

Dissociation energies and potential energy functions for the ground $X^1\Sigma^+$ and “avoided-crossing” $A^1\Sigma^+$ states of NaH

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A direct-potential-fit analysis of all accessible data for the $A^1\Sigma^+ - X^1\Sigma^+$ system of NaH and NaD is used to determine analytic potential energy functions incorporating the correct theoretically predicted long-range behaviour. These potentials represent all of the data (on average) within the experimental uncertainties and yield an improved estimate for the ground-state NaH well depth of $\mathcal{D}_e = 15797.4(\pm 4.3) \text{ cm}^{-1}$, which is $\sim 20 \text{ cm}^{-1}$ smaller than the best previous estimate. The present analysis also yields the first empirical determination of centrifugal (non-adiabatic) and potential-energy (adiabatic) Born-Oppenheimer breakdown correction functions for this system, with the latter showing that the A -state electronic isotope shift is $-1.1(\pm 0.6) \text{ cm}^{-1}$ going from NaH to NaD. © 2015 AIP Publishing LLC. [<http://dx.doi.org/10.1063/1.4906086>]

I. INTRODUCTION

Alkali hydrides are among the simplest diatomic molecules, and as such, they are of great interest as testing grounds for theory. The one with the smallest reduced mass is LiH, for which 80 years of experimental studies^{1–3} culminated the direct-potential fit (DPF) analysis by Coxon and Dickinson⁴ which yielded quantum-mechanically accurate analytic potential energy function and Born-Oppenheimer breakdown (BOB) correction functions that explain all of the available data for all four isotopologues (on average) to within the experimental uncertainties. The next smallest alkali hydride is NaH, and determining analogous quantum-mechanically accurate potential energy and BOB correction functions for it was the objective of the present work.

While NaH has piqued interest as an observable molecule in cool stars in recent years,^{5,6} it has been the object of study since the early 1930s.^{7,8} As is illustrated by Fig. 1, the potential energy functions of the lower $^1\Sigma^+$ states of the alkali halides are governed by a series of avoided crossings as the outer walls of each of the lowest $^1\Sigma^+$ states in turn each takes on a substantial degree of ionic character.^{2,9} In particular, the avoided crossings between the X - and A -state potentials give rise to the unusual shapes of the A -state potential minima that are more quartic than quadratic, which is the reason that their vibrational spacings and B_v values initially increase with v . In addition, the analogous avoided crossings with the C states have the effect of causing the A -state potentials to be abruptly cut off in order to allow them to approach the $\text{H}(1s) + \text{alkali}(np)$ asymptotes from below.

All previous empirical analyses of data for the $A^1\Sigma^+ - X^1\Sigma^+$ system of NaH were based on classical Dunham expansion fits,¹⁰ sometimes followed by application of the first-order semiclassical Rydberg-Klein-Rees (RKR) inversion procedure^{11,12} to obtain potential function points. In contrast, the present paper uses the fully quantum-mechanical “DPF” procedure of Refs. 13–15 to determine analytic

potential energy functions for both the $X^1\Sigma^+$ and $A^1\Sigma^+$ states that incorporate the correct theoretically known long-range behaviour,¹⁶ together with centrifugal and potential energy BOB functions for both states.

II. OVERVIEW OF PREVIOUS WORK

NaH was first observed in 1930 by Hori who reported absorption bands of the $A^1\Sigma^+ - X^1\Sigma^+$ system spanning four vibrational levels of the X state and 18 levels of the A state.⁷ In a paper on the NaH emission spectrum published the next year, he reported data for an additional band that he believed to be $0 \leftarrow 0$, increased all of his previous the A -state v' values by one unit⁸ and extended the range of the observed bands to span what was later shown to be¹⁷ the range $v'(A) = 3-20$ for $v''(X) = 0-3$. The first NaD data were obtained by Olsson in 1934,¹⁷ rovibrational bands for $v''(X) = 0-1$ and $v'(A) = 7-17$, which he reported together with new NaH measurements for $v''(X) = 0-1$ and $v'(A) = 7-17$. His first-order semiclassical analysis showed that Hori’s revised v' numbering for NaH had to be increased by 3 units. Following Pesl *et al.*,¹⁸ we have used uncertainties of $\pm 0.3 \text{ cm}^{-1}$ to weight most of the 2430 NaH transitions reported by Hori^{7,8} and Olsson¹⁷ and $\pm 0.05 \text{ cm}^{-1}$ for Olsson’s 604 NaD transitions.¹⁷

After a decade of inactivity during World War II, work on this system resumed with Pankhurst’s 1949 report¹⁹ of high temperature emission spectra of bands with $v''(X) = 3-8$ and $v'(A) = 1-7$ which had estimated line position uncertainties of $\pm 0.1 \text{ cm}^{-1}$, a factor of three smaller than that those for the earlier work.^{7,8,17} Unfortunately, most of his data have been lost, and all we were left with is a Deslandres table of band origins, a list of A -state rotational constants, and tables of combination differences for $v''(X) = 3-8$ and $v'(A) = 1-7$ with $J = 1-32$. While the latter comprise remarkable sets of pseudo-microwave $S(J)$ data spanning a large range of J values for a wide range of $v''(X)$ and $v'(A)$ levels, they seem to have been overlooked in some later studies.¹⁸

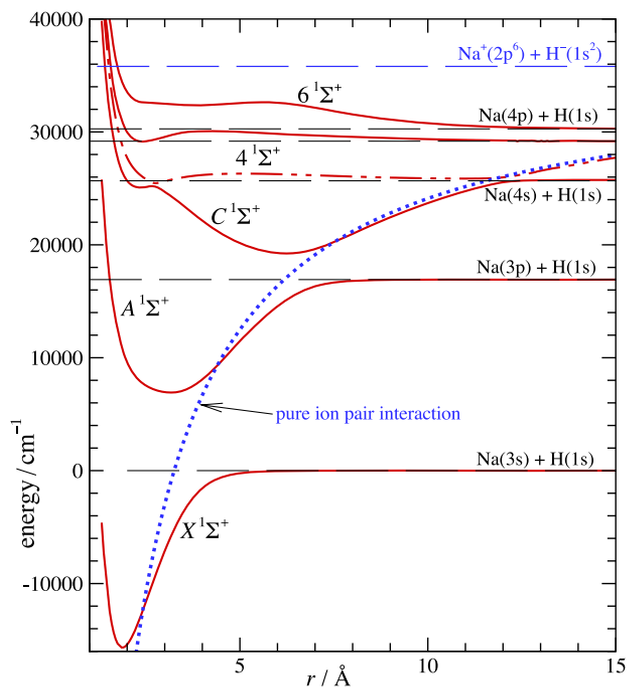


FIG. 1. Potential energy functions of the lowest six $1\Sigma^+$ states of NaH as calculated by Aymar, Deiglmayr, and Dulieu.⁹ solid red curves are states dissociating to neutral fragments, while the dashed-dotted-dotted curve is the lowest state dissociating to the ion-pair limit (blue long-dashed line) and the blue dotted curve shows the inward extrapolation of its limiting long-range behaviour.

After a further interregnum of 30 years, work on this system began in earnest in 1980 with Orth and Stwalley's observation of the $(v', v'') = (0, 6) - (0, 8)$ and $(1, 6) - (1, 8)$ bands in emission.²⁰ Although their $\pm 0.1 \text{ cm}^{-1}$ line position uncertainties were the same as those of Pankhurst,¹⁹ they did provide the first direct observation of the lowest vibrational level of the $A^1\Sigma^+$ state. A year later, the first microwave data for this system were reported by Sastry *et al.*²¹ While their data consisted of only the $R(0)$ line for each of $v''(\text{NaH}) = 0-3$, the $R(1)$ and $R(2)$ lines for $v''(\text{NaD}) = 0$ and the $R(2)$ line for $v''(\text{NaD}) = 1-3$, its very high precision led to much improved low-order Dunham constants and the first examination of Born-Oppenheimer breakdown effects and deviations from first-order semiclassical reduced-mass scaling for this system. In 1987, this work was extended by Leopold *et al.*²² who measured $R(J)$ transitions for $J = 0-6$ in $v = 0$, for $J = 0-7$ in $v = 1$, and for $J = 0-5$ in $v = 3$. Their work led to the determination of low-order molecular constants that were "... typically an order of magnitude more accurate than the best values previously available ..."²²

The first infrared vibrational measurements for NaH were performed in 1988 by Magg and Jones using a tunable diode laser spectrometer.²³ The $\pm 0.001 \text{ cm}^{-1}$ uncertainties of their data for 19 transitions of the vibrational fundamental and 7 transitions of the first hot band meant that when combined with the pure rotational data they yielded "... the most complete set of accurate values of the (low-order) ground-state Dunham parameters ..." to date.²³ A year later, Maki and Olson²⁴ measured the infrared spectrum of NaH using a Fourier transform spectrometer and extended

the data set to include transitions in the $3 \leftarrow 2$ band with uncertainties of $\pm 0.002 \text{ cm}^{-1}$. This led to the determination of a further-improved set of Dunham constants and the first systematic determination of BOB parameters for this system. However, as with all such parameter-expansion analyses, it was impossible to delineate between true BOB effects and those due to breakdown of first-order semiclassical reduced-mass scaling.

