

Modeling of diatomic predissociation resonances within the Optimizer project

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ABSTRACT

The simulation of the predissociation spectrum of the $\text{Na}_2 2^3\Pi_g \sim 3^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) transition, observed earlier with the Perturbation Facilitated Optical–Optical Double Resonance (PFOODR) experimental method, is done within the Optimizer project with our Split-operator computational package using Padé approximations for all functions involved in the model. This simulation reproduces the experimental spectrum satisfactory well. The parameters of the model functions involved in the simulations are determined and reported. A brief description of our computational methods and programs is presented.

Introduction

We have developed the Optimizer project [1,2] in various versions over time during the last three decades or more. Although we believe that its main application is the analysis of molecular spectra it can also be used more broadly including problems that require optimization in various fields of science such as engineering and technology, for example. Most of our scientific results (e.g., [3–19] and references therein) were possible by using the Optimizer project.

The Optimizer project facilitates construction and optimization of complicated mathematical models of a versatile nature. The optimization process relies on using the familiar Levenberg–Marquardt algorithm [20,21] based on Singular Value Decomposition (SVD) [22] of the design matrix, which when needed, can be switched to any other one. It supports various methods of problem regularization, robust estimators and tools to parallelize the computation, for example.

The model construction can be done block-by-block from relatively simple “standard” modeling programs (blocks) with a convenient

universal mechanism of transferring data from block to block. The program codes are subjected to some conventional regulations enabling them to be easily embedded into the entire Optimizer model. A library of such modeling programs is available for general approximation problems in addition to computational molecular spectroscopy along with the main Optimizer package, which is continually improved and extended.

All program components are written in MATLAB algorithmic language and can be executed using the MATLAB [23] or OCTAVE [24] environments. They are available with detailed instruction manuals and test samples for a free download [2] on the Internet.

The predissociation spectrum of the $\text{Na}_2 2^3\Pi_g \sim 3^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) transition was observed and reported by Liu et al. in [25]. The experimental technique was based on the Perturbation Facilitated Optical–Optical Double Resonance (PFOODR) technique [3,7,26–36] to probe excited triplet states and detect fluorescence to the lower lying electronic states. The general scheme of the PFOODR experiment

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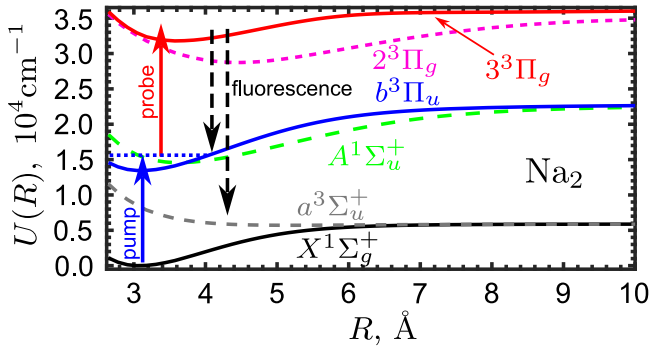


Fig. 1. (Color online) The scheme of the PFOODR experiment [25]. The pump laser excites directly the intermediate “window” level, whose approximate position is shown with the horizontal dotted line, and the probe laser excites the states of the mixed $2^3\Pi_g \sim 3^3\Pi_g$ complex above the dissociation limit of $2^3\Pi_g$ and below the one of $3^3\Pi_g$.

Ref. [25] is shown in Fig. 1 with the *ab initio* adiabatic potential energy functions of Ref. [37].

The spectrum in Fig. 2 is characterized by broad fluorescence spectral features caused by the non-adiabatic interaction between the $3^3\Pi_g$ and $2^3\Pi_g$ states as well as by additional narrower spectral lines attributed to the participation of the $4^3\Sigma_g^+$ and 2^3A_g states. Predissociation is an example of quantum resonances, which are among the most important and intriguing phenomena in quantum physics [38,39]. During predissociation a molecule in a bound rovibrational level below the dissociation limit of its electronic state can dissociate into atoms by coupling to a continuum level that belongs to a lower lying electronic state above its dissociation limit. Inverse predissociation can also play a role in the formation of molecules in the interstellar molecular clouds, for example (see Refs. [40,41]).

A preliminary simulation (with a participation of some of us) of the predissociation spectrum of the Na_2 $2^3\Pi_g \sim 3^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) transition by using the multichannel Split-Operator method [42–48] was shown at the conference [49]. A few years later Narevicius et al. [50] reported a simulation of this spectrum using the complex scaling method [51] with *ab initio* potential functions [37]. The spectrum was not reproduced quantitatively by shifting the *ab initio* potentials of the $2^3\Pi_g$ and $3^3\Pi_g$ states relative to each other to optimize the widths of the resonance peaks in the simulation compared to the experiment. To the best of our knowledge, no other attempts to simulate or analyze this spectrum have been performed since then.

In the present work we have improved our multichannel Split-operator computer program and adapted it to meet the requirements of the Optimizer project. We checked its ability to simulate predissociation resonances in the predissociation spectrum being discussed in this work. For this purpose, we constructed the model of the two-channel coupling of the $3^3\Pi_g$ and $2^3\Pi_g$ states using Padé approximations [52–65] for all the functions involved. We optimized the parameters of the model to get the best reproduction of the experimental spectrum.

In the Appendix we present brief description of our computational methods and programs.

