

2D Vibron Model Program Suite

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Contents

1 Abstract

This is the documentation of the 2DVM_U3 program suite that is composed of mainly Fortran 90 but also Perl and Python scripts to perform calculations using the 2D limit of the Vibron model (2DVM).

2 Theory Outline

The original paper where the 2DVM was presented is Ref.~[**Iachello1996**]. We also make extensive use of the detailed description of the model provided in Ref.~[**PBernal2008**]. Other references deal mainly with the application of the 2DVM to molecular bending vibrations [**Ishikawa2002**, **Iachello2003**, **PBernal2005**], quantum phase transitions [**Caprio2008**, **PBernal2010**, **PFernandez2011**, **PRA86**, **Santos2015**, **santos2016**, **Castanos2016**, **PB_Santos2016**], or both [**Larese2011**, **algonod1**]. New developments also consider the coupling of several $U(3)$ dynamical algebras for the modeling of systems with several degrees of freedom [**mp_u3xu3**, **pla_376**, **cocoyoc**, **Larese2014**, **Calixto2014**].

We proceed to review the main theory results used in this program suite.

The 2DVM has an $U(3)$ dynamical algebra whose nine generators are built as bilinear products of one scalar σ^\dagger and two circular τ_\pm^\dagger bosonic creation operators with the corresponding annihilation operators, σ, τ_\pm [**PBernal2008**]. The nine generators are recasted to provide a deeper physical insight as [**Iachello1996**]

$$\begin{aligned} \hat{n} &= \tau_+^\dagger \tau_+ + \tau_-^\dagger \tau_- & , \quad \hat{n}_s &= \sigma^\dagger \sigma \\ \hat{\ell} &= \tau_+^\dagger \tau_+ - \tau_-^\dagger \tau_- & , \quad \hat{Q}_\pm &= \sqrt{2} \tau_\pm^\dagger \tau_\mp \\ \hat{D}_\pm &= \sqrt{2} (\pm \tau_\pm^\dagger \sigma \mp \sigma^\dagger \tau_\mp) & , \quad \hat{R}_\pm &= \sqrt{2} (\tau_\pm^\dagger \sigma + \sigma^\dagger \tau_\mp) . \end{aligned} \tag{1}$$

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There are two possible subalgebra chains starting from the dynamical algebra $U(3)$ and ending in the symmetry algebra $SO(2)$. The symmetry algebra has the 2D angular momentum $\hat{\ell}$ as generator, which imply that this is a conserved quantity:

$$\begin{array}{ccc}
 & U(2) & \text{Chain(I)} \\
 U(3) & \nearrow & \searrow \\
 & SO(2) & \\
 & \nwarrow & \nearrow \\
 & SO(3) & \text{Chain(II)}
 \end{array} \quad (2)$$

The corresponding subalgebras are composed by the following elements

$$\begin{array}{ll}
 U(2) & \{\hat{n}, \hat{\ell}, \hat{Q}_+, \hat{Q}_-\} \\
 SO(3) & \{\hat{l}, \hat{D}_+, \hat{D}_-\} \\
 SO(2) & \{\hat{l}\}
 \end{array} \quad (3)$$

where the $SO(3)$ elements satisfy the usual angular momentum commutation relations. Because of an automorphism of the Lie algebra $U(3)$ constructed with τ_{\pm} , σ there is an alternative $SO(3)$ subalgebra of $U(3)$, called $\overline{SO(3)}$, with elements

$$\overline{SO(3)} \quad \{\hat{l}, \hat{R}_+, \hat{R}_-\} . \quad (4)$$

Another ingredient of the algebraic approach are the Casimir (or invariant) operators associated to each subalgebra chain [frank, bookalg]. The first and second order Casimir operators for the subalgebras in Eqs. \ (??) and (??) are

$$\begin{aligned}
 \hat{C}_1[U(2)] &= \hat{n} \quad , \quad \hat{C}_2[U(2)] = \hat{n}(\hat{n} + 1) \\
 \hat{C}_2[SO(3)] &= \hat{W}^2 = (\hat{D}_+ \hat{D}_- + \hat{D}_- \hat{D}_+)/2 + \hat{l}^2 \\
 \hat{C}_1[SO(2)] &= \hat{l} \quad , \quad \hat{C}_2[SO(2)] = \hat{l}^2
 \end{aligned} \quad (5)$$

and

$$\hat{C}_2[\overline{SO(3)}] = \hat{\bar{W}}^2 = (\hat{R}_+ \hat{R}_- + \hat{R}_- \hat{R}_+)/2 + \hat{\ell}^2 . \quad (6)$$

We proceed to state the basis quantum numbers and branching rules for each dynamical symmetry.