Since 1990, all experimental studies of NaH or NaD have involved measurements of their ultraviolet $A^1\Sigma^+ - X^1\Sigma^+$ spectra. First of all, in 1993, Rafi *et al.*²⁵ reported new measurements of $v''(X) = 0$ bands for $v'(A) = 13-22$ and of $v''(X) = 1$ bands for $v'(A) = 12-25$, in the form of band origins and A -state B_v and D_v rotational constants. Although their original line lists were lost,²⁶ since these measurements are the only observations of levels lying above $v'(A) = 20$, we chose to include their band origin data and to use their rotational constants to construct a set of synthetic pure rotational $\Delta J = 1-6$ A -state transitions for each of their bands, which we included in our data set. The uncertainties we associated with these synthetic data were $\pm 0.2 \text{ cm}^{-1}$, which is roughly the size of the uncertainties that Rafi *et al.*²⁵ had assigned to their band origins. Three years later, Lochbrunner *et al.*²⁷ reported measurements of NaH band absorption from $v''(X) = 0-3$ into $v'(A) = 2-15$ and of NaD bands for $v''(X) = 0-2$ into $v'(A) = 5-17$. However, their results were only reported in the form of lists of band origins and $B_{v'}$ and $D_{v'}$ values and a new set of Dunham coefficients. Fortunately, the data for their $(v', 0) = (8, 0) - (15, 0)$ bands were obtained privately for inclusion in the later analysis of Pesl *et al.*,^{18,28} but the data for their nine $v(A) > 0$ NaH bands and all of their twenty-five NaD bands are represented here only by their band origins. However, the precision of their results was only at the same *ca.* $\pm 0.3 \text{ cm}^{-1}$ level as the early data of Hori.^{7,8} The first high resolution ($\pm 0.006 \text{ cm}^{-1}$) electronic data for this system were those of Bahns *et al.* who reported the measurement of 141 transitions from $v(X) = 0$ and 1 into $v(A) = 3-9$ with $J'' = 0-30$.²⁹ Additional high-precision ($\pm 0.001 - 0.02 \text{ cm}^{-1}$) UV data were later obtained by Pesl *et al.*¹⁸ who applied a new variant of Doppler spectroscopy to laser-induced fluorescence from NaH molecules formed by reactive scattering of crossed beams of Na_2 and H atoms. They observed 280 transitions from bands for $v''(X) = 2-8$ and $v'(A) = 6-9$ and combined their results with all accessible earlier data to obtain a further improved sets of Dunham constants and RKR turning points.

Most recently, Huang *et al.*³⁰ used fluorescence depletion spectroscopy and stimulated emission pumping to delineate the properties of levels in the upper portion of the ground-state potential well. However, their analysis ignored the information about the $A^1\Sigma^+$ state that was contained in their data. In particular, they added the *differences* between their pump and probe laser frequencies to estimates of $X^1\Sigma^+$ state term values for the lower levels of their pump transitions that were calculated from the spectroscopic constants of Pesl *et al.*¹⁸ and fitted the resulting set of term values to conventional Dunham $(v + 1/2)$ polynomials for G_v and B_v (ignoring centrifugal distortion). Their observation and inclusion of data for $v(X) = 21$ certainly made their X -state

analysis the most comprehensive to date. However, their neglect of the older data and of the information about the $A^1\Sigma^+$ state and their reliance on term values generated from previously reported molecular constants¹⁸ rather than on the actual data on which they were based were weaknesses of their approach.

In all previous studies of this system, the analyses were based on historic parameter-fitting Dunham-type treatments,^{10,31} and with two exceptions,^{21,24} when isotope effects were considered at all it was assumed that they could be accounted for by first-order semiclassical reduced-mass scaling.^{10,13} In contrast, the present work reports the first DPF analysis of this system, which is fully quantum-mechanical, and whose basic results are analytic expressions for the potential energy curves and any relevant BOB functions for the state(s) in question. BOB effects are particularly important in molecules with small reduced mass, especially hydrides, and a combined-isotopologue DPF analysis allows them to be taken into account properly. In addition, the $A^1\Sigma^+$ of the alkali hydrides is known to have an abnormal “truncated” potential energy curve, and for such cases, no finite Dunham expansions can readily represent all of its level energies, while a DPF analysis using an appropriate model potential form can. The purpose of this paper is therefore to present a combined-isotopologue DPF analysis of all available data for this system that yields highly accurate and compact analytic representations of the potential energy curves for the $X^1\Sigma^+$ and $A^1\Sigma^+$ of NaH.

III. SELECTION AND TREATMENT OF DATA

The properties and sources of the data used in the present analysis are summarized in Table I. The uncertainties shown there were taken from the appropriate experimental paper and were used for all lines from that source, with the following exceptions. In the case of the electronic band data, if a line had a discrepancy much larger than its neighbours and another line in that data set had the same energy, its uncertainty was multiplied by a factor of three. This modification affected 58 of the Hori NaH data and 11 of the Olsson NaD data. Similarly, if a relatively large discrepancy occurred for a line associated with the highest observed J -level in a given branch, its uncertainty was multiplied by a factor of two. This second adjustment only affected 3 of the Hori data.^{7,8} Finally, if a line had a discrepancy from our best-fit model of more than 8 times the average uncertainty for its band/group, we concluded that it was an outlier (possibly mis-assigned) and de-weighted it out of the final fits. This led to removal of 1 Hori NaH datum, 13 Olsson NaD data, 5 Pesl *et al.* NaH data, 4 of the Pankhurst “combination difference” pure rotational data, and 15 of the Bahns *et al.* data, including the three assigned to $v' = 10$ and 12 for which the discrepancies of $>10\text{ cm}^{-1}$ show that they were completely mis-identified. Moreover, while all of the synthetic $A^1\Sigma^+$ -state pure rotational Rafi²⁵ data were consistent with our best models (within the assumed 0.2 cm^{-1} uncertainties), the discrepancies of his band origins for $v(A) = 0-20$ led us to assign them uncertainties of $\pm 1\text{ cm}^{-1}$, while the much larger

TABLE I. Experimental data used in the final analyses of the present work.

Isotope	Type	Uncertainty (cm^{-1})	$v(A^1\Sigma^+)$	$v(X^1\Sigma^+)$	J range	No. Data	Source
NaH	Electronic	0.3–0.9	3–20	0–3	0–37	1905	Hori ^{7,8}
	Electronic	0.3	4–13	0–1	0–24	524	Olsson ¹⁷
	Electronic	1.0	1–7	3–8	0	27	Pankhurst ¹⁹
	Pure rotational	0.1	...	3–8	0–32	149	Pankhurst ¹⁹
	Pure rotational	0.1	1–7	...	0–32	186	Pankhurst ¹⁹
	Electronic	0.1	0–1	6–8	0–33	261	Orth ²⁰
	Pure rotational	7×10^{-6}	...	0–3	0–1	4	Sastry ²¹
	Pure rotational	$(3-15) \times 10^{-6}$...	0–3	0–8	27	Leopold ²²
	Vib-rotational	0.001–0.009	...	0–2	0–15	26	Magg ²³
	Vib-rotational	0.001–0.002	...	0–3	0–21	58	Maki ²⁴
	Electronic	1.0	12–20	0–1	0	17	Rafi ²⁵
	Synthetic MW	0.2	13–25	0–1	0–6	144	Rafi ²⁵
	Electronic	0.3–0.6	8–15	0	0–25	298	Lochbrunner ²⁷
	Electronic	0.3–0.4	2–9	1–3	0	9	Lochbrunner ²⁷
	Electronic	0.006 & 0.31	3–9	0–1	0–22	126	Bahns ²⁹
	Electronic	0.001–0.017	2–8	6–9	1–29	275	Pesl ¹⁸
	Electronic “pump”	3.0	7–12	0 & 9	0–14	50	Huang ³⁰
	Electronic “probe”	2.0	7–12	10–21	1–14	277	Huang ³⁰
Vib-rotational	1.0	...	0, 9–21	1–14	286	Huang ³⁰	
						Total:	4642
NaD	Electronic	0.05–0.15	7–17	0–1	0–29	591	Olsson ¹⁷
	Pure rotational	0.000 007	...	0–3	1–3	5	Sastry ²¹
	Electronic	0.30–0.50	4–17	0–4	0	25	Lochbrunner ²⁷
						Total:	621
Total overall:						5263	

discrepancies (increasing from -4 to -60 cm^{-1}) for his 7 $v(A) = 21$ – 25 band origins led us to deweight them completely out of the final data set. All of the other older (pre 2010) data retained in our analysis were weighted by the uncertainties shown in column-3 of Table I, while numbers shown in its column-7 are the numbers of each type of data used in the final analysis.