Theory and algorithms

In the diabatic representation [66], the two-channel potential is a matrix

$$U_{\text{diab}}(R) = \begin{bmatrix} V_1(R) & W(R) \\ W(R) & V_2(R) \end{bmatrix}, \quad (1)$$

where R is the interatomic distance, $V_1(R)$ and $V_2(R)$ are the diabatic potential functions of the molecular electronic states, $W(R)$ is the interaction matrix element. The eigenvalues of this matrix are the

adiabatic potential functions $U_1(R)$ and $U_2(R)$ (differ by the “+” and “-” signs below):

$$U_{1,2}(R) = \frac{1}{2} \left[V_1(R) + V_2(R) \pm \sqrt{(V_1(R) - V_2(R))^2 + 4W^2(R)} \right], \quad (2)$$

$$V_{1,2}(R) = \frac{1}{2} \left[U_1(R) + U_2(R) \pm \epsilon \sqrt{(U_1(R) - U_2(R))^2 - 4W^2(R)} \right] \quad (3)$$

(ϵ is a conventional sign factor, which should be inverted at cross points), while the normalized eigenvectors being arranged column-wise comprise the transformation matrix from the adiabatic to the diabatic representation:

$$F_{a \rightarrow d}(R) = \begin{bmatrix} f_1^{(1)}(R) & f_2^{(1)}(R) \\ f_1^{(2)}(R) & f_2^{(2)}(R) \end{bmatrix} \quad (4)$$

with

$$\begin{cases} f_{1,2}^{(1)}(R) = t_{1,2}(R)f_{1,2}^{(2)}(R), \\ f_{1,2}^{(2)}(R) = (t_{1,2}^2 + 1)^{-1/2}, \\ t_{1,2}(R) = [\delta V(R) \mp \sqrt{\delta V^2(R) + 1}]^{-1}, \\ \delta V(R) = \frac{V_2(R) - V_1(R)}{2W(R)}. \end{cases} \quad (5)$$

As is well known [67], the diabatic potential functions can cross each other, while the adiabatic ones cannot and instead give rise to avoided crossing.

There are number of ambiguities in the above set of equations that need to be addressed before they can be used for specific calculations. First, the columns in Eq. (4) can be swapped, resulting in a switch of the order of the adiabatic potential functions. We choose to assign the first column of the matrix to the state with lower adiabatic potential. Second, the sign of every column of (4) is undefined, reflecting the well known arbitrariness of eigenvector phases and resulting in the change of sign of the function being transformed with the help of this matrix. The signs of the interaction function $W(R)$ relative to the signs of the transition moment functions $M_1(R)$ and $M_2(R)$ in either diabatic or adiabatic representation reflects the choice of the signs of the electronic eigenfunctions in the same representation, while a switch of a sign of a column in (4) implies a switch of the sign of a (transformed) electronic eigenfunction in one of these representations. In principle, with any consistent choice of signs at every point R independently the theory remains correct, however, it is evident that smooth functions are easier to interpret and compute. We adopted the rule that the signs of the diagonal elements $f_1^{(1)}(R)$ and $f_2^{(2)}(R)$ to coincide with the sign of $W(R)$; notice, that our final $W(R)$ occurred negative in the entire range, causing no problems with the switch of all relevant signs at a point of $W(R) = 0$.

Our computations were performed in the following way. First, we linearly transformed (warped) the *ab initio* adiabatic potentials [37] of the $3^3\Pi_g$ and $2^3\Pi_g$ states placing the long-range limit to the approximate atomic $3s + 3d$ and $3s + 4p$ energies [68] (neglecting the fine and hyperfine splitting), and the bottom at the approximate experimental energy of the potential minimum [69]. We deperturbed these potentials using Eq. (3) with the implied cross point at $R_c \approx 2.8 \text{ \AA}$ and the constant value for $W(R) = -153 \text{ cm}^{-1}$ (following the rough model of [49]). Then, we fitted the diabatic potential energy curves to the Padé functions and used them as the zeroth order approximations. In contrast, the zeroth order approximations for the interaction $W(R)$ and the transition moment functions $M_1(R)$ and $M_2(R)$ were just constants.

To generate the initial wavepacket, we computed the wavefunction of the Na_2 $b^3\Pi_u$ ($v = 14, J = 14$) level with our programs of the

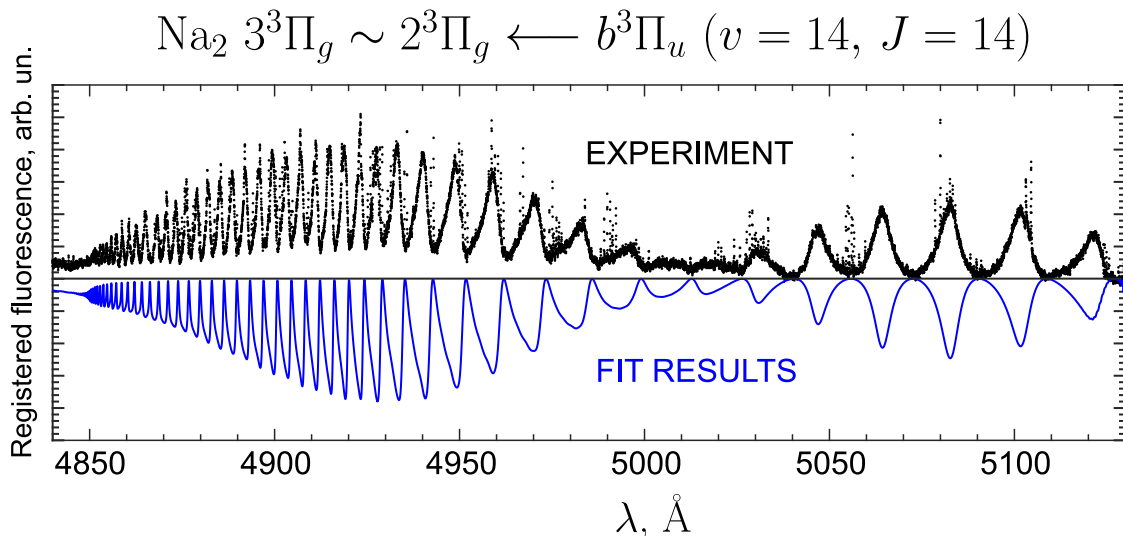


Fig. 2. (Color online) The experimental [25] and fitted predissociation spectra of the $\text{Na}_2 3^3\Pi_g \sim 2^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) transition.