2.1 $U(2)$ or Cylindrical Oscillator Dynamical Symmetry

In this chain states are labeled by the quantum numbers n and ℓ

$$\left| \begin{array}{ccc} U(3) & \supset & U(2) & \supset & SO(2) \\ [N] & & n & & \ell \end{array} \right\rangle , \quad (7)$$

and the basis is $|[N]; n, \ell\rangle$. An alternative notation, often used in molecular physics, is $|[N]; n^\ell\rangle$. The quantum number N labels the totally symmetric representation of $U(3)$, $[N]$, and it is related to the total number of bound states of the system, n is the vibrational quantum number, and ℓ is the vibrational angular momentum. The branching rules are

$$\begin{aligned}
 n &= N, N-1, N-2, \dots, 0 \\
 l &= \pm n, \pm(n-2), \dots, \pm 1 \text{ or } 0, \quad (n = \text{odd or even}) .
 \end{aligned} \quad (8)$$

This is the natural basis to fit experimental bending vibration data in the case of linear and quasilinear molecules.

2.2 $SO(3)$ or Displaced Oscillator Dynamical Symmetry

States in this chain are characterized by the quantum numbers

$$\left| \begin{array}{c} U(3) \\ [N] \end{array} \supset \begin{array}{c} SO(3) \\ \omega \end{array} \supset \begin{array}{c} SO(2) \\ \ell \end{array} \right\rangle . \quad (9)$$

and will be labelled as $|[N]; \omega, \ell\rangle$. The branching rules are

$$\begin{aligned} \omega &= N, N-2, N-4, \dots, 1 \text{ or } 0, \quad (N = \text{odd or even}), \\ l &= \pm\omega, \pm(\omega-1), \dots, 0 . \end{aligned} \quad (10)$$

In this case, it is convenient to introduce a vibrational quantum number ν , which can be identified with the number of quanta of excitation in the displaced oscillator:

$$\nu = \frac{N - \omega}{2} . \quad (11)$$

The branching rules in this case are

$$\begin{aligned} \nu &= 0, 1, \dots, \frac{N-1}{2} \text{ or } \frac{N}{2}, \quad (N = \text{odd or even}), \\ l &= 0, \pm 1, \pm 2, \dots, \pm(N-2\nu) . \end{aligned} \quad (12)$$

This is the natural basis to fit experimental bending vibration data in the case of quasi-rigid and rigidly-bent molecules.

2.3 2DVM Hamiltonians included in the program suite

We present the three different Hamiltonians that are considered in the present programs. The simplest Hamiltonian, used mainly in the study of QPTs and ESQPTs, is the so called model Hamiltonian, that includes the $\hat{n} = \tau_+^\dagger \tau_+ + \tau_-^\dagger \tau_-$ operator from the dynamical symmetry (I) (DS-I) and the Pairing operator \hat{P} which is built adding a diagonal contribution to the $SO(3)$ second order Casimir operator $\hat{W}^2 = (\hat{D}_+ \hat{D}_- + \hat{D}_- \hat{D}_+)/2 + \hat{\ell}^2$. The Pairing operator is $\hat{P} = N(N+1) - \hat{W}^2$. the $SO(3)$ algebra).

The model Hamiltonian $\hat{\mathcal{H}}$ is then

$$\hat{\mathcal{H}} = \varepsilon \left[(1 - \xi) \hat{n} + \frac{\xi}{N-1} \hat{P} \right] . \quad (13)$$

The Hamiltonian has two parameters: a global energy scale ε and a control parameter ξ such that for $\xi = 0.0$ the system is in the DS-I and for $\xi = 1.0$ the system is in the DS-II. This is specially adequate to characterize the ground state and excited state QPTs as there is a second order ground state QPT for the critical value $\xi_c = 0.2$ [PBernal2008].

A second Hamiltonian is \hat{H}_{2b} , the most general one- and two-body Hamiltonian of the 2DVM defined in [Iachello1996].

$$\hat{H}_{2b} = E_0 + \varepsilon \hat{n} + \alpha \hat{n}(\hat{n} + 1) + \beta \hat{\ell}^2 + A \hat{P} , \quad (14)$$

where the operators \hat{n} and $\hat{n}(\hat{n} + 1)$ belong to the DS-I and the pairing operator \hat{P} to the DS-II while the vibrational angular momentum, $\hat{\ell}$, is common to both dynamical chains. In fact in all the cases considered the angular momentum is a constant of the motion, ℓ is a good quantum number, and the Hamiltonian are block diagonal in ℓ , a fact that simplifies the numerical calculations reducing the dimension of the considered matrices. In addition to this, as in the absence of external fields, the first

order angular momentum operator $\hat{\ell}$ is not included there is a degeneracy between positive and negative ℓ values for $\ell \neq 0$ and only $\ell \geq 0$ are considered. The formula for the dimension of a subspace of the totally symmetric representation N with angular momentum ℓ is $(N - |\ell|)/2 + 1$, if N and ℓ have the same parity, and $(N - |\ell| + 1)/2$ if they have different parity. Both cases are considered simultaneously in the programs as

$$\dim(N, \ell) = \frac{N - |\ell| + \text{mod}(N - |\ell|, 2) + 1}{2} . \quad (15)$$