The present work utilizes the data of Huang *et al.*³⁰ in a much more direct way than they did.³² First of all, following their approach, we use the *differences* between their pump and probe transition energies to define a set of 286 pseudo-vib-rotation transitions from levels with $v(X) = 0$ and $J = 0$ – 14 into those with $v(X) = 9$ – 21 . However, rather than represent the properties of the lower levels of the X state by a previously reported set of Dunham expansion parameters, we have relied in our inclusion of all of the earlier data for this system to define the lower portion of that potential energy well. Moreover, rather than neglect information about the A state, we have also included their 50 “pump” transitions from $v(X) = 0$ and 9 into $v(A) = 7$ – 12 and their 277 $v(A) = 7$ – 12 into $v(X) = 9$ – 21 “probe” transitions as separate data. As shown in Table I, we have assigned uncertainties of 1.0, 3.0, and 2.0 cm^{-1} to these three components of their data, within which they are in good agreement with our best models. Listings of our entire data set and of the differences between those data and our recommended model are included in the supplementary material associated with this paper.³³

IV. DIRECT-POTENTIAL-FIT DATA ANALYSIS

A. The radial Hamiltonian

Most spectroscopic data may be described in terms of differences between eigenvalues of potential energy function(s) for the electronic state(s) involved in the transition. However, for species of small reduced mass, especially hydrides, BOB effects give rise to differences between the effective potential energy functions for different isotopologues and introduce isotopologue-dependent corrections to the “mechanical” centrifugal potential. The present work uses Watson’s formulation of this problem,^{34,35} in which the effective radial Schrödinger equation for isotopologue- α of a diatomic molecule A-B may be written in the form³⁶

$$\begin{aligned} \hat{H} \psi_{v,J}(r) &= \left\{ -\frac{\hbar^2}{2\mu_\alpha} \frac{d^2}{dr^2} + \left[V_{\text{ad}}^{(1)}(r) + \Delta V_{\text{ad}}^{(\alpha)}(r) \right] \right. \\ &\quad \left. + \frac{[J(J+1)]\hbar^2}{2\mu_\alpha r^2} \left[1 + g^{(\alpha)}(r) \right] \right\} \psi_{v,J}(r) \\ &= E_{v,J} \psi_{v,J}(r). \end{aligned} \quad (1)$$

Here, $V_{\text{ad}}^{(1)}(r)$ is the effective adiabatic internuclear potential for a chosen reference isotopologue (denoted $\alpha = 1$), $\Delta V_{\text{ad}}^{(\alpha)}(r)$ is the difference between the effective adiabatic potential for isotopologue- α and that for the chosen reference isotopologue, μ_α is the normal reduced mass of atoms A and B with masses $M_A^{(\alpha)}$ and $M_B^{(\alpha)}$, and $g^{(\alpha)}(r)$ is the effective non-adiabatic correction term to the centrifugal potential for isotopologue α . As was shown by Watson, $\Delta V_{\text{ad}}^{(\alpha)}(r)$ and $g^{(\alpha)}(r)$ may each be written a sum of terms associated with the two atoms.^{34,35}

We find it most convenient to express those terms using the mass-difference and mass-ratio representation of Ref. 36,

$$\Delta V_{\text{ad}}^{(\alpha)}(r) = \frac{\Delta M_A^{(\alpha)}}{M_A^{(\alpha)}} \tilde{S}_{\text{ad}}^A(r) + \frac{\Delta M_B^{(\alpha)}}{M_B^{(\alpha)}} \tilde{S}_{\text{ad}}^B(r), \quad (2)$$

$$g^{(\alpha)}(r) = \frac{M_A^{(1)}}{M_A^{(\alpha)}} \tilde{R}_{\text{na}}^A(r) + \frac{M_B^{(1)}}{M_B^{(\alpha)}} \tilde{R}_{\text{na}}^B(r), \quad (3)$$

in which $\Delta M_{A/B}^{(\alpha)} \equiv M_{A/B}^{(\alpha)} - M_{A/B}^{(1)}$ is the difference between the masses of atom A or B in isotopologue- α and in the reference isotopologue, $\tilde{S}_{\text{ad}}^{A/B}(r)$ represents the “adiabatic” potential-energy BOB correction function for atom A or B, and $\tilde{R}_{\text{na}}^{A/B}(r)$ is the (dimensionless) “non-adiabatic” centrifugal BOB radial strength function for atom A or B.

B. Data analysis procedure

The present analysis involved application of a standard DPF procedure.^{14,15,37} Observed transition energies are compared to eigenvalue differences generated by solving Eq. (1) using parameterized analytic trial potential energy and BOB functions, and a standard non-linear least-squares procedure is used to optimize those parameters.¹⁴ The partial derivatives of the eigenvalues with respect to the parameters of the model required by this procedure are readily calculated using the standard Hellmann-Feynman theorem expression: $\frac{\partial E_{v,J}}{\partial p_j} = \left\langle \psi_{v,J}(r) \left| \frac{\partial V(r)}{\partial p_j} \right| \psi_{v,J}(r) \right\rangle$. The preliminary trial potential-function parameters required to initiate such fits may be generated by fitting the chosen functional form to a set of turning points generated by application of the semiclassical “RKR” inversion procedure^{11,12} to a conventional set of molecular constants, or by *ab initio* calculations.³⁸

For a given model, we characterize the quality of fit by the “dimensionless root-mean-square deviation,”^{14,15,37} A “good” fit is one for which $\overline{dd} \lesssim 1$. However, if \overline{dd} is persistently significantly greater than unity for a particular subset of the data, independent of the number of free parameters in the model, it usually means that the estimated uncertainties for that group of data are too small and should be increased. This was the basis of the data-weighting/data-selection procedure described in Sec. III.

C. The potential energy function form

It is well known that the outer walls of the potential energy wells of the $A^1\Sigma^+$ states of all of the alkali hydrides change shape abruptly, just before they rise past the energy of the alkali $nP_{1/2}$ atomic state because of an avoided crossing between a strongly attractive $^1\Sigma^+$ “diabatic” ion-pair state and a $^1\Sigma^+$ van der Waals state that dissociates to the alkali $nP_{1/2} + H(2S)$ threshold (see Fig. 1).^{2,9,39} Because this unusual behaviour would seem to preclude normal merging to inverse-power-sum long-range behaviour, we have chosen to represent the potential energy function for this state by an “Expanded Morse Oscillator” (EMO) function. This function has the form of a simple Morse potential whose exponent coefficient is a

function of distance

$$V_{\text{EMO}}(r) = \mathcal{D}_e \left(1 - e^{-\beta_{\text{EMO}}(r) \cdot (r - r_e)} \right)^2, \quad (4)$$

in which \mathcal{D}_e is the well depth, r_e is the equilibrium internuclear distance, and the exponent coefficient

$$\beta_{\text{EMO}}(r) = \sum_{i=0}^{N_\beta} \beta_i y_q^{\text{ref}}(r)^i \quad (5)$$

is expressed as a power series in a dimensionless variable

$$y_q^{\text{ref}}(r) = \frac{r^q - r_{\text{ref}}^q}{r^q + r_{\text{ref}}^q} \quad (6)$$

that is defined by a selected expansion centre r_{ref} and power q .

The free fitting parameters of this potential form are \mathcal{D}_e , r_e , and the expansion coefficients β_i of Eq. (5), while the values of N_β , q , and r_{ref} are optimized manually. There are no rules governing the choice of q and r_{ref} other than that experience shows that “too small” values of the former ($q \lesssim 2$) sometimes lead to potential functions that behave badly in the extrapolation region(s) outside the “data range,” the region bounded by the inner and outer turning points of the highest vibrational level involved in the analysis, and that we would normally expect the latter to lie within this “data range.” The geometric mean of those two turning points is usually a good initial trial value for r_{ref} .