Numerov method [70], already included into the Optimizer project [2]. For the potential function of the $b^3\Pi_u$ state we took the one reported in [71]. In the course of the computations we eliminated a misprint in the table II of [71]: the decimal order of the parameter a_5 of the $b^3\Pi_u$ state must be “4” in place of the reported value “5”; otherwise it was impossible to reproduce the ro-vibrational term values with this potential.

After that, the following iteratively repeated procedure was applied:

1. The diabatic potential energy functions of both states, the interaction function, and the transition moment functions were computed from their current Padé parameters at the required spatial grid points.
2. The two-channel diabatic initial wavepacket was formed by multiplying the $b^3\Pi_u$ ($v = 14, J = 14$) wavefunction by the transition moment functions of the corresponding channels.
3. The adiabatic potential functions were computed from the diabatic ones using Eq. (2), and the set of transformation matrices was computed using Eqs. (4) and (5).
4. Next, the two-channel evolution (the time autocorrelator) of the wavepacket in the field of the $3^3\Pi_g \sim 2^3\Pi_g$ electronic state complex was computed with our Split-operator programs.
5. The predissociation spectrum was computed using a Fourier transform of the time autocorrelator and compared with the experimental one.

Besides that, we added to the data being fitted and simulated the *ab initio* calculated [37] adiabatic potential functions and the adiabatic transition moment functions, although with rather smaller weights. Then, model parameters were corrected and the procedure was repeated until a satisfactory reproduction of the experimental spectrum was achieved.

In most computations, the spatial grid was set between 2 Å and 26.564 Å with step 0.012 Å, and the time grid spanned the 0 to 30 fs range with step of 0.001 fs (permitting computation of the time autocorrelator in the range from -60 fs to 60 fs [46]). In the long-range region, the wavepacket was damped by the cosine-type absorbing function described in our work [72] with the recommended parameters (in the designations of [72]) $E_0 = 100 \text{ cm}^{-1}$, $k = 3$, $l = 4$, $\alpha = 0.266$, $l_+ = 1$, $\gamma = 1$, with its right boundary located at the very end of the spatial grid and the corresponding left boundary at $R_b = 22.76 \text{ \AA}$. When the main computation was finished, we tried altered parameters: reduced spatial and time steps twice, extended the time grid to 50 fs, set the reference energy of the absorbing function to $E_0 = 30 \text{ cm}^{-1}$,

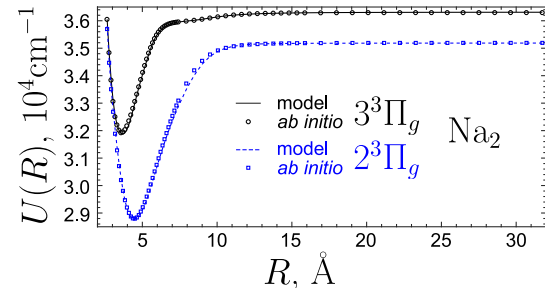


Fig. 3. (Color online) The *ab initio* and the final model adiabatic potential energy curves of the $\text{Na}_2 3^3\Pi_g$ and $2^3\Pi_g$ states. The origin of the energy scale is at the ground state $\text{Na}_2 X^1\Sigma^+$.

extended the absorbing region to $l = 6$ with $\alpha = 0.306$ [72],—and observed no noticeable change in the computed spectrum.

Results

The final simulation of the $\text{Na}_2 3^3\Pi_g \sim 2^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) predissociation spectrum is compared with the experimental one in Fig. 2.

The final Padé approximant parameters of all diabatic functions that were used to reproduce the results of the present paper, are reported in Tables 1 and 2. We would like to emphasize that the simulated spectrum in Fig. 2 was computed with the parameters in Tables 1 and 2 exactly, *i.e.* being already rounded.

The general view of the final adiabatic $3^3\Pi_g$ and $2^3\Pi_g$ state potential functions along with the *ab initio* ones is shown in Fig. 3, and a magnified view of the region of the (avoided) crossings is shown in Fig. 4.

The function of the interaction matrix element is shown in Fig. 5

The transition moment functions of the $\text{Na}_2 3^3\Pi_g \leftarrow b^3\Pi_u$ and $2^3\Pi_g \leftarrow b^3\Pi_u$ optical transitions in both diabatic and adiabatic representations are shown in Fig. 6 along with the *ab initio* [37] ones.

Discussion

In our opinion, the reproduction of the experimental spectrum with our final simulation in Fig. 2 is exceptionally good, with the quality

Table 1

Parameters of the Padé functions approximating the $\text{Na}_2 3^3 \Pi_g$ and $2^3 \Pi_g$ diabatic potentials involved in the model. All values are measured in Å and cm^{-1} . The origin of the energy scale is at the bottom of the ground state $\text{Na}_2 X^1 \Sigma_g^+$. Uncertainties of the fitted model parameters are of the order of magnitude of the last decimal digit reported.