The third and last Hamiltonian considered is \hat{H}_{4b} , the most general 1-, 2-, 3-, and 4-body Hamiltonian, that can be expressed in terms of the Casimir operators \hat{n} , \hat{n}^2 , $\hat{\ell}^2$, \hat{W}^2 , and $\hat{\bar{W}}^2$ can be expressed as follows

$$\begin{aligned} \hat{H}_{4b} = & P_{11}\hat{n} \\ & + P_{21}\hat{n}^2 + P_{22}\hat{\ell}^2 + P_{23}\hat{W}^2 \\ & + P_{31}\hat{n}^3 + P_{32}\hat{n}\hat{\ell}^2 + P_{33}(\hat{n}\hat{W}^2 + \hat{W}^2\hat{n}) \\ & + P_{41}\hat{n}^4 + P_{42}\hat{n}^2\hat{\ell}^2 + P_{43}\hat{\ell}^4 + P_{44}\hat{\ell}^2\hat{W}^2 \\ & + P_{45}(\hat{n}^2\hat{W}^2 + \hat{W}^2\hat{n}^2) + P_{46}\hat{W}^4 + P_{47}(\hat{W}^2\hat{\bar{W}}^2 + \hat{\bar{W}}^2\hat{W}^2)/2 . \end{aligned} \quad (16)$$

The Hamiltonian has fourteen spectroscopic constants P_{ij} , where the subindices indicate that this parameter corresponds to the j -th operator of the i -body operators.

From the matrix elements of the creation and annihilation sigma and tau bosons in the two dynamical symmetries published in [PBernal2008], the ensuing operator matrix elements can be derived.

2.3.1 Operator Matrix Elements in the Dynamical Symmetry (I)

The diagonal operators in this dynamical symmetry are

Operator \hat{n}^p $\langle [N]; n^\ell | \hat{n}^p | [N]; n^\ell \rangle = n^p$ for $p = 1, 2, 3, 4$.

Operator $\hat{\ell}^{2q}$ $\langle [N]; n^\ell | \hat{\ell}^{2q} | [N]; n^\ell \rangle = \ell^{2q}$ for $q = 1, 2$.

Operator $\hat{n}^p \hat{\ell}^{2q}$ $\langle [N]; n^\ell | \hat{n}^p \hat{\ell}^{2q} | [N]; n^\ell \rangle = n^p \ell^{2q}$ for $p = 1, 2$ and $q = 1$.

The non-diagonal matrix elements in this basis are

$SO(3)$ Casimir Operator \hat{W}^2

$$\begin{aligned} \langle [N]; n_2^l | \hat{W}^2 | [N]; n_1^l \rangle = & [(N - n_1)(n_1 + 2) + (N - n_1 + 1)n_1 + l^2] \delta_{n_2, n_1} \\ & - \sqrt{(N - n_1 + 2)(N - n_1 + 1)(n_1 + l)(n_1 - l)} \delta_{n_2, n_1 - 2} \\ & - \sqrt{(N - n_1)(N - n_1 - 1)(n_1 + l + 2)(n_1 - l + 2)} \delta_{n_2, n_1 + 2} . \end{aligned} \quad (17)$$

Note that this is the main nondiagonal operator in this case and it is a band matrix as the non-zero matrix elements are located in the main and first diagonals only.

Operator $\hat{n}\hat{W}^2 + \hat{W}^2\hat{n}$ As the operator \hat{n} is diagonal the matrix is also band diagonal with matrix elements

$$\begin{aligned} \langle [N]; n_2^l | \hat{n}\hat{W}^2 + \hat{W}^2\hat{n} | [N]; n_1^l \rangle = & 2n_1 [(N - n_1)(n_1 + 2) + (N - n_1 + 1)n_1 + l^2] \delta_{n_2, n_1} \\ & - (2n_1 - 2) \sqrt{(N - n_1 + 2)(N - n_1 + 1)(n_1 + l)(n_1 - l)} \delta_{n_2, n_1 - 2} \\ & - (2n_1 + 2) \sqrt{(N - n_1)(N - n_1 - 1)(n_1 + l + 2)(n_1 - l + 2)} \delta_{n_2, n_1 + 2} . \end{aligned} \quad (18)$$

Operator $\hat{\ell}^2\hat{W}^2$ This operator is computed for $\ell \neq 0$ multiplying the \hat{W}^2 operator matrix times ℓ^2 .