In contrast with the $A^1\Sigma^+$ state, the potential energy function for the $X^1\Sigma^+$ state of NaH is expected to show the normal merging to inverse-power-sum long-range behaviour, so we represent it by the Morse/Long-Range (MLR) function that has the general form of a Morse-type potential with two key differences: (i) the exponent distance factor $(r - r_e)$ is replaced by a dimensionless variable that is also linear in r near r_e , but approaches a constant as $r \rightarrow \infty$, and (ii) there is a pre-factor in front of the exponential term that defines the long- and very-short-range behaviour of the potential. This function has the form

$$V_{\text{MLR}}(r) = \mathcal{D}_e \left(1 - \frac{u_{\text{LR}}(r)}{u_{\text{LR}}(r_e)} e^{-\beta_{\text{MLR}}(r) \cdot y_p^{\text{eq}}(r)} \right)^2, \quad (7)$$

in which \mathcal{D}_e and r_e are as defined above, while the radial distance variable has the same form as that of Eq. (6), except that its expansion centre is fixed at r_e and it is defined by a different integer power, p ,

$$y_p^{\text{eq}}(r) = \frac{r^p - r_e^p}{r^p + r_e^p}. \quad (8)$$

The exponent coefficient function in Eq. (7) is then defined by the expression

$$\begin{aligned} \beta_{\text{MLR}}(r) &= \beta_{p,q}^{\text{ref}}(r) \\ &= y_p^{\text{ref}}(r) \beta_\infty + [1 - y_p^{\text{ref}}(r)] \sum_{i=0}^{N_\beta} \beta_i y_q^{\text{ref}}(r)^i, \end{aligned} \quad (9)$$

in which

$$\beta_\infty = \lim_{r \rightarrow \infty} \beta_{\text{MLR}}(r) \equiv \ln \left\{ \frac{2\mathcal{D}_e}{u_{\text{LR}}(r_e)} \right\} \quad (10)$$

and the expansion variable $y_q^{\text{ref}}(r)$ is again defined by Eq. (6), while the variable $y_p^{\text{ref}}(r)$ has that same form, but is defined

by the same power p used to characterize the radial distance variable of Eq. (8), rather than by q (although one could choose to set $q = p$). The three dimensionless radial variables are defined as they are because of the effect of the integer powers p and q on the long-range behaviour of the exponential term in Eq. (7).⁴⁰

The pre-factor to the exponential term in Eq. (7) is defined by the expression

$$u_{\text{LR}}(r) = \sum_{m=m_1}^{m_{\text{last}}} D_m(\rho r) \frac{C_m}{r^m} \quad (11)$$

in which the powers $\{m\}$ characterize the inverse-power terms that theory dictates define the limiting long-range behaviour of the particular molecular state, and the associated coefficients C_m are (usually) known from theory. In the present work, the “damping functions” $D_m(\rho r)$ are represented by the “ $s = -1$ generalized Douketis-type functions” recommended by Ref. 40, and the system-dependent parameter used to account for the sizes of the overlapping electron clouds of the interacting atoms is defined (following Refs. 40 and 41) as $\rho = \rho_{\text{NaH}} = 0.69$. For the interaction of ground-state H and Na atoms, the leading terms in Eq. (11) correspond to $m = 6, 8$, and 10 , and we use the excellent values for those C_m coefficients reported by Mitroy and Bromley¹⁶ to define the long-range tail function $u_{\text{LR}}(r)$ of our MLR potential for ground-state NaH.

The free fitting parameters in the MLR model are \mathcal{D}_e , r_e , and the β_i expansion parameters of Eq. (9), while the values of N_β , p , q , and r_{ref} are set manually and the C_m coefficients are fixed at their theoretical values.¹⁶ The only firm constraint on the values of the integer powers p and q is that the power p must be larger than the difference between the last and first powers in the (damped or undamped) inverse-power sum of Eq. (11), $p > (m_{\text{last}} - m_i)$, in order to prevent the long-range behaviour of the exponential term in Eq. (7) from affecting that defined by the expansion of Eq. (11). However, as with the EMO potential form, “small” values of q ($q \lesssim 2$) sometimes tend to lead potential functions that behave badly in the extrapolation region(s) outside the “data range,” while too-large values make the $y_q^{\text{ref}}(r)$ expansion variable somewhat “stiff,” so that larger values of the power N_β are required to give an equivalently “good” potential energy function.

D. BOB radial strength function forms

Following Ref. 42, the radial strength functions characterizing the atom-dependent potential-energy and centrifugal BOB corrections of Eqs. (2) and (3) are expanded in the same form utilized for the exponent coefficient-function of the MLR potential,

$$\tilde{S}_{\text{ad}}^{\text{A}}(r) = y_{p_{\text{ad}}}^{\text{eq}}(r) u_{\infty}^{\text{A}} + [1 - y_{p_{\text{ad}}}^{\text{eq}}(r)] \sum_{i=0}^{N_{\text{ad}}^{\text{A}}} u_i^{\text{A}} y_{q_{\text{ad}}}^{\text{eq}}(r)^i, \quad (12)$$

$$\tilde{R}_{\text{na}}^{\text{A}}(r) = y_{p_{\text{na}}}^{\text{eq}}(r) t_{\infty}^{\text{A}} + [1 - y_{p_{\text{na}}}^{\text{eq}}(r)] \sum_{i=0}^{N_{\text{na}}^{\text{A}}} t_i^{\text{A}} y_{q_{\text{na}}}^{\text{eq}}(r)^i. \quad (13)$$

Since NaH is not an ion, $t_{\infty}^{\text{A}} = t_{\infty}^{\text{B}} = 0$ for both the $X^1\Sigma^+$ and $A^1\Sigma^+$ states.⁴² Moreover, since we adopt the convention of

defining the absolute zero of energy as the energy of ground state atoms at infinite separation, by definition, $u_\infty^A = u_\infty^B = 0$ for all molecular states that dissociate to that limit, and since Na has only one stable isotope, this is also true for the $A^1\Sigma^+$ state.⁴² Within this convention, the values of $u_0^{A/B}(X)$ define the isotopologue-dependence of the ground state well depth. However, since Na has only one isotope, $u_0^{\text{Na}}(X) = 0$, and since the NaD data for the $X^1\Sigma^+$ state span only a small fraction ($\lesssim 23\%$) of the well, we cannot expect to be able to determine the H/D isotope dependence of that well depth from these data, so we also fixed $u_0^{\text{H}}(X) = 0$. However, our fitted value of $u_0^{\text{H}}(A)$ (see below) provides the best current estimate of the H/D electronic isotope shift for the $A-X$ system.

Other than the general caveat that too small values may lead to unphysical behaviour in the extrapolation regions outside the “data range,” while too large values may tend to a need for higher polynomial orders in Eqs. (12) and (13),⁴² there are no explicit constraints on the values of p_{na} , q_{na} , q_{ad} . However, since p_{ad} defines the limiting inverse-power long-range behaviour of the $S_{\text{ad}}^A(r)$ functions, it should normally be set equal to the power of the leading term in Eq. (11), $p_{\text{ad}} = m_1$ to ensure that the effective adiabatic potentials $V_{\text{ad}}^{(\alpha)}$ have the same limiting long-range functional behaviour for all isotopologues.

V. RESULTS

A. Preliminary $X^1\Sigma^+$ single-state analysis

Because of the unusual shape of the $A^1\Sigma^+$ -state potential function, we began our analysis with fits that treated all of its vibration-rotation levels as independent term values. This allowed us to optimize our representation of the potential energy and centrifugal BOB functions for the $X^1\Sigma^+$ state, independent of any assumptions about a representation for the $A^1\Sigma^+$ state. Moreover, since the highest observed levels of the $A^1\Sigma^+$ state lie more than 1000 cm^{-1} below its asymptote, and the distance between the potential asymptotes is the accurately known atomic excitation energy of Na, this will have no significant effect on the determination of the ground-state dissociation energy.

