to the calculated line positions, which are selected to have a rotational structure identical to that of the observed system, and then a merge fit made. Perhaps as few as 500 repetitions of this will adequately define the non-$t$ and non-$\chi^2/f$ distributions of $(\hat{\beta}_i - \beta_{i, \text{true}})/(\hat{\delta}_{ii}^M)$ and $\delta M^2$ for the system at hand, and from these, proper confidence limits can then be established. For a small band system with a low number of lines per band, this simulation will not be expensive.

It may seem that the concern over the properties of $\delta M$ and $\delta M^2$ is a complicated feature of the merge method alone, and therefore the method has an unattractive complexity. However, such concerns are fundamental to all least-squares applications. The reason that simple and familiar unweighted least-squares fits appear to have avoided such concerns is that an assumption has been made (often implicitly) that all subblocks of the data have the same measurement precision. Thus, its simplicity is only a result of accepting an untested assumption.

Finally, it should be noted that just because the occurrence of $\delta M^2$ values from merged least-squares fits is described by a $\chi^2/f$ distribution for sufficiently large degrees of freedom in the fit to each band, it does not mean that the same can be said for the $\delta w^2$ values from the weighted simultaneous least-square fit to the same data. In fact, Eq. (18) shows that the scatter of the $\delta M^2$ and $\delta w^2$ values about unity must be very different, because $f_w$ is generally much larger than $f_M$. In particular, the scatter of the $\delta w^2$ values about unity is much smaller than that of the $\delta M^2$ values. Thus, the probability of occurrence for $\delta w^2$ is a distribution much narrower than a $\chi^2/f_w$. If one is not aware of this, it can be quite misleading, since, as will be shown below, $\delta^2$ values near unity are desirable indicators of the absence of serious systematic measurement or model problems. Thus, it should be understood that a $\chi^2/f_w$ distribution does not describe $\delta w^2$ values and that tests (described below) employing that distribution should not be applied to
a $\delta_w^2$ value without the scaling implied by Eq. (18). These same remarks apply to $t$-tests of $\hat{\beta}^w(=\hat{\beta}^M)$ using $\hat{\Theta}^w = \delta_w^W \hat{W}^w$, since $(\hat{\beta}^w - \beta^{true})/(\hat{\Theta}^wi^w)$) does not follow a $t(f_w)$ distribution, but rather a narrower distribution.

DETECTION OF RELATIVE SYSTEMATIC ERROR

The preceding section showed that, when the degrees of freedom of the fit to each band are in the range of most spectroscopic applications, the merged variance $\delta_{M}^2$ can be expected, with a chosen confidence, to lie in an asymmetric range about unity with limits given by the percentage points of the $\chi^2/f_M$ distribution, where $f_M$ are the degrees of freedom of the second step of the merged fit. This statement rests on the assumption of normally distributed measurement errors, which is frequently the case, and on the assumption of no significant relative systematic errors in either the measurements or the Hamiltonians used in fitting the measurements. This assumption, of course, often can be very inappropriate. Any test that can indicate the presence of such relative systematic errors would be extremely valuable, since most statistical indicators gauge only random measurement error. As examples, it would be particularly useful to know whether relative systematic measurement errors exist in an extensive band system, where the wide wavelength range creates the strong possibility of such errors, or whether undetected perturbations cause some of the Hamiltonians to be incomplete. We show in this section that such types of errors cause $\delta_M^2$ to lie outside of the range expected for purely random errors and that this can be used as a convenient test for the likelihood of their occurrence.

To test the sensitivity of the test, the synthetic data methods described above were used, except that various types of systematic errors were superimposed on the random errors. The first type of error considered is a constant 0.05 cm$^{-1}$ shift in the wave-numbers of band 1 of the model system described in Fig. 1. For this electronic band, therefore, the systematic shift is equal to one standard deviation of the random error. An error of this type is, of course, absorbed into and alters only the value of the origin of the band. Thus, for the band-by-band results given in Table II, the only change is that $\nu_0(0,0)$ changes from 12 000.007(10) to 12 000.057(10) cm$^{-1}$. The former value missed the "true" value by 0.007 cm$^{-1}$ due to the random errors. Now the latter value missed the "true" value by an additional 0.05 cm$^{-1}$ due to the systematic error. Since the three bands are cyclically interrelated (see Fig. 1), this additional 0.05 cm$^{-1}$ will perturb the entire merge fit over and above that expected from random errors alone.