Operator $\hat{n}^2\hat{W}^2 + \hat{W}^2\hat{n}^2$ This is computed as the $\hat{n}\hat{W}^2 + \hat{W}^2\hat{n}$ operator.

$$\begin{aligned} \langle [N]; n_2^l | \hat{n}^2\hat{W}^2 + \hat{W}^2\hat{n}^2 | [N]; n_1^l \rangle = & 2n_1^2 [(N - n_1)(n_1 + 2) + (N - n_1 + 1)n_1 + l^2] \delta_{n_2, n_1} \\ & - [n_1^2 + (n_1 - 2)^2] \sqrt{(N - n_1 + 2)(N - n_1 + 1)(n_1 + l)(n_1 - l)} \delta_{n_2, n_1 - 2} \\ & - [n_1^2 + (n_1 + 2)^2] \sqrt{(N - n_1)(N - n_1 - 1)(n_1 + l + 2)(n_1 - l + 2)} \delta_{n_2, n_1 + 2} . \end{aligned} \quad (19)$$

Operator \hat{W}^4 This operator is computed as the matrix product of the \hat{W}^2 operator matrix times itself.

Operator $\hat{W}^2\hat{\bar{W}}^2 + \hat{\bar{W}}^2\hat{W}^2$ In this basis the only difference between the matrix elements of the \hat{W}^2 and $\hat{\bar{W}}^2$ operators is the sign of the non-diagonal contribution, which is positive in this case. The full operator is computed via matrix multiplication.

2.3.2 Operator Matrix Elements in the Dynamical Symmetry (II)

The diagonal operators in this dynamical symmetry are

$$SO(3) \text{ Casimir Operator } \hat{W}^2 \quad \langle [N]; \omega\ell | \hat{W}^2 | [N]; \omega\ell \rangle = \omega(\omega + 1).$$

$$\text{Operator } \hat{\ell}^{2q} \quad \langle [N]; \omega\ell | \hat{\ell}^{2q} | [N]; \omega\ell \rangle = \ell^{2q} \text{ for } q = 1, 2.$$

$$\text{Operator } \hat{\ell}^2\hat{W}^2 \quad \langle [N]; \omega\ell | \hat{\ell}^2\hat{W}^2 | [N]; \omega\ell \rangle = \ell^2\omega(\omega + 1).$$

$$\text{Operator } \hat{W}^4 \quad \langle [N]; \omega\ell | \hat{W}^4 | [N]; \omega\ell \rangle = \omega^2(\omega + 1)^2.$$

The non-diagonal matrix elements in this basis are

Operator \hat{n}

$$\begin{aligned}
\langle [N]; w_2^l | \hat{n} | [N]; w_1^l \rangle &= \left\{ \frac{(N - w_1) [(w_1 - l + 2)(w_1 - l + 1) + (w_1 + l + 2)(w_1 + l + 1)]}{2(2w_1 + 1)(2w_1 + 3)} \right. \\
&\quad \left. + \frac{(N + w_1 + 1) [(w_1 + l)(w_1 + l - 1) + (w_1 - l)(w_1 - l - 1)]}{2(2w_1 + 1)(2w_1 - 1)} \right\} \delta_{w_2, w_1} \quad (20) \\
&\quad + \sqrt{\frac{(N - w_1)(N + w_1 + 3)(w_1 - l + 2)(w_1 - l + 1)(w_1 + l + 2)(w_1 + l + 1)}{(2w_1 + 1)(2w_1 + 3)^2(2w_1 + 5)}} \delta_{w_2, w_1 + 2} \\
&\quad + \sqrt{\frac{(N - w_1 + 2)(N + w_1 + 1)(w_1 - l)(w_1 - l - 1)(w_1 + l)(w_1 + l - 1)}{(2w_1 - 3)(2w_1 - 1)^2(2w_1 + 1)}} \delta_{w_2, w_1 - 2} \quad (21)
\end{aligned}$$

Note that this is the main nondiagonal operator in this case and it is again a band matrix with non-zero matrix elements are located in the main and first diagonals only. The \hat{n} matrix element in this basis are taken from [PBernal2008] with a typo that has been corrected.

Operators \hat{n}^2 , \hat{n}^3 , and \hat{n}^4 These three operators are computed by matrix multiplication of the basic operator (??)

Operator $\hat{n}\ell^2$ This operator is computed for $\ell \neq 0$ multiplying the \hat{n} operator matrix times ℓ^2 .

Operator $\hat{n}\hat{W}^2 + \hat{W}^2\hat{n}$ As the operator \hat{n} is band diagonal \hat{W}^2 is diagonal this operator matrix is also band diagonal with matrix elements