The lower panel of Fig. 2 shows how our quality-of-fit parameter \overline{dd} varies with the order of the MLR exponent polynomial $N = N_\beta$ and the location of expansion the centre r_{ref} for two different combinations of the expansion coefficient powers p and q . As expected, we see that for *any* choice of r_{ref} increasing N_β improves the quality-of-fit, and that both families of models seem to be converging to the same optimum value of \overline{dd} . Following the arguments presented at the end of Sec. IV C, all of the cases considered here involve

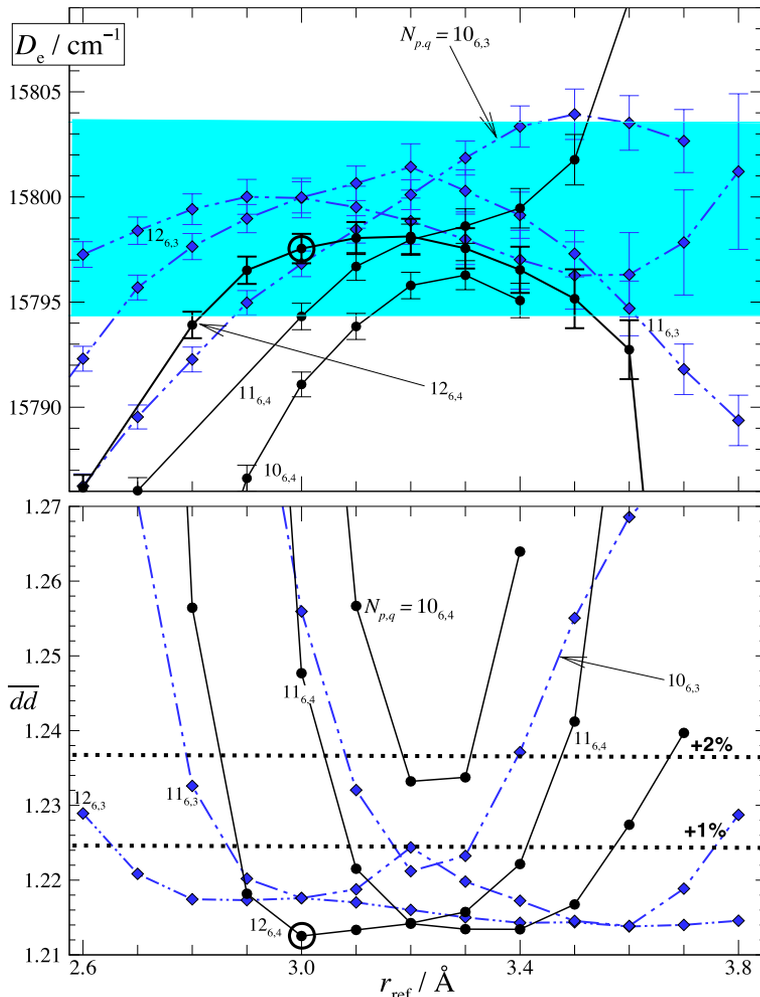


FIG. 2. Lower panel: dependence of \overline{dd} on the exponent polynomial order $N = N_\beta$ and the expansion centre location r_{ref} for expansion function variables defined by $\{p, q\} = \{6, 3\}$ and $\{6, 4\}$, in single-state fits to data for the $X^1\Sigma^+$ state. Upper panel: fitted values of D_e and their associated 95% confidence limit uncertainties for cases considered in the lower panel.

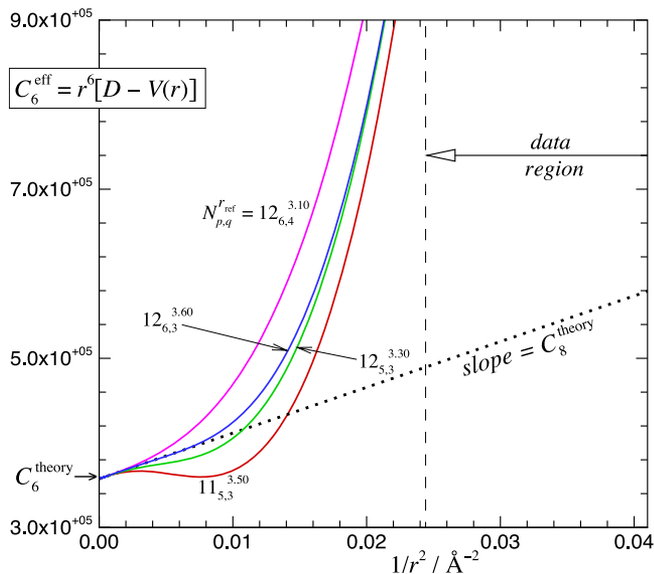


FIG. 3. Test of the long-range extrapolation behaviour of selected optimized model potential energy functions for the $X^1\Sigma^+$ state of NaH. In the symbols $N_{p,q}^{ref}$, $N = N_\beta$ indicates the exponent polynomial expansion order, r_{ref} is the expansion centre location, while p and q are the integers defining the radial variables of Eqs. (7)–(9).

$p > (m_{last} - m_1) = 4$ (in particular, $p = 5$ or 6), and the values of q are all > 2 . However, there are no obvious trends with p or q .⁴³

In addition to minimizing \overline{dd} for the smallest possible number of parameters, a key criterion in selecting an optimum potential function model is that it should behave sensibly in the extrapolation interval past the outer end of the data region. A simple way of testing for that is to examine the behaviour of a plot of $\{C_6^{eff}(r) \equiv r^6[D - V_{MLR}(r)]\}$ vs. $1/r^2$. The structure of Eq. (11) means that as $1/r^2 \rightarrow 0$ ($r \rightarrow \infty$), plots of this type must approach an intercept of C_6^{theory} with a slope of C_8^{theory} , where C_m^{theory} are the values of the fixed coefficients defining the two leading terms in the expansion of Eq. (11). Figure 3 shows that all of the potentials considered there do have this expected limiting behaviour, but that the models for which $p = 5$ tend to dip down and approach the limiting slope from below. This is unphysical, since the leading deviation from the limiting (linear) behaviour should show positive curvature, due to the positive (attractive) C_{10}^{theory} term in the long-range potential. We therefore select a MLR function with $N = N_\beta = 12$, $p = 6$, $q = 4$, and $r_{ref} = 3.0$ (circled points on Fig. 2) as our recommended model for the potential energy function of the $X^1\Sigma^+$ state of NaH.

The upper panel of Fig. 2 shows that the model-dependence of the fitted values of the ground-state well depth \mathcal{D}_e is far larger than the uncertainties in virtually any of the values yielded by the individual fits.⁴³ In order to obtain the best possible estimate of the overall uncertainties in the physically interesting parameters \mathcal{D}_e and r_e , we therefore adopted the averaging-over-models procedure of Eqs. (6)–(8) of Ref. 44 (or Eqs. (21)–(23) of Ref. 15). On averaging over the results for the 63 cases for which the value of \overline{dd} was no more than 1% larger than that for the recommended model (points below the +1% line in the lower panel of Fig. 2), we obtain $\text{unc}\{\mathcal{D}_e\} = 4.3 \text{ cm}^{-1}$ and $\text{unc}\{r_e\} = 0.0000015 \text{ \AA}$. Extending this averaging to include result for the 82 models

whose \overline{dd} values were within 2% of that for the recommended model (points below the +2% line in the lower panel of Fig. 2) only changed $\text{unc}\{\mathcal{D}_e\}$ in its second decimal place and raised $\text{unc}\{r_e\}$ to 0.0000020 \AA . The shaded region in the upper panel of Fig. 2 indicates the band of uncertainty associated with averaging over the 63 models associated with the 1% limit.⁴³

With an optimum X -state potential function model selected based on the fits to NaH data alone described above, we undertook a preliminary attempt to determine the number of BOB terms required to simultaneously explain those data. This study showed that either three or four potential-energy BOB terms and one centrifugal BOB term would be required. However, because of the expected high degree of correlation between the numbers and values of the BOB parameters for the X and A states, the delineation of optimum BOB models for the $X^1\Sigma^+$ state had to await the global two-state analysis described below.