$3^3 \Pi_g$		$2^3 \Pi_g$	
Parameter	Value	Parameter	Value
x_0	3.686	b_0	1
C	36 311.4	b_1	0
		b_2	2840
a_0	-4467	b_3	1285
a_1	0	b_4	220
a_2	-1.2685e+07	b_5	330
a_3	-5.62e+06	b_6	130
a_4	5.25e+06	b_7	-300
a_5	400 000	b_8	-27
a_6	-90 000	b_9	99
a_7	-8.167e+05	b_{10}	29.02
a_8	1.235e+05	b_{11}	-3.5
a_9	15 000	b_{12}	-2.21
a_{10}	-3450	b_{13}	0.332
a_{11}	-1200	b_{14}	0.021

Table 2

Parameters of the Padé functions approximating the $\text{Na}_2 2^3 \Pi_g \sim 3^3 \Pi_g$ interaction matrix element W and $3^3 \Pi_g \leftarrow b^3 \Pi_u$, $2^3 \Pi_g \leftarrow b^3 \Pi_u$ transition moments M involved in the model. All values are measured in Å, cm^{-1} , and D. Uncertainties of the fitted model parameters are of the order of magnitude of the last decimal digit reported.

$2^3 \Pi_g \sim 3^3 \Pi_g, W$		$3^3 \Pi_g \leftarrow b^3 \Pi_u, M$		$2^3 \Pi_g \leftarrow b^3 \Pi_u, M$	
Parameter	Value	Parameter	Value	Parameter	Value
x_0	2.906	x_0	5	x_0	5
C	0	C	0	C	5
a_0	-140	a_0	0.48	a_0	-16.3
a_1	-530	a_1	-0.55	a_1	-9
a_2	2.6	a_2	-0.4	a_2	-2.16
b_0	1	b_0	1	b_0	1
b_1	-0.4	b_1	0.9	b_1	0.49
b_2	0.2	b_2	0.2143	b_2	0.2

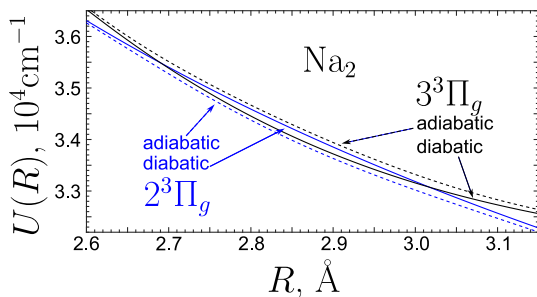


Fig. 4. (Color online) The final model diabatic (solid lines) and adiabatic (dashed lines) potential energy curves of the $\text{Na}_2 3^3 \Pi_g$ and $2^3 \Pi_g$ states in the region of the (avoided) crossings. The origin of the energy scale is at the bottom of the ground state $\text{Na}_2 X^1 \Sigma_g^+$.

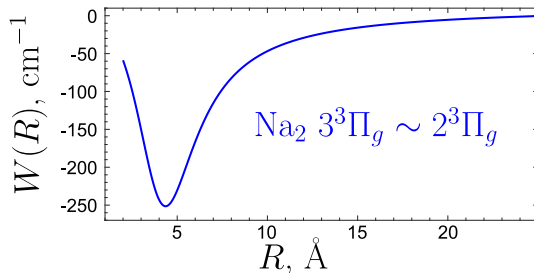


Fig. 5. (Color online) The final model interaction matrix element of the $\text{Na}_2 3^3 \Pi_g \sim 2^3 \Pi_g$ state complex.

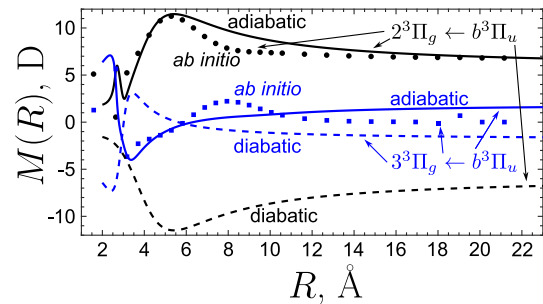


Fig. 6. (Color online) The moment functions of the $\text{Na}_2 3^3 \Pi_g \leftarrow b^3 \Pi_u$ and $2^3 \Pi_g \leftarrow b^3 \Pi_u$ optical transitions: dots—*ab initio*, solid lines—model adiabatic, dashed lines—model diabatic.

close to the accuracy of the experimental measurements. The wavelengths, widths, and intensities of the resonance peaks are accurately reproduced. The remaining discrepancies in the forms of some of the line profiles can be attributed to both experimental inaccuracies and restrictions of our model, including ignoring the triplet character of the states, spin-orbit mixing of the initial $b^3 \Pi_u$ state with the $A^1 \Sigma_u^+$ state, and perturbations from other near-lying states.

The most important factor determining the quality of this reproduction is the outer branch of the $3^3 \Pi_g$ state potential above the $3s + 3d$ (lower) dissociation limit, which can be considered the most reliable quantitative result of our investigation. Other parameters are less influential but necessary for the model.

It is interesting and rather unexpected that our final potential functions have two crossing points at about 2.69 Å and 3.01 Å—see Fig. 4. The previous simulations [49,50] implied the existence of only one crossing point. It was noted [25,50] that the position of the crossing point influences the widths of the resonance peaks. Probably, ignoring the possible existence of an additional crossing point can explain the difficulty in the simulation of the spectral line widths in [50].

Our final transition moment functions in Fig. 6 are relatively close to the *ab initio* ones, however, we must recognize that the deviations remain significant, especially in the short-range region, which is the most important in the formation of the initial wavepacket.