Table VI shows the resulting merged constants. These should be compared to those in Table IV, which were determined without the systematic error. The values for $T_0'$, $B_0'$, and $D_0'$ are changed substantially, but the values for the other molecular constants are changed very little. This pattern can be understood in terms of the correlation coefficients. $T_0'$, which critically depends on the input value of $\nu_0(0,0)$, is rather strongly

\[11\] The discussion is limited to relative systematic errors. An example would be an error that varied systematically on wavelength. A constant systematic error in all of the measurements to be fitted is, of course, undetectable by statistical tests.

\[12\] Such an error would not extensively perturb a merged fit to a set of absorption bands with no common upper vibrational levels. Only the merged value for $\nu_0(0,0)$ would change. The reason is, of course, that there is no cyclic relation to any of the $\nu_0$ values of such a system.
correlated with $B'_0$ and $D'_0$ ($-0.515$ and $-0.500$, respectively), but only weakly correlated with the other constants ($0.069$ maximum, which is with $T_{1''}$, the only other constant that is slightly changed). The reason for this is that the $(0, 0)$ vibration-rotation band dominates the merged fit for the unchanged constants and the correlation coefficients reflect this fact. Although it is interesting to see how and where this systematic error propagates into the merged constants, these changes cannot be used as an indication of the presence of a systematic error because, of course, we would not know what the values for the constants would have been if there were no error!

Our interest is in the value of $\delta_M = 2.301$, or more particularly, $\delta_M^2 = 5.295$. In the preceding section, it was shown that one could reasonably (95% confidence) expect $\delta_M^2$ to lie in the range from $0.241$ to $2.29$, when the three assumptions discussed above are met. Not only is this one value outside of these limits, but $971$ of the $\delta_M^2$ values from the 1000 randomly different sets of measurement errors with the 0.05 cm$^{-1}$ systematic error in band 1 were also greater than $2.29$. Furthermore, the mean of the 1000 $\delta_M^2$ values is $4.90$, rather than the mean of unity of a $\chi^2/f$ distribution. Since the systematic error used in the example here is only one standard deviation, it should be clear that the test is a fairly sensitive one for this type of error. If the error had been two standard deviations, $\delta_M^2$ would have been $17.6$ for the present example. Thus, the sensitivity here is stronger than linear.

The second type of relative error considered here is one that systematically varies with wavenumber or rotational level. These could occur, for example, from differential emulsion shrinkage or other plate problems, or a change in the degree of blending along the branch, or an unrecognized perturbation not included in the Hamiltonian. We
simulate this type of error by introducing into the synthetic data of the (0, 0) electronic band of the model system an error that linearly increases with wavenumber from zero at the lowest wavenumber, with an average of 0.1 cm\(^{-1}\) (i.e., 2\(\sigma\)). In the fit to these data, \(\sigma_1 = 0.0528 \text{ cm}^{-1}\), hence is virtually unaffected by this type of relative systematic error and thus could give no indication of its presence, as it could if it had been several times larger than the expected measurement precision. Furthermore, the largest change in the estimated molecular constants was for \(\hat{B}_0' = 1.49981 \text{ cm}^{-1}\), which was a change of only one standard error. Comparison of this value to the value obtained for this constant from the (0, 1) electronic band would not indicate the possibility of an error. Thus, examination of the results of the band-by-band fits would not straightforwardly reveal that anything was awry.

However, the merged fit yields \(\hat{\sigma}_M^2 = 2.48\), which is outside of the range expected with 95\% confidence from \(\hat{\sigma}_M^2\) when the errors are purely random. In fact, 502 of the 1000 \(\hat{\sigma}_M^2\) values obtained from the randomly different fits with this type of error were larger than 2.29. The mean of these \(\hat{\sigma}_M^2\) values was 2.49, rather than unity.