$$\begin{aligned}
\langle [N]; w_2\ell | \hat{n}\hat{W}^2 + \hat{W}^2\hat{n} | [N]; w_1\ell \rangle &= 2\omega_1(\omega_1 + 1) \left\{ \frac{(N - w_1) [(w_1 - l + 2)(w_1 - l + 1) + (w_1 + l + 2)(w_1 + l + 1)]}{2(2w_1 + 1)(2w_1 + 3)} \right. \\
&\quad \left. + \frac{(N + w_1 + 1) [(w_1 + l)(w_1 + l - 1) + (w_1 - l)(w_1 - l - 1)]}{2(2w_1 + 1)(2w_1 - 1)} \right\} \delta_{w_2, w_1} \quad (22) \\
&\quad + [\omega_1(\omega_1 + 1) + (\omega_1 + 2)(\omega_1 + 3)] \\
&\quad \times \sqrt{\frac{(N - w_1)(N + w_1 + 3)(w_1 - l + 2)(w_1 - l + 1)(w_1 + l + 2)(w_1 + l + 1)}{(2w_1 + 1)(2w_1 + 3)^2(2w_1 + 5)}} \delta_{w_2, w_1 + 2} \\
&\quad + [(\omega_1 - 2)(\omega_1 - 1) + \omega_1(\omega_1 + 1)] \\
&\quad \times \sqrt{\frac{(N - w_1 + 2)(N + w_1 + 1)(w_1 - l)(w_1 - l - 1)(w_1 + l)(w_1 + l - 1)}{(2w_1 - 3)(2w_1 - 1)^2(2w_1 + 1)}} \delta_{w_2, w_1 - 2}
\end{aligned}$$

Operator $\hat{n}^2\hat{W}^2 + \hat{W}^2\hat{n}^2$ This is computed in the same way that the $\hat{n}\hat{W}^2 + \hat{W}^2\hat{n}$ operator but taking into account that the \hat{n}^2 operator is double banded. Therefore the operator matrix elements can be expressed as follow

$$\begin{aligned}
\langle [N]; w_2\ell | \hat{n}^2\hat{W}^2 + \hat{W}^2\hat{n}^2 | [N]; w_1\ell \rangle &= 2\omega_1(\omega_1 + 1) [\hat{n}^2]_{w_1, w_1} \delta_{w_2, w_1} \\
&\quad + [\omega_1(\omega_1 + 1) + (\omega_1 + 2)(\omega_1 + 3)] [\hat{n}^2]_{w_1, w_1 + 2} \delta_{w_2, w_1 + 2} \\
&\quad + [(\omega_1 - 2)(\omega_1 - 1) + \omega_1(\omega_1 + 1)] [\hat{n}^2]_{w_1, w_1 - 2} \delta_{w_2, w_1 - 2} \quad (23) \\
&\quad + [\omega_1(\omega_1 + 1) + (\omega_1 + 4)(\omega_1 + 5)] [\hat{n}^2]_{w_1, w_1 + 4} \delta_{w_2, w_1 + 4} \\
&\quad + [(\omega_1 - 4)(\omega_1 - 3) + \omega_1(\omega_1 + 1)] [\hat{n}^2]_{w_1, w_1 - 4} \delta_{w_2, w_1 - 4} ,
\end{aligned}$$

where $[\hat{n}^2]_{w_i, w_j}$ are the \hat{n}^2 operator matrix elements.

Operator $\hat{W}^2 \hat{\bar{W}}^2 + \hat{\bar{W}}^2 \hat{W}^2$ In this basis we need first to compute the matrix elements of the $\hat{\bar{W}}^2$ making use of Eqs. (37) and (38) of Ref. [PBernal2008].

$$\langle [N]; w_2 \ell_2 | \hat{R}_- | [N]; w_1 \ell_1 \rangle = A_{w_1, \ell_1} \delta_{w_2, w_1} \delta_{\ell_2, \ell_1 - 1} + B_{w_1, \ell_1} \delta_{w_2, w_1 - 2} \delta_{\ell_2, \ell_1 - 1} + C_{w_1, \ell_1} \delta_{w_2, w_1 + 2} \delta_{\ell_2, \ell_1 - 1}, \quad (24)$$

where

$$\begin{aligned} A_{w, \ell} &= \frac{(2N+3)(2\ell+1)}{(2w-1)(2w+3)} \sqrt{(w+\ell)(w-\ell+1)/2} \\ B_{w, \ell} &= -\sqrt{\frac{2(N+w+1)(N-w+2)(w+\ell)(w-\ell)(w+\ell-1)(w+\ell-2)}{(2w+1)(2w-1)^2(2w-3)}} \\ C_{w, \ell} &= \sqrt{\frac{2(N+w+3)(N-w)(w+\ell+1)(w-\ell+1)(w-\ell+2)(w-\ell+3)}{(2w+1)(2w+3)^2(2w+5)}}. \end{aligned}$$

The previous result can be used for the obtention of an expression for the \hat{R}_+ operator matrix elements

$$\begin{aligned} \langle [N]; w_2 \ell_2 | \hat{R}_- | [N]; w_1 \ell_1 \rangle^\dagger &= \langle [N]; w_1 \ell_1 | \hat{R}_+ | [N]; w_2 \ell_2 \rangle \\ &= A_{w_2, \ell_2 + 1} \delta_{w_1, w_2} \delta_{\ell_1, \ell_2 + 1} + B_{w_2 + 2, \ell_2 + 1} \delta_{w_1, w_2 - 2} \delta_{\ell_1, \ell_2 + 1} \\ &\quad + C_{w_2 - 2, \ell_2 + 1} \delta_{w_1, w_2 + 2} \delta_{\ell_1, \ell_2 + 1}. \end{aligned} \quad (25)$$