The number of parameters of the $3^3 \Pi_g$ state potential in Table 1 exceeds the analogous number of parameters of the $2^3 \Pi_g$ state potential. In the beginning we tried to describe it with the same number of parameters as $2^3 \Pi_g$ but were unsuccessful. Fig. 7 provides the explanation. The initial version of the function (smaller set of parameters) was unable to reproduce the bend at around 8 Å, which is clearly seen in the *ab initio* potential energy curve (Fig. 7(a)), causing significant shifts

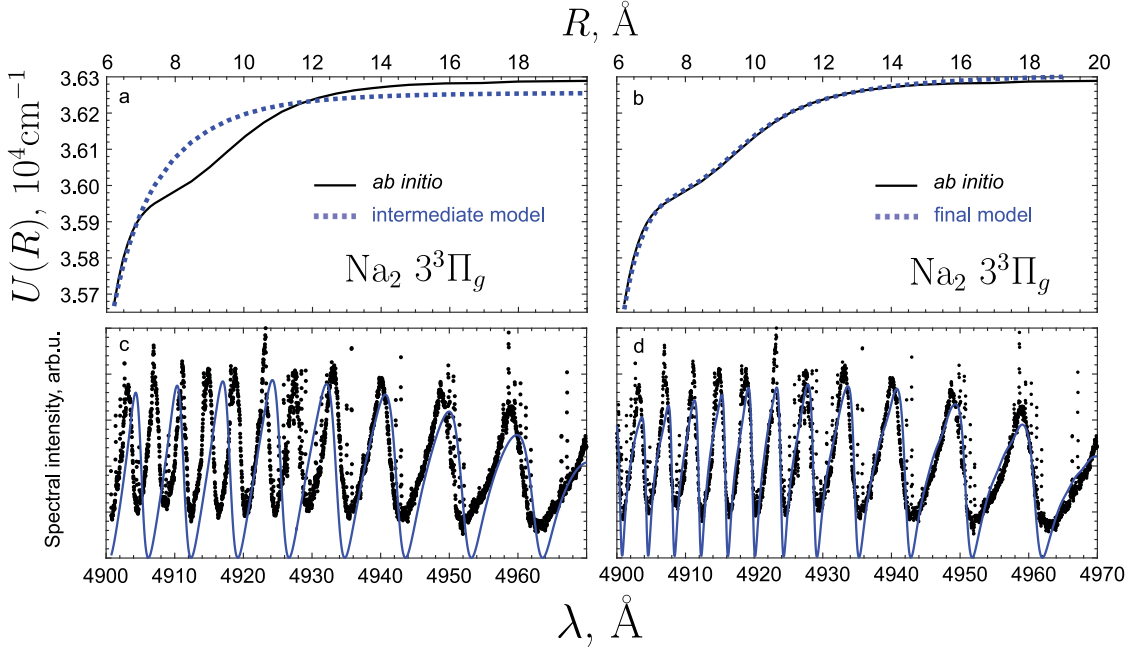


Fig. 7. (Color online) Influence of the $\text{Na}_2 3^3\Pi_g$ potential function bends on the predissociation spectrum: (a) adiabatic *ab initio* and intermediate model potentials; (b) adiabatic *ab initio* and final model potentials; (c) experimental (dots) and intermediate model (line) spectra; (d) experimental [25] (dots) and final model (line) spectra.

of the computed peaks in the range of wavelengths shorter than 4935 Å (Fig. 7(c)). The final version of the function (bigger set of parameters) reproduced this bend (Fig. 7(b)) and improved the positions of the computed peaks (Fig. 7(d)). On the one hand, this confirms once more the importance of even subtle features of the $3^3\Pi_g$ state potential in our model, and on the other hand, this confirms the quality of the long-range *ab initio* potentials [37], exhibiting those subtle features.

We also explored the ability of the Fourier grid method (see [73–76]) in our realization of it for the Optimizer project (see the brief description in [72] with more detail given in the manual in [2]) to predict properties of the predissociation resonances in terms of the complex energies. For this purpose, we took our final potential energy functions of the $3^3\Pi_g$ and $2^3\Pi_g$ states and the interaction function $W(R)$, and embedded the optical potential with the same parameters as of the absorbing function used in the main computation above. Embedding an optical potential makes the Fourier-grid Hamiltonian complex-type with complex-type eigenenergies. The inverse of the imaginary part of this eigenenergy is proportional to the lifetime of the decaying state, located at the energy equal to the real part of it. In Fig. 8, circles connected with the solid line (red in the online color version) are those “lifetimes” versus the real-type energies of the states.

Although some weak peaks are observed near the positions of resonances, they are not much stronger than the neighboring “non-resonant” peaks. Besides, some artifacts (deep dips) are observed, which can confuse the interpretation of the results. We suppose that the long lifetimes of the non-resonant states are caused by a long distance of the region before the absorbing function, so that the non-resonant parts of the wavepacket indeed lives there for a relatively long time. To check this, we shifted the left boundary of the absorbing region from $R_b = 22.76 \text{ \AA}$ to $R_b = 10 \text{ \AA}$ and repeated the computation. The result is shown in Fig. 8 as diamonds connected by the dashed line (blue in the online color version). This indeed made the resonance peaks much more prominent. This allowed us to separate the most intense peaks (biggest lifetimes) and plot them in Fig. 9 on the same scale as the experimental spectrum. Although some correspondence between the observed and calculated resonances takes place, the positions of the computed ones are noticeably shifted in the short-wavelength region.

For a comparison, we computed the unperturbed eigenenergies of the $3^3\Pi_g$ state in the diabatic representation with the Numerov method, and showed them in Figs. 8 and 9 as squares located at the abscissa. It is seen, that the positions of the model complex-type-energy resonances are closer to the experimental peaks in the long-wavelength region but deviate in the short-wavelength region, while the unperturbed eigenenergies are closer to the experimental peaks in the short-wavelength region, but deviate in the long-wavelength region. This can be explained by the fact that at the higher energies the perturbation of the $3^3\Pi_g$ states becomes weaker (in part, that is reflected by the narrower peaks), while the influence of the absorbing optical potential distorts more strongly the Fourier-grid Hamiltonian.