B. Two-state analysis and determining the $A^1\Sigma^+$ state potential energy function

Using the recommended model for the $X^1\Sigma^+$ -state potential determined above, we then proceeded to perform full two-state, 2-isotopologue fits to determine an optimum model for the $A^1\Sigma^+$ potential energy function. As discussed at the beginning of Sec. IV C, we have chosen to use an EMO function to represent the potential function for this state. While the values of \mathcal{D}_e , r_e , and the well-shape parameters β_i are all determined automatically using our least-squares fitting program DPotFit,¹⁴ as with the $X^1\Sigma^+$ -state fits described above, optimum values for the expansion centre r_{ref} and power q defining the dimensionless radial variable of Eq. (6) were determined manually. One general finding was that for $q = 2$ or 3 , all of the fitted EMO potentials had an inflection point on the inner wall above the potential asymptote, which led to a spurious inner maximum and potential function turnover in the short-range extrapolation region, so those models were physically unacceptable. This was also true for the $q = 4$,

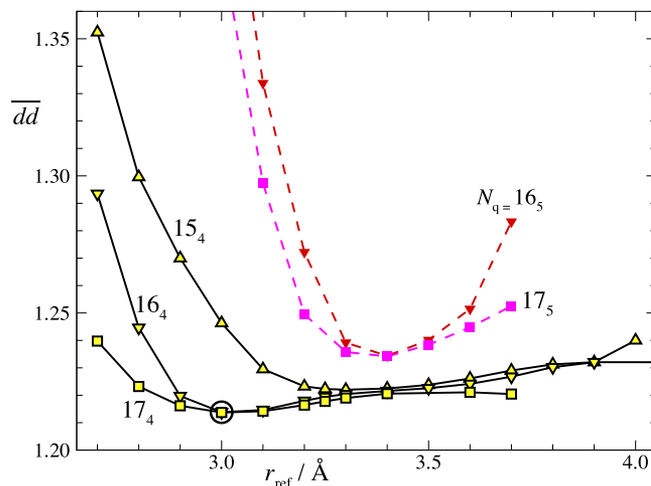


FIG. 4. Dependence of \overline{dd} on the exponent polynomial order $N = N_\beta$ and the expansion centre location r_{ref} for two choices of the expansion function parameter q for the $A^1\Sigma^+$ state in two-state fits with the $X^1\Sigma^+$ -state model fixed, but its parameters free.

$N_\beta = 17$ models for $r_{\text{ref}} < 2.9$ Å. However, the remaining $N_\beta = 17$, $q = 4$ models, all of the $q = 4$, $N_\beta = 15$ and 16 models, and all of the $q = 5$ models considered, had physically sensible short-range walls with strictly positive curvature.

Figure 4 shows how the quality-of-fit parameter \overline{dd} for the global fit varies with the EMO exponent polynomial order $N = N_\beta$ and with the expansion variable parameters q and r_{ref} . As is seen there, the $q = 5$ models (solid points joined by dashed lines) require more β_i expansion parameters to achieve a given quality of fit than do those for $q = 4$ (open filled points joined by solid lines), and do not even approach the $q = 4$ results for the range of $N = N_\beta$ values considered. Because of the close convergence of the $N_\beta = 16$ and 17 models at their common \overline{dd} minima, we choose the $N_\beta = 16$, $q = 4$ EMO function for $r_{\text{ref}} = 3.0$ Å as our recommended potential energy function for the $X^1\Sigma^+$ state of NaH. Figure 4 is a much simpler picture than is the full version of Fig. 2,⁴³ because the EMO potential has only one integer expansion-variable parameter to vary, while the MLR form has two. Moreover, since the distance between their potential asymptotes is determined by the precisely known $\text{Na}(3s \rightarrow 3p)$ atomic excitation energy,⁴⁵ both the value of and the uncertainty in the $A^1\Sigma^+$ well depth are determined by those for the $X^1\Sigma^+$ state, so the analog of the upper panel of Fig. 2 would carry little new information.

The final stage of the analysis consisted of our optimizing the forms of the expansion variables and numbers of expansion parameters defining the BOB functions for the two states. This was a relatively straightforward procedure, since the fact that their effects are quite small relative to those of the potential function parameters means that at worst, only order-of-magnitude trial parameters are required to initiate stable, unique least-squares fits (indeed, trial parameter values of 0.0 usually suffice!). Note, however that the limited vibrational range of the deuteride data does not allow a physically significant determination of distinct well depths for the deuterium isotopologues, so the values of u_0^{H} for the two states are 100% correlated. In particular, either one (or their difference) could serve to define the single experimental observable, the electronic isotope shift. In the analysis reported here, we chose arbitrarily to fix $u_0^{\text{H}}(X) = 0.0$, which means that all of this isotope shift is attributed to the isotopologue-dependence of the A -state well depth. However, this ansatz could readily be changed if additional data were obtained that would allow a reliable independent determination of one of the deuteride well depths. Since the centrifugal BOB corrections for the two states both affect high- J electronic data in similar ways, one might expect these functions to also be fairly highly correlated. However, a trial-and-error procedure quickly led to an optimum model here too.

The parameters defining our recommended analytic potential energy functions for the $X^1\Sigma^+$ and $A^1\Sigma^+$ states of NaH are presented in Table II. Numbers in parentheses are 95% confidence limit uncertainties in the last digits shown, obtained by application of the averaging-over-models procedure described above to models whose \overline{dd} values lie within 1% of the minimum value. After application of the “sequential rounding and refitting” (SRR) scheme of Ref. 46 minimized the numbers of digits required to define the model with no significant loss of accuracy (<0.04% increase in

\overline{dd}), the final fit to 5263 data, and yielded $\overline{dd} = 1.216$. A Fortran subroutine for generating the effective potential energy and centrifugal potential functions for either isotopologue in either state is included with the supplementary material accompanying this manuscript.³³ This supplementary material also includes listings of the eight leading “band constants” (the vibrational energies and first 7 rotational constants) for all vibrational levels of both isotopologues supported by these two potentials, together with sample input data files for the standard vibrational-eigenvalue/Franck-Condon “LEVEL.”⁴⁷

The lower panel of Fig. 5 compares our spectroscopically accurate empirical potential energy functions for the $X^1\Sigma^+$ and $A^1\Sigma^+$ states of NaH with the *ab initio* potentials of Aymar, Deiglmayr, and Dulieu⁹ (points). On the scale of this

TABLE II. Parameters defining the recommended MLR potential energy function for the $X^1\Sigma^+$ state of NaH, the EMO function for the $A^1\Sigma^+$ state, and the associated BOB functions, as determined from the present DPF analysis. The analysis also relied on the $X^1\Sigma^+$ -state dispersion coefficients of Mitroy and Bromley:¹⁶ $C_6 = 3.57502 \times 10^5$, $C_8 = 5.41796 \times 10^6$, and $C_{10} = 1.12920 \times 10^8$, all in units [$\text{cm}^{-1} \text{Å}^m$], the damping function scaling parameter $\rho_{\text{AB}} = 0.69$, and the value of the $\text{Na}(3s^2S \rightarrow 3p^2P_{1/2})$ excitation energy (taken from the NIST www site⁴⁵) that defines the energy of the potential asymptote $\text{VLIM}(A^1\Sigma^+)$. Units of energy and length are cm^{-1} and Å.

	$X^1\Sigma^+$	$A^1\Sigma^+$
VLIM	0.0	16956.17025
T_e	0.0	22712.57025 (12200)
\mathcal{D}_e	15797.4 (43)	10041.0 (43)
r_e	1.887023 (15)	3.1927 (5)
$\{p, q\}/\{q\}$	{6,4}	{4}
r_{ref}	[3.0]	[3.0]
β_0	0.06582161	0.3620684
β_1	-4.1671619	0.131948
β_2	-5.775325	0.155548
β_3	-6.25871	0.08981
β_4	-6.34291	0.41019
β_5	-6.2056	-0.2531
β_6	-5.0814	-2.8482
β_7	-2.205	-0.0395
β_8	-1.678	13.606
β_9	-8.36	2.618
β_{10}	-14.93	-39.17
β_{11}	-11.18	-8.34
β_{12}	-3.1	66.76
β_{13}	...	10.8
β_{14}	...	-61.7
β_{15}	...	-5.
β_{16}	...	24.
$\{p_{\text{ad}}, q_{\text{ad}}\}/\{q\}$	{6,4}	{3,3}
u_0	0.0	-2.26 (124)
u_1	17.36	-8.
u_2	28.2	21.4
u_3	69.2	70.
u_4	-220.	...
u_∞	[0.0]	[0.0]
$\{p_{\text{na}}, q_{\text{na}}\}$	{3,3}	{3,3}
t_0	[0.0]	[0.0]
t_1	0.00005	0.00016
t_2	-0.003	...
t_∞	[0.0]	[0.0]