The upper diagonal matrix elements of the Casimir operator $\hat{\bar{W}}^2 = \hat{R}_+ \hat{R}_- + \hat{\ell}^2$ can then be expressed as

$$\begin{aligned} \langle [N]; w_2 \ell | \hat{\bar{W}}^2 | [N]; w_1 \ell \rangle &= A_{w_1, \ell_1}^2 + B_{w_1, \ell_1}^2 + C_{w_1, \ell_1}^2 \delta_{w_2, w_1} \\ &\quad + (A_{w_1, \ell} B_{w_1 + 2, \ell} + C_{w_1, \ell} A_{w_1 + 2, \ell}) \delta_{w_2, w_1 + 2} \\ &\quad + C_{w_1, \ell} B_{w_1 + 4, \ell} \delta_{w_2, w_1 + 4}, \end{aligned} \quad (26)$$

and the lower diagonal matrix elements can be computed from this equation.

The $\hat{W}^2 \hat{\bar{W}}^2 + \hat{\bar{W}}^2 \hat{W}^2$ operator is then computed as in Eq. (??)

$$\begin{aligned} \langle [N]; w_2 \ell | \hat{W}^2 \hat{\bar{W}}^2 + \hat{\bar{W}}^2 \hat{W}^2 | [N]; w_1 \ell \rangle &= 2\omega_1(\omega_1 + 1) [\hat{\bar{W}}^2]_{w_1, w_1} \delta_{w_2, w_1} \\ &\quad + [\omega_1(\omega_1 + 1) + (\omega_1 + 2)(\omega_1 + 3)] [\hat{\bar{W}}^2]_{w_1, w_1 + 2} \delta_{w_2, w_1 + 2} \\ &\quad + [(\omega_1 - 2)(\omega_1 - 1) + \omega_1(\omega_1 + 1)] [\hat{\bar{W}}^2]_{w_1, w_1 - 2} \delta_{w_2, w_1 - 2} \\ &\quad + [\omega_1(\omega_1 + 1) + (\omega_1 + 4)(\omega_1 + 5)] [\hat{\bar{W}}^2]_{w_1, w_1 + 4} \delta_{w_2, w_1 + 4} \\ &\quad + [(\omega_1 - 4)(\omega_1 - 3) + \omega_1(\omega_1 + 1)] [\hat{\bar{W}}^2]_{w_1, w_1 - 4} \delta_{w_2, w_1 - 4}, \end{aligned} \quad (27)$$

where $[\hat{n}^2]_{w_i, w_j}$ are the \hat{n}^2 operator matrix elements.

3 Programs included in the package

We proceed to describe the different programs provided. Compiled Fortran programs have names with the suffix `_gfortran` or `_ifort` depending on the compiler used.

The subroutine used for matrix diagonalization is `LA_SYEVR` from the Lapack95 [LAPACK95], the Fortran 95 interface to Lapack [laug].

3.1 Fortran programs to compute eigenvectors, eigenvalues, and other quantities

3.1.1 Programs avec_U3_U2_mh and avec_U3_S03_mh

The programs avec_U3_XX_mh compute, and eventually save, eigenvalues and eigenvectors of the model Hamiltonian (??) using the XX = U2 -Eq. (??)- or S03 -Eq. (??)- basis. The input file for this program is of namelist type with the following syntax

```
#
#      INPUT FILE FOR 2DVM U(3) Calculations
#      Model Hamiltonian
&par_aux Iprint = 1, Eigenvec_Log = .T., Excitation_Log = .F., Save_avec_Log = .T. /
&par_0   N_val = 20, L_val = 0 /
&par_1   epsilon = 1.0,   xi = 0.4 /
```

And the different variables are:

- *Iprint Integer variable.* Control program verbosity.
- *Eigenvec_Log Logical variable* If the variable is .True. eigenvalues and eigenvectors are computed.
- *Excitation_Log Logical variable* If the variable is .True. excitation energies wrt the ground state energy are reported.
- *Save_avec_Log Logical variable* If the variable is .True. eigenstates are saved in a file with the name eigvec_XX_NYY_LZZ.dat where XX = u2/so3, YY = N_val, and ZZ = L_val.
- *N_val Integer variable* Totally symmetric irrep of $U(3)$. Determines the Hilbert space size.
- *L_val Integer variable* Vibrational (2D) angular momentum ℓ value.
- *epsilon Real (DP) variable* Model Hamiltonian global energy scale ε in Eq.(??).
- *xi Real (DP) variable* Model Hamiltonian control parameter, $0 \leq \xi \leq 1$ defined in Eq. (??).