Overall, the approach based on the complex-type eigenenergies of the Fourier-grid Hamiltonian with the embedded optical potential seems not to be very reliable even if it can show some characteristics of the resonances. Its results strongly depend on the parameters of the optical potential and other technical details. We expect that more sophisticated methods that use the complex-type eigenenergies, such as the method of the complex scaling [50,51], are able to give much better results. However, these methods are not included in the Optimizer project yet and are currently outside of the scope of our research.

The only works investigating the lower-lying bound levels of the sodium dimer states under consideration, which we are aware of, are Refs. [28,69,77,78]. However, we have not found estimates of the parameters of the interaction between the $\text{Na}_2 2^3\Pi_g$ and $3^3\Pi_g$ states in them, which could be compared to our results.

Conclusions

We have demonstrated the efficiency of the multichannel Split-operator method programs, embedded into the Optimizer project for description of resonance phenomena, such as molecular predissociation spectra. The experimental $\text{Na}_2 2^3\Pi_g \sim 3^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) spectrum was simulated with an accuracy close to the experimental results. We also demonstrated the ability of the Padé approximants to describe all important features of the potential energy and other functions involved in the model, including subtle bends. The fitted parameters that enabled the reproduction of the results of the present

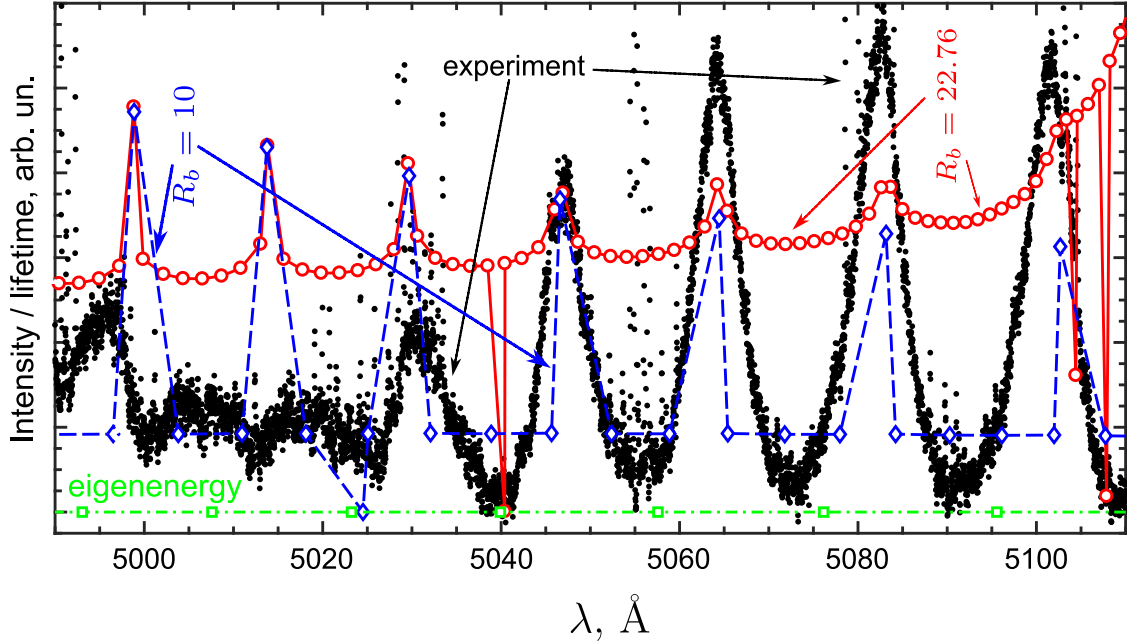


Fig. 8. The experimental [25] $\text{Na}_2 3^3\Pi_g \sim 2^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) predissociation spectrum (dots), lifetimes estimated with the Fourier grid method with the optical potential starting at $R_b = 22.76 \text{ \AA}$ (circles connected with a solid line) and $R_b = 10 \text{ \AA}$ (diamonds connected with a dashed line), and the eigenenergies of the $\text{Na}_2 3^3\Pi_g$ ($J = 14$) state (squares at the abscissa). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

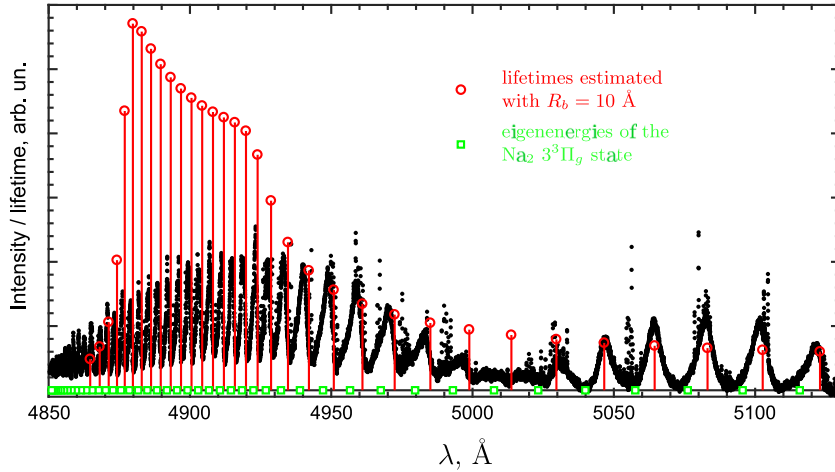


Fig. 9. (Color online) The experimental [25] $\text{Na}_2 3^3\Pi_g \sim 2^3\Pi_g \leftarrow b^3\Pi_u$ ($v = 14, J = 14$) predissociation spectrum (dots), lifetimes estimated with the Fourier grid method with the optical potential starting at $R_b = 10 \text{ \AA}$ (circles with stems), and the eigenenergies of the $\text{Na}_2 3^3\Pi_g$ ($J = 14$) state (squares at the abscissa).

paper, are reported. We also showed that the straightforward computation of the complex-type eigenenergies of the Fourier-grid Hamiltonian with the embedded optical potential can give results in an approximate description of the resonances but is not very reliable quantitatively. A brief introduction of the computer programs used in the present research and embedded into the Optimizer project is presented in the Appendix.