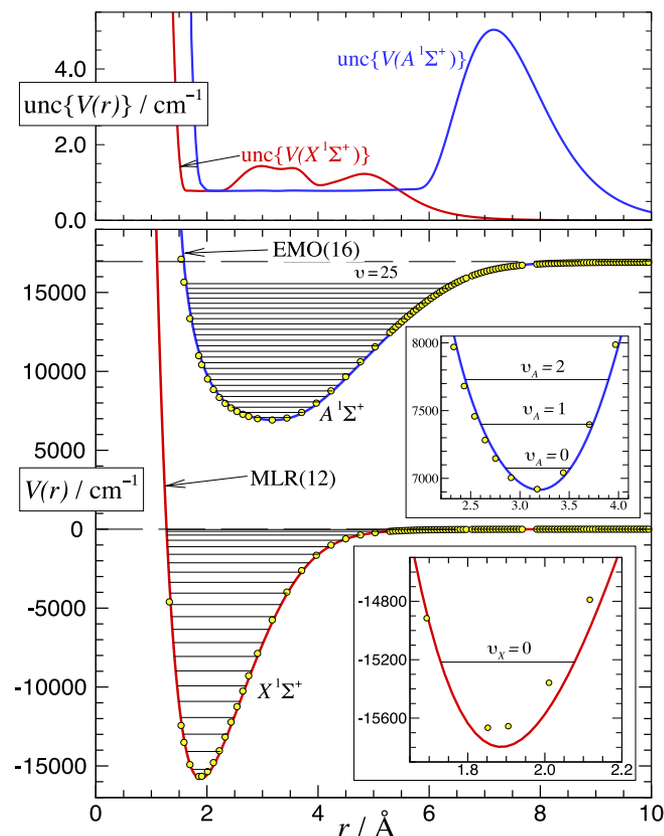


FIG. 5. Lower panel: comparison of the *ab initio* potentials of Aymar *et al.* (round points) with the present empirical potentials (solid red and blue curves). The horizontal line segments indicate the energies of the observed vibrational levels used in the analysis, while the diagram inserts provide additional detail near the potential minima. Upper panel: predicted uncertainties in the fitted potentials due to the uncertainties in the parameters determined by the fit.

figure, the degree of agreement is remarkably good, although the minimum and inner wall of the *ab initio* potential for the *A*-state appear to lie at slightly smaller distances than are dictated by the experimental data. Indeed, the *ab initio* estimates of the equilibrium distances for the *X* and *A* states, 1.87 Å and 3.16 Å,⁹ are both about 1% smaller than our recommended empirical values. The upper panel of Fig. 5 then shows the 95% confidence limit uncertainties in the final potentials, as calculated from the parameter uncertainties and correlation matrix of the fit using standard expressions (see, e.g., Eq. (7) of Ref. 46). As would be expected, the uncertainty in the *A*-state potential goes through a maximum in the extrapolation region between the outer turning point of the highest observed level and the asymptote. This does not occur for the ground state, as the outer end of its data region lies at relatively large *r* where the overall interaction energy is going to zero. As may be expected, the uncertainties in both potentials become quite large beyond the inner ends of their respective data regions which lack the limiting constraint of having to approach an asymptote.

VI. CONCLUSION AND DISCUSSION

A global DPF analysis of all available data for the $A^1\Sigma^+ - X^1\Sigma^+$ systems of NaH and NaD has yielded accurate

analytical potential energy functions and Born-Oppenheimer breakdown functions for these two states that (on average) reproduce all of the data within the experimental uncertainties. Figure 5 shows that these potential functions are in excellent agreement with the best available *ab initio* calculations for these states. It is particularly gratifying to see the close agreement of our fitted *A*-state EMO function with the *ab initio* points in the extrapolation region past the outer turning point for $v_A = 25$, which marks the outer end of the *A*-state data region, as this demonstrates the ability of this functional form to accommodate the abrupt change of potential function character associated with the avoided crossing that takes this state to the $\text{Na}(3p) + \text{H}(1s)$ limit (see Fig. 1) and gives rise to the abrupt cutoff of the $\Delta G_{v+1/2}$ plot seen in Fig. 6. These new potentials, together with new *ab initio* dipole moment and transition moment functions, are now being used to generate comprehensive rotational-vibrational and rotational-vibrational-electronic line lists for the *X* state and the *A*-*X* systems of NaH and NaD.⁴⁸

Our analysis confirms that $v = 21$ (bound by 18.9 cm^{-1}) is indeed the last bound level of the ground state of NaH and predicts that $v = 30$ (bound by 1.26 cm^{-1}) is the highest vibrational level of ground-state NaD. While the uncertainty in \mathcal{D}_e may make the significance of these estimates somewhat uncertain, the scattering lengths implied by our potentials for ground-state NaH and NaD are 0.392 and 7.76 Å, respectively. Similarly, our analysis predicts that the highest vibrational levels of the $A^1\Sigma^+$ states of NaH and NaD are $v = 32$ (bound by 26.8 cm^{-1}) and $v = 45$ (bound by 4.5 cm^{-1}), respectively, and our estimates of the energies and rotational constants of all unobserved levels of both states are included in the supplementary material associated with this paper.³³

Table III compares our present estimate of the potential energy well depth of ground-state NaH with selected previous values. It is noteworthy that our value differs from that of Huang *et al.* by far more than the sum of their uncertainties, although they were based on very similar data sets. We believe that this improvement illustrates the strength of our

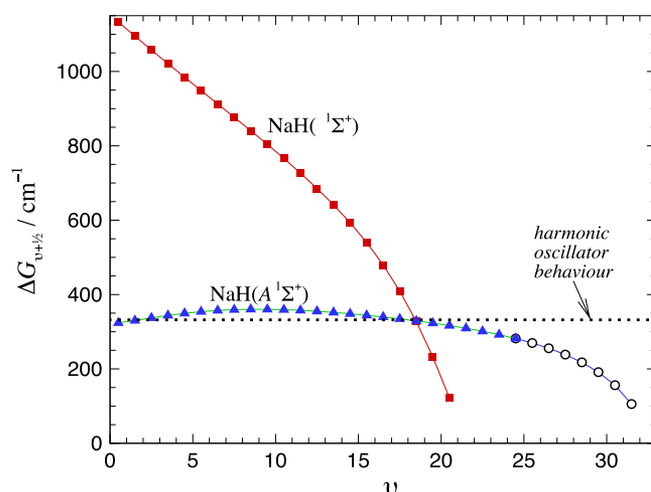


FIG. 6. Birge-Sponer plots for the observed vibrational levels of NaH($X^1\Sigma^+$) (red square points) and of NaH($A^1\Sigma^+$) (blue triangular points) and predicted values for the unobserved higher levels of the latter (open round points).

TABLE III. Comparison of present $\text{NaH}(X^1\Sigma^+)$ well depth (shown in bold font) with some previous values.

Reference	D_e/cm^{-1}	Method
Present work (2013)	15 797.4 ± 4.3	Direct fit of data to analytic potential functions
Huang <i>et al.</i> ³⁰	15 815 ± 5	Polynomial Birge-Sponer extrapolation from $v'' = 21$
Aymar <i>et al.</i> ⁹	15 671.3	<i>Ab initio</i> points: see Fig. 5
Stwalley <i>et al.</i> ³⁹	15 900 ± 100	Add <i>ab initio</i> tail to vibrational energy at $v'' = 19$
Nedelec and Giroud ⁴⁹	15 785 ± 20	Polynomial Birge-Sponer extrapolation from $v = 19$
Giroud and Nedelec ⁵⁰	16 300 ± 500	Polynomial Birge-Sponer extrapolation from $v = 16$
Pankhurst ¹⁹	17 410	Linear Birge-Sponer extrapolation from $v'' = 8$
Hori ⁸	18 100	Polynomial Birge-Sponer extrapolation from $v'' = 20$

DPF procedure. As illustrated by the upper panel of figure 2, our final estimate of the uncertainty in D_e is almost an order of magnitude larger than the uncertainty yielded by the final fit to our particular recommended model. This is a common effect of taking model-dependence into account. However, this averaging over models responsible for less than half of our estimated uncertainties in the physically significant A -state parameters T_e , r_e , and u_0 .

A widely noted fact about the $A^1\Sigma^+$ states of all alkali hydrides is that their vibrational spacings and B_v values initially increase with v , a property that reflects the broader-than-quadratic shape of their potential minima, which in turn is associated with the influence of the transient ion-pair character of their attractive outer walls (see Fig. 1) and the associated (distant) avoided crossing with the ground state. However, Figs. 5 and 6 illustrate another interesting property, which is that over much of its well, the vibrational spacings of the $A^1\Sigma^+$ state of NaH are approximately constant, even though Fig. 5 shows that its shape is strongly anharmonic. This provides an unusual physical illustration of the separate roles of vibrational and rotational data in determining a potential energy function, which are so clearly delineated by the two equations of the semiclassical RKR inversion procedure:¹¹ the vibrational spacings only depend on, and hence determine, how the width of the potential well increases with energy, and not its overall shape, so that over a substantial energy range a pure harmonic oscillator and a strongly anharmonic potential energy function can have approximately the same vibrational spectrum.

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¹F. Crawford and J. T. Jorgenson, *Phys. Rev.* **47**, 358 (1935).