Lastly, the effects of an error in the Hamiltonian of one of the bands were explored. A realistic problem of this type would be whether or not to include the constant \(H_r\). Values of \(H_r = 7 \times 10^{-13}, 6 \times 10^{-12}, \text{and } 5 \times 10^{-12} \text{ cm}^{-1}\) for \(H_y', H_1', \text{and } H_3'\), respectively, were added to the “true” constants in Table I that were used to calculate the line positions of the three bands. For bands 1 and 3, \(H_y'\) and \(H_1'\) were included in the Hamiltonian used in the least-square fit to each band. However, for band 2, it was not included. The repeated fits showed that this “model error” for band 2 produced no deleterious effects on either \(\hat{B}_y\), \(\hat{B}_y\), or \(\hat{\sigma}_M^2\). This is expected, of course, because these \(H_r\) values cannot produce significant effects for the relatively low-\(J\) lines of band 2. In this case, it should not matter whether \(H_r\) is included or omitted in the Hamiltonian of the second band. Hence, while the equations given herein have been expressed in terms of one Hamiltonian for each vibrational level, it is not necessary in practice to include statistically insignificant constants.\(^13\)

On the other hand, if the lines of band 2 are extended to include the first 50 \(R\) lines and \(P\) lines and the “true” standard errors of the three bands are decreased by a factor of 10, the role of \(H_r\) in band 2 becomes more important and the omission of \(H_y'\) and \(H_1''\) from the Hamiltonian produces erroneous results. However, this is not immediately obvious from the fit (without \(H_y'\) and \(H_1'\)) to band 2. For example, the other rotational constants “absorb” the absence of \(H_y'\) and \(H_1''\). This is expected, of course, because these \(H_r\) values cannot produce significant effects for the relatively low-\(J\) lines of band 2. In this case, it should not matter whether \(H_r\) is included or omitted in the Hamiltonian of the second band. Hence, while the equations given herein have been expressed in terms of one Hamiltonian for each vibrational level, it is not necessary in practice to include statistically insignificant constants.\(^13\)

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985 of the $\delta_M$ values were larger than the 2.29 limit expected if all of the least-squares assumptions were valid. The mean $\delta_M^2$ value was 5.2.

These tests show that certain types of relative systematic errors cause the expected $\delta_M^2$ distribution to skew toward larger values. Thus, if a fitted $\delta_M^2$ value for a particular set of laboratory data is larger than the upper value of a selected pair of confidence limits, one rejects the hypothesis that the errors are randomly produced and considers the possibility of systematic problems.

The size of the confidence limits are, of course, a factor in this decision and reflect a personal choice, in which one balances the desire to be specific against the fear of being wrong. The limits used above as examples corresponded to a 5% risk of erroneously concluding that something was wrong when in fact it was not. Wider limits, say 0.141 and 2.90, would reduce the risk of this type of erroneous conclusion to 1%, but would, of course, increase the risk of failing to detect that something was wrong when in fact it was. Hamilton (11, pp. 45–49) provides an elaboration on this balance.

The merge fit of the model system has only seven degrees of freedom. Usually, spectroscopic band systems are more extensive than this model and the merged degrees of freedom are much higher. For example, the application of the merged fit in the accompanying paper (6) has 95 degrees of freedom, for which the 95% confidence limits are much narrower, 0.74 and 1.31.

Of course, many other types of systematic errors can certainly occur. These examples suggest that the merged variance test should prove useful in indicating the presence of many of them. The test is, of course, blind to the exact location of the problem, but a strong indication that a problem exists can provide the needed incentive for what is often a tedious rechecking. Finally, even though a recheck of the measurements or Hamiltonians does not discover the source of the problem, we feel that being aware of the existence of the problem is nevertheless valuable information. One would know, for example, not to completely trust the statistical indicators, like standard errors and the confidence limits derived from them.

The extent to which the statistical meaning of spectroscopic standard errors is damaged by the presence of relative systematic errors is an interesting and clearly worthwhile subject for general study, but is beyond the present scope. We only note here that for the types and magnitudes of errors examined above, the confidence limits (Eq. (22b)) proved fairly reasonable; namely, the use of the fitted value $\delta_M^2$, which is larger than normal because of the relative systematic errors, to compute $\hat{\Theta}^M$ (Eq. (8)), yields standard errors14 large enough to encompass the "true" value with roughly the probability associated with the selected multiplicative $t$ factor. However, this should only be taken to suggest that some of the usual meaning of the standard error still exists even when moderate systematic errors are present. Without a detailed study, safety seems to dictate that when the $\delta_M^2$ test indicates that moderate systematic errors are present, somewhat more conservative confidence limits are desirable, e.g., from 99 to 99.99% limits rather than the popular 95%. (This means standard deviation multiples from 2.5 to 3.9, respectively, rather than the popular 2.0, for high degrees of freedom.) Ramsey (12) also discusses this point.