The result of running the program using the U2 basis with the previous input file is

```
$ ../bin/avec_U3_U2_mh_gfortran < u3_modham.inp
L_val =                0
                0    4.7178381890544925
-6.6913473582377064E-002 |                0 >
0.26757490446002280      |                2 >
-0.50918508872650692     |                4 >
0.59782591356876802     |                6 >
-0.47489555175329112     |                8 >
0.26470245150057914      |               10 >
-0.10411984650838124     |               12 >
2.8307962444698111E-002 |               14 >
-5.0262214812812631E-003 |               16 >
```



```

5.1057145041144211E-004 | 18 >
-1.9360623933001603E-005 | 20 >
.
.
.
20 20.560646361706880
4.8846592744607648E-008 | 0 >
7.4751184587909931E-007 | 2 >
6.4855302773322584E-006 | 4 >
4.1763856737063440E-005 | 6 >
2.2180851797258004E-004 | 8 >
1.0276743631749022E-003 | 10 >
4.3044673503380383E-003 | 12 >
1.6740682624946063E-002 | 14 >
6.2024723264416194E-002 | 16 >
0.22776900283209650 | 18 >
0.97158341333025411 | 20 >

```

As `Save_avec_Log = .T.` a file named `eigvec_u2_N20_L0.dat` is created. This file includes the energies, the Hamiltonian matrix diagonal¹, and the eigenstates in column-major order. It has the following structure, where `XX = u2` or `so3`, depending on the basis selected.

```

# N =          N_val ; L =          L_val ; XX basis
# Eigenvalues
E_1          E_2          ..... E_dim
# Hamiltonian Diagonal
H_11          H_22          ..... H_dimdim
Psi_1_1      Psi_2_1      ..... Psi_dim_1
.....
Psi_dim_1 Psi_dim_2 .... Psi_dim_dim

```

3.1.2 Programs avec_U3_U2 and avec_U3_S03

The Programs `avec_U3_XX` compute the 2D Vibron Model energies and eigenstates using the `XX` chain basis for a given set of parameters of the most general one- and two-body Hamiltonian in Eq. (??). The input file is similar to the input in the previous section ??, the main difference being the Hamiltonian parameters:

```

#
#          INPUT FILE FOR U(3) 2 Body Hamiltonian Calculations
#
&par_aux Iprint = 0, Eigenvec_Log = .T., Excitation_Log = .T., Save_avec_Log = .F. /
&par_0   N_val = 10, L_val = 0 /
&par_1   epsilon_p=0.8, alpha_p=-0.000667779632721202, beta_p=0.0, A_p=0.0003339 /

```

The new elements in this input file are ϵ , α , β and A of the -Eq. (??). To run this program using a UNIX terminal:

¹In case `Excitation_Log = .True.` the ground state energy is also subtracted from the Hamiltonian matrix diagonal.

```
$ [...]/bin/avec_U3_XX_gfortran < u3_ham.inp
```

where [...] is the path of the program's bin folder. The output of running it with the given input is:

```
0 0.000000000000000000
2 1.5859642569051831
4 3.1719656273638952
6 4.7579771193315414
8 6.3439717420554338
10 7.9299225026296547
```

3.1.3 Programs ipr_U3_U2_mh and ipr_U3_S03_mh

The programs ipr_U3_XX_mh compute the energies and the Inverse Participation Ratio (IPR) in the XX chain for -Eq.(??) Hamiltonian. The input looks like the last one:

```
#
# INPUT FILE FOR U(3) Model Hamiltonian Calculations
#
&par_aux Iprint = 2, Eigenvec_Log = .T., Excitation_Log = .T. /
&par_0 N_val = 15, L_val = 0 /
&par_1 epsilon = 1.0, xi = 0.75 /
```

Introducing

```
$ [...]/bin/ipr_U3_U2_mh_gfortran < ipr_u3_modham.inp
```

in the command line, it is given back the following output:

```
Reading N_val, L_val
Reading epsilon, xi
Iprint = 2; Eigenvec_LOG = T; Excitation_Log = T
N_val = 15; L_val = 0
epsilon = 1.000000E+00; xi = 7.500000E-01
Operator number 1
Operator number 2
GS_energy = 1.7321844551756294
L_val = 0
1 0.000000000000000000 3.5165700354623559
2 3.0929130257955548 4.4667740770364324
3 5.7488037073509926 4.8502861663079422
4 7.9586934507799825 5.5371533436156843
5 9.6925122152480512 4.1265672238584008
6 10.908945553417363 4.9730348558314166
7 12.086226648623684 5.5191888106052103
8 13.654429757379329 2.7538257245146016
```

In this case, the verbosity has been increased using IPRINT=2.