²R. S. Milliken, *Phys. Rev.* **36**, 1017 (1936).

³W. C. Stwalley and W. T. Zemke, *J. Phys. Chem. Ref. Data* **22**, 87 (1993).

⁴J. A. Coxon and C. S. Dickinson, *J. Chem. Phys.* **121**, 9378 (2004).

⁵J. Tennyson, in 68th International Symposium on Molecular Spectroscopy at the Ohio State University, Columbus, Ohio, 2013, Paper TB05.

⁶A. S. Rajpurohit, C. Reyle, F. Allard, D. Homeier, M. Schultheis, M. S. Bessell, and A. C. Robin, *Astron. Astrophys.* **556**, A15 (2013).

⁷T. Hori, *Z. Phys.* **62**, 352 (1930).

⁸T. Hori, *Z. Phys.* **71**, 478 (1931).

⁹M. Aymar, J. Deiglmayr, and O. Dulieu, *Can. J. Phys.* **87**, 543 (2009).

¹⁰G. Herzberg, *Spectra of Diatomic Molecules* (Van Nostrand, New York, 1950).

¹¹R. Rydberg, *Z. Phys.* **73**, 376 (1931); O. Klein, *Z. Phys.* **76**, 226 (1932); R. Rydberg, *Z. Phys.* **80**, 514 (1933); A. L. G. Rees, *Proc. Phys. Soc.* **59**, 998 (1947).

¹²R. J. Le Roy, RKR1 2.0: A Computer Program Implementing the First-Order RKR Method for Determining Diatomic Molecule Potential Energy Curves, University of Waterloo Chemical Physics Research Report CP-657, 2003; see <http://leroy.uwaterloo.ca/programs/>.

¹³R. J. Le Roy, “Determining equilibrium structures and potential energy functions for diatomic molecules,” of *Equilibrium Structures of Molecules* edited by J. Demaison and A. G. Csaszar, (Taylor & Francis, London, 2011), Chap. 6, pp. 159-203.

¹⁴R. J. Le Roy, J. Seto, and Y. Huang, DPotFit 2.0: A Computer Program for fitting Diatomic Molecule Spectra to Potential Energy Functions, University of Waterloo Chemical Physics Research Report CP-667, 2013; see <http://leroy.uwaterloo.ca/programs/>.

¹⁵R. D. E. Henderson, A. Shayesteh, J. Tao, C. C. Haugen, P. F. Bernath, and R. J. Le Roy, *J. Phys. Chem. A* **117**, 13373 (2013).

¹⁶J. Mitroy and M. W. J. Bromley, *Phys. Rev. A* **68**, 052714 (2003); Erratum, *A71*, 019902(E) (2005).

¹⁷E. Olsson, *Z. Phys.* **93**, 206 (1934).

¹⁸F. P. Pesl, S. Lutz, and K. Bergmann, *Eur. Phys. J. D* **10**, 247 (2000).

¹⁹R. C. Pankhurst, *Proc. Phys. Soc., Sect. A* **62**, 191 (1949).

²⁰F. B. Orth, W. C. Stwalley, S. C. Yang, and Y. K. Hsieh, *J. Mol. Spectrosc.* **79**, 324 (1980).

²¹K. V. L. N. Sastry, E. Herbst, and F. C. de Lucia, *J. Chem. Phys.* **75**, 4753 (1981).

²²K. R. Leopold, L. R. Zink, K. M. Evenson, and D. A. Jennings, *J. Mol. Spectrosc.* **122**, 150 (1987).

²³U. Magg and H. Jones, *Chem. Phys. Lett.* **146**, 415 (1988).

²⁴A. G. Maki and W. B. Olson, *J. Chem. Phys.* **90**, 6887 (1989).

²⁵M. Rafi, N. Ali, K. Ahmad, I. A. Khan, M. A. Baig, and Z. Iqbal, *J. Phys. B: At., Mol. Opt. Phys.* **26**, L129 (1993).

²⁶M. A. Baig, private communication (2013).

²⁷S. Lochbrunner, M. Motzkus, G. Pichler, K. L. Kompa, and P. Hering, *Z. Phys. D: At., Mol. Clusters* **38**, 35 (1996).

²⁸J. Pesl, Private Communication (2012); we are grateful to Dr. Pesl for providing us with these unpublished data of Lochbrunner *et al.* (Ref. 27).

²⁹J. T. Bahns, C. C. Tsai, B. Ji, J. T. Kim, G. Zhao, W. C. Stwalley, J. C. Bloch, and R. W. Field, *J. Mol. Spectrosc.* **186**, 222 (1997).

³⁰H.-Y. Huang, T.-L. Lu, T.-J. Whang, Y.-Y. Chang, and C.-C. Tsai, *J. Chem. Phys.* **133**, 044301 (2010)

³¹J. L. Dunham, *Phys. Rev.* **41**, 713-720 (1932); **41**, 721-731 (1932).

³²Our initial analyses found that in Huang *et al.*'s original data set,³⁰ the transition energies for the (10,16) band appeared to be systematically displaced by approximately 10 cm^{-1} . Dr. Huang has kindly provided us with the correct values for these data (T.-J. Huang, private communication (2012)), and they were used in the present analysis.

- ³³See supplementary material at <http://dx.doi.org/10.1063/1.4906086> for ASCII files containing: (i) a list of all of the data used in the present work, (ii) a list of [calc.-obs] values for our final recommended model, (iii) tabulations of the vibrational energies and rotational constants all bound levels of the $X^1\Sigma^+$ and $X^1\Sigma^+$ states of NaH and NaD that were generated from our final recommended potential energy functions, (iv) a digital listing of the complete fitted parameters set from Table II, together with a band-by-band summary of the final set of RMS discrepancies, (v) sample data files for use in performing bound-state/Franck-Condon calculations using Program LEVEL,⁴⁷ (vi) a FORTRAN subroutine for generating the recommended MLR(X) and DELR(A) potential energy functions, and (vii) the *full* version of Fig. 2 including the results for *four* families of $\{p, q\}$ cases.
- ³⁴J. K. G. Watson, *J. Mol. Spectrosc.* **80**, 411 (1980).
- ³⁵J. K. G. Watson, *J. Mol. Spectrosc.* **223**, 39 (2004).
- ³⁶R. J. Le Roy, *J. Mol. Spectrosc.* **194**, 189 (1999).
- ³⁷N. Dattani and R. J. Le Roy, *J. Mol. Spectrosc.* **268**, 199 (2011).
- ³⁸R. J. Le Roy, betaFIT 2.1: A Computer Program to Fit Potential Function Points to Selected Analytic Functions, University of Waterloo Chemical Physics Research Report CP-666, 2013; see <http://leroy.uwaterloo.ca/programs/>.
- ³⁹W. C. Stwalley, W. T. Zemke, and S. C. Yang, *J. Phys. Chem. Ref. Data* **20**, 153 (1991).
- ⁴⁰R. J. Le Roy, C. C. Haugen, J. Tao, and H. Li, *Mol. Phys.* **109**, 435 (2011).
- ⁴¹C. Douketis, G. Scoles, S. Marchetti, M. Zen, and A. J. Thakkar, *J. Chem. Phys.* **76**, 3057 (1982).
- ⁴²R. J. Le Roy and Y. Huang, *J. Mol. Struct.: THEOCHEM* **591**, 175 (2002).
- ⁴³A somewhat “busier” version of the present Fig. 2 which also includes the results of fits with $\{p, q\} = \{5, 3\}$ and $\{5, 4\}$, is included with the supplementary information associated with this paper.³³
- ⁴⁴R. J. Le Roy, *J. Chem. Phys.* **101**, 10217 (1994).
- ⁴⁵See http://physics.nist.gov/PhysRefData/ASD/levels_form.html for a table of Na atom level energies.
- ⁴⁶R. J. Le Roy, *J. Mol. Spectrosc.* **191**, 223 (1998).
- ⁴⁷R. J. Le Roy, Level 8.2: A Computer Program for Solving the Radial Schrödinger Equation for Bound and Quasibound Levels, University of Waterloo Chemical Physics Research Report CP-663, 2014; see <http://leroy.uwaterloo.ca/programs/>.
- ⁴⁸T. Rivlin, L. Lodi, S. N. Yurchenko, J. Tennyson, and R. J. Le Roy, “ExoMol molecular line lists: The spectrum of sodium hydride” (unpublished).
- ⁴⁹O. Nedelec and M. Giroud, *J. Chem. Phys.* **79**, 2121 (1983).
- ⁵⁰M. Giroud and O. Nedelec, *J. Chem. Phys.* **73**, 4151 (1980).