14 The increase of $\delta_M^2$ due to relative systematic problems may, of course, entirely offset the improvement of precision associated with the decrease of $\hat{\Theta}^M$ when the merged data set is enlarged.
SUMMARY AND CONCLUSIONS

The two-step merged band-by-band method is a way to determine the most precise, single values for the molecular constants of a system of \( n \) interconnected bands. Its first step, defined by Eqs. (1)-(4), is the set of familiar band-by-band least-squares fits to the \( n \) bands of the system, the output quantities of which are the \( n = 1, 2, 3, \ldots \) sets of molecular constants \( \hat{\beta}_k \) and accompanying variance–covariance matrices \( \hat{\Theta}_k \). These quantities serve as input to the second step, defined by Eqs. (5)-(10), which is a correlated least-squares fit that yields the minimum-variance single values \( \hat{\beta}^m \) for the molecular constants. The advantages offered by this method are as follows:

1. The first step is one that is already commonly done in most two-state reductions (see footnote 1) of a set of bands. However, one need not now stop at this point. The redundancy of most band systems can be utilized to obtain more precise estimates of the molecular constants by storing the values of \( \hat{\beta}_k \) and \( \hat{\Theta}_k \) from each of the final band-by-band fits and using them as input to the merge fit.

2. It is not necessary to go back to the line position measurements and fit all of them in a massive weighted simultaneous multiband fit (Eqs. (11)-(15)). The merged fit gives the equivalent results with much less data handling and computer storage requirements. However, the merged results are not equivalent to the unweighted simultaneous multiband fit, which is less desirable because it makes the unnecessary (and often incorrect) assumption that all of the bands are of the same measurement precision.

3. For most spectroscopic applications in which the degrees of freedom of the band-by-band fits are sizable (say, greater than about 40), the merged estimates \( \hat{\beta}^m \), \( \hat{\Theta}^m \), and \( \hat{\sigma}_M^2 \) have the usual statistical properties. This means that the standard errors \( (\hat{\Theta}^m)^{-1/2} \) can be used to construct confidence limits for the estimates \( \hat{\beta}^m \) (Eq. (22b)) that have their usual meaning. Similarly, confidence limits can be constructed for \( \hat{\sigma}_M^2 \) (Eq. (25)). Thus, these limits can be used in the customary way for model testing, rejection of spurious measurements, etc.

4. One of the more valuable of these tests is whether a fitted value \( \hat{\sigma}_M^2 \) is reasonable. In particular, if it lies substantially outside of the confidence limits expected for \( \hat{\sigma}_M^2 \) values whose variation about unity is due only to random measurement errors, then one can conclude that it is likely that relative systematic errors are present in some or all of the data sets or Hamiltonians.

5. Although the merge method has been presented here in the context of reducing a band system to molecular constants, it is actually a very general method having much wider application within spectroscopy and elsewhere. A few other applications can be mentioned. The procedure obviously can merge the fitted results from investigations of all types: bands from different or the same electronic systems, vibration–rotation, microwave, Raman, etc. However, when the set of results to be merged are from more and more heterogeneous sources, problems may naturally arise; e.g., the \( \hat{\Theta}_k \) matrices may not be available and systematic measurement and Hamiltonian differences are much more likely. For example, such problems can make the treatment of all of the inter-

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16 In storing the values of \( \hat{\beta}_k \) and \( \hat{\Theta}_k \), care must be taken to avoid the rounding off of digits that have relative significance (see Ref. (4), Sections G-1-d and G-2-a).
connected data for a molecule, with the goal of obtaining the best single constants for all of the vibrational levels, a frustrating will-o’-the-wisp. A final noteworthy application is that the method can merge band-by-band results into, not just molecular constants, but the best Dunham coefficients $Y_{ij}$ (i.e., $\omega_x, \omega_y x, \omega_y y, \ldots, B_e, \alpha_e, \gamma_e, \ldots$) for the data set, or any combination of molecular constants and Dunham coefficients. An example is given in the accompanying paper (6).

In conclusion, it should be clear that all possibilities and problems of the merged technique have not been explored here. However, enough has been done, hopefully, to demonstrate the potential of the technique to (a) properly utilize the redundancy of a band system and (b) to indicate the presence of relative systematic errors in a data set, which are two common problems faced routinely in the fitting of models to spectroscopic data.

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