CRediT authorship contribution statement

Vladimir B. Sovkov: Writing – original draft, Software, Methodology, Investigation, Conceptualization. **Jizhou Wu:** Writing – review & editing, Investigation. **Jie Ma:** Writing – review & editing, Investigation. **Feng Xie:** Writing – review & editing, Investigation. **A.M. Lyrya:** Writing – review & editing, Investigation. **Ergin Ahmed:** Writing – review & editing, Investigation.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

The data that support the findings of this study are openly available in the repository Source Forge at <https://sourceforge.net/projects/optimizer-sovkov/>, Ref. [2]. They include the open-source codes in Matlab of all the programs referred to and used here, their manuals, and examples enabling a reproduction of all the numerical results.

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Appendix A. Padé approximants in the optimizer project

The Padé approximant [52] of a dependence $y(x)$ can be defined as:

$$y(x) = C + \frac{P_n(x)}{P_d(x)} \equiv C + \frac{\sum_{k=0}^{N_n} a_k (x - x_0)^k}{\sum_{k=0}^{N_d} b_k (x - x_0)^k} \quad (6)$$

Equivalent forms parameterize the polynomials $P_n(x)$ and $P_d(x)$ via their nodes $z_k^{n/d}$ in place of the coefficients a_k and b_k :

$$P_{n/d} = C_{n/d} \prod_{k=1}^{N_n/d} (x - z_k^{n/d}). \quad (7)$$

For real-type functions, the nodes $z_k^{n/d}$ can either be real-type or pairs of mutually conjugated complex-type nodes. Our program supports both types of the parameterization.

A polynomial is a partial case of the Padé approximant with $P_d(x) = 1$, but in general, it is an approximation tool that has greater power. A relatively simple analytical form allows one to easily manage analytical properties such as poles, zeros, extrema, inflection points, asymptotes, etc.

Prospects of using the Padé approximants to describe potential energy curves of molecules have been discussed for several decades [53–65]. Nonetheless, their application in this field is still rather limited. One of the reasons is that, sometimes in the course of the optimization (fit) process, they can produce unexpected features, such as undesirable poles, bends, etc., between the nodes of a computational grid, while at the nodes everything looks fine; we encountered such problems with first versions of our programs. Murrel et al. [58], based on their test computations, did not recommend using Padé approximants for this purpose. A recent review [79] presenting 50 most popular analytical representations of potential energy functions did not even include the Padé approximants in this list.

However, these difficulties are easily overcome within the mechanisms of the Optimizer project. At every step of the optimization process, the desirable properties (absence of poles, absence or existence of predefined number of extrema and inflection points in any predefined range, etc.) can be checked either analytically or numerically, in a case of their violation causing a break of the computation and an automatic reduce of the optimization step until the desired properties are fulfilled. In part, we successfully used this strategy in the present work.

The potential energy functions of molecules at large interatomic distances $R \equiv x$ are expected [66] to exhibit the inverse-power character:

$$U(R \sim \infty) \sim C + C_1/R^{N_1} + C_2/R^{N_2} + \dots$$

with the leading powers depending on the state of a molecule (rather often, $N_1 = 3$, $N_2 = 6$, ...), while the so-called dispersion coefficients (C_1 , C_2 , ...) are either precomputed or estimated from experimental data. The Padé approximants can easily ensure a desirable asymptotic behavior.

Indeed, making the leading asymptotic term to be $\sim 1/R^{N_1}$ is achieved via making the power N_d of the polynomial $P_d(x)$ in Eq. (6) bigger than the power N_n of $P_n(x)$ by N_1 , i.e., $N_1 = N_d - N_n$, with $C_1 = a_{N_n}/b_{N_d}$. Then, assuming $x_0 = 0$ and substituting $q = 1/x$,

$$\frac{\sum_{k=0}^{N_n} a_k x^k}{\sum_{k=0}^{N_d} b_k x^k} - \frac{a_{N_n}}{b_{N_d}} \frac{1}{x^{N_d - N_n}} = q^{N_d - N_n} \times \frac{\sum_{k=0}^{N_n-1} a_k q^{N_n-k} - (a_{N_n}/b_{N_d}) \sum_{k=0}^{N_d-1} b_k q^{N_d-k}}{\sum_{k=0}^{N_d} b_k q^{N_d-k}}.$$

Hence, for the next asymptotic term:

$$N_{next} = N_d - N_n + 1, \\ C_{N_{next}} = \frac{a_{N_n-1} - (a_{N_n}/b_{N_d}) b_{N_d-1}}{b_{N_d}}.$$

The latter relation enables setting this term to any desirable value (including the zero one) via fixing the approximant structure and equating a part of its coefficients to some definite values.

For example, in order to the leading asymptotic term to be $\sim 1/x^3$, the power of the polynomial in the denominator must be bigger than the power of the polynomial in the numerator by 3: $N_d = N_n + 3$. The easiest way to ensure the behavior of $\sim 1/x^6$ for the next asymptotic term is (with $x_0 = 0$) to equate to zero two penultimate coefficients a_k and three penultimate coefficients b_k . If $b_{N_d} = 1$, then the leading dispersion coefficient C_1 will be just equal to a_{N_n} and the next one C_2 will be equal to the last nonzero penultimate coefficients $a_{k'}$ (with $k' = N_n - 3$ in our example). It is easy to extend this reasoning to asymptotic terms of higher orders.

Our program can handle the following problems:

- Compute values y_i of the Padé function (6), (7) at a predefined grid $\{x_i\}$ using a predefined set of parameters.
- Compute values of the function and its derivatives of predefined orders at a set of some extra points.
- Impose constraints of the equality of the function itself and/or its derivatives of predefined orders at predefined points to predefined values via correction of a predefined set of parameters.
- Compute the local design matrix (Jacobi matrix) of the dependence

$$y_i = f(\{a_k\}, \{b_k\}, x_0, C_{add}, C_{mul})$$

or the equivalent dependence on the nodes $z_k^{n/d}$ in place of a_k , b_k , in accordance with the regulations of the package Optimizer [2].

- Compute the polynomial coefficients a_k , b_k Eq. (6) from the nodes $z_k^{n/d}$ Eq. (7) or vice versa.
- Compute coordinates and function values of the extrema and the inflection points.
- Estimate coefficients at $1/x^p$ (including positive and negative integer p) in the long-range asymptote $x \rightarrow \infty$.
- Support working with a composite function, when the formal variable x is a predefined function of a physical variable R .

A more comprehensive manual is accessible online [2], with the program codes and executable samples.

Appendix B. Split-operator method in the optimizer project

The Split-operator method [42–48] is a kind of the so-called pseudospectral methods. It is aimed at a simulation of the dynamics of a quantum system and, based on it, of a probability (intensity) distribution in a spectrum of a transition in such a system.

The quantum evolution operator at a short time interval δt can be approximated with:

$$\begin{aligned} \mathcal{T}(\delta t) &= \exp \left\{ -\frac{i}{\hbar} H \delta t \right\} = \exp \left\{ -\frac{i}{\hbar} K \delta t - \frac{i}{\hbar} U \delta t \right\} \approx \\ &\exp \left\{ -\frac{i}{\hbar} K \delta t / 2 \right\} \exp \left\{ -\frac{i}{\hbar} U \delta t \right\} \exp \left\{ -\frac{i}{\hbar} K \delta t / 2 \right\} \approx \\ &\exp \left\{ -\frac{i}{\hbar} U \delta t / 2 \right\} \exp \left\{ -\frac{i}{\hbar} K \delta t \right\} \exp \left\{ -\frac{i}{\hbar} U \delta t / 2 \right\}, \end{aligned} \quad (8)$$

where H is the Hamiltonian, $K = -\frac{\hbar^2}{2m} \frac{\partial^2}{\partial R^2}$ is the operator of the system kinetic energy and $U(R)$ is the potential energy (generally, multichannel as in Eq. (1)). These forms are unitary, symmetrical and keep an important property of the time reversibility. Further on, we name these two approximations the KUK and UKU forms.

The operator $\exp \left\{ -\frac{i}{\hbar} U \delta t \right\}$ is diagonal relative the coordinate R in the coordinate representation and relative the channel index in the adiabatic representation, i.e., its action on a wavepacket is easily computed

(as just a multiplication) in the coordinate adiabatic representation. Analogously, the action of the operator $\exp\left\{-\frac{i}{\hbar}K\delta t\right\}$ is easily computed in the momentum diabatic representation. As is well known from Quantum Mechanics, the transform between the coordinate and momentum representations and *vice versa* is just the Fourier transform, while the transform between the adiabatic and diabatic representations is done with the transform matrices analogous to Eq. (4). This means that the evolution of a wavepacket $\psi(t=0, R)$ in an entire time range can be computed as a sequence of its multiplication by one diagonal operator in one of these representation at a reasonably short time interval δt , transform to the other representation, multiplication by another diagonal operator, transform back to the first representation, and so on until the desired final time is achieved.

For a computing of the evolution of a bound state, no extra complications are needed. However, if a free (decaying) state is considered, some artificial features arise, which should be damped. Indeed, the spatial computational grid is anyway finite, so, when the modeled wavepacket arrives at its boundary, it is partially reflected from it and partially enters the grid from the opposite boundary (in the theory of the Fourier transform, it is called the aliasing effect), demonstrating nonphysical behavior. To overcome this drawback, absorbing boundary conditions or an imaginary-type “optical” potential [72] are usually embedded.

When an optical transition of a molecule from an initial state with the wavefunction $\varphi(R)$ into a (multichannel) system of final states is considered, the initial multichannel wavepacket $\psi(t=0, R)$ is constructed via a multiplication of $\varphi(R)$ by a corresponding transition moment function $M_k(R)$ in every k th channel: $\psi_k(t=0, R) = M_k(R)\varphi(R)$. Then the evolution $\psi(t, R)$ is calculated the way described above with the estimate at every time instant of a time autocorrelation function

$$S(t) = \langle \psi(t=0, R) | \psi(t, R) \rangle$$

and the distribution of the transition probability (the spectrum of the transition) is estimated as the Fourier transform of the autocorrelator $S(t)$ (see also [46]).

We wrote and adapted to the Optimizer project several variants of the Split-operator programs. First, we realized both KUK and UKU schemes Eq. (8). Our test computations have shown that these two schemes possess approximately equivalent accuracy, however the KUK scheme is a little bit faster (see the manual [2] for an explanation why). Then, every program is written in both pure Matlab codes (vMatlab versions) and with blocks written in C and embedded with the help of the Matlab mex-mechanism (vMatlabC versions). The vMatlabC versions are several times faster than the vMatlab version but have a somewhat poorer functionality, that is not nevertheless important for the majority of prospective applications. The absorbing boundary conditions described in our recent work [72] are supported by all the versions.

A more detailed manual is available on the internet [2], along with the program codes and executable samples.

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