3.1.4 Program ipr_Husimi_U3_U2_mh

3.1.5 Programs ipr_U3_U2 and ipr_U3_S03

The ipr_U3_XX program is similar to ipr_U3_XX_mh. The main difference is the input file, because of the Hamiltonian used is -Eq. (??) instead of -Eq.(??). In this case, an example of the input file could be:

```
#
#      INPUT FILE FOR U(3) Model Hamiltonian Calculations
#
&par_aux Iprint = 2, Eigenvec_Log = .T., Excitation_Log = .T., Save_avec_Log = .F./
&par_0   N_val = 30, L_val = 0 /
&par_1   epsilon_p = 1.0, alpha_p = 0.0, beta_p = 0.0, A_p = 0.0 /
```

Running it in the $U(2)$ chain, we can analyze the output to test the program:

```
$ [...] /bin/ipr_U3_U2_gfortran < ipr_u3_ham.inp
```

```
Reading  N_val, L_val
Reading  Hamiltonian Paramenters
  Iprint =  2; Eigenvec_LOG =  T; Excitation_Log =  T; Save_Avec_Log =  F
N_val =      30; L_val =      0
epsilon =  1.0000000E+00; alpha =  0.0000000E+00 ; beta =  0.0000000E+00; A =  0.0000000
Operator number      1
Operator number      2
Operator number      3
Operator number      4
GS_energy =  0.000000000000000000
L_val =      0
      1  0.000000000000000000      1.000000000000000000
      2  2.000000000000000000      1.000000000000000000
      3  4.000000000000000000      1.000000000000000000
      4  6.000000000000000000      1.000000000000000000
      5  8.000000000000000000      1.000000000000000000
      6 10.000000000000000000      1.000000000000000000
      7 12.000000000000000000      1.000000000000000000
      8 14.000000000000000000      1.000000000000000000
      9 16.000000000000000000      1.000000000000000000
     10 18.000000000000000000      1.000000000000000000
     11 20.000000000000000000      1.000000000000000000
     12 22.000000000000000000      1.000000000000000000
     13 24.000000000000000000      1.000000000000000000
     14 26.000000000000000000      1.000000000000000000
     15 28.000000000000000000      1.000000000000000000
     16 30.000000000000000000      1.000000000000000000
```

The output consist of three columns: an integer one, the eigenvalues of \hat{n} and the IPR. The last column was expected because \hat{n} eigenstates are well located in the $U(2)$ basis (they match basis' components). Now, if we compute it using $SO(3)$ basis,

```
$ [...]/bin/ipr_U3_S03_gfortran < ipr_u3_ham.inp
```

```
Reading  N_val, L_val
Reading  Hamiltonian Paramenters
  Iprint =  2; Eigenvec_LOG =  T; Excitation_Log =  T; Save_Avec_Log =  F
N_val =    30; L_val =    0
epsilon =  1.0000000E+00; alpha =  0.0000000E+00 ; beta =  0.0000000E+00; A =  0.0000000
Operator number      1
Operator number      2
Operator number      3
Operator number      4
GS_energy =  -7.1054273576010019E-015
L_val =
      0
      1  0.0000000000000000      6.2158795395844608
      2  2.0000000000000142      7.8800707358483129
      3  4.0000000000000107      8.8145249866016897
      4  6.0000000000000213      9.3394577175011282
      5  8.0000000000000284      9.5645594435444661
      6  10.000000000000000      9.5200527843596721
      7  12.000000000000014      9.1667631466188073
      8  14.000000000000014      8.2391939350606123
      9  16.000000000000014      8.2352287186286670
     10  18.000000000000007      9.1475889006614128
     11  20.000000000000021      9.4784447604423558
     12  22.000000000000011      9.4915382955106420
     13  24.000000000000014      9.2204419120130368
     14  26.000000000000011      8.6203298376166941
     15  28.000000000000004      7.5323034205731147
     16  30.000000000000007      5.3098659835781525
```

we obtain the same result excluding the IPR column. In the $SO(3)$ basis, the \hat{n} eigenstates are not well located.

3.1.6 Program ipr_4b_U3_U2 and ipr_4b_U3_S03

If the reader needs more precision, ipr_4b_U3_XX program is available to compute the energies, eigenvectors and IPR in both chains considering higher interactions of the Hamiltonian. Here, the difference is the same than in the last case: the input file.

```
&INP1 N_val=10, L_val=1, IPRINT=1, Eigenvec_log = .TRUE., Excitation_Log = .True., Save_
# Hamiltonian Parameters
&INP1b P11=2378.0D0 /
&INP2b P21=-37.597D0, P22=19.312D0, P23=-9.6422D0 /
&INP3b P31=0.0D0, P32=0.0D0, P33=0.0D0 /
&INP4b P41=0.0D0, P42=0.0D0, P43=0.0D0, P44=0.0D0, P45=0.0D0, P46=0.0D0, P47=0.0D0 /
```

P_{ij} elements correspond to the parameters of the Hamiltonian in the -Eq. (??). To run this program in the $SO(3)$ chain, for example, write the following commands in the terminal:

```
$ echo ipr_4b.inp | [...]/bin/ipr_4b_U3_S03_gfortran
```

The output in this case is

```
GS_energy = -200.60192731579809
L_val = 0
1 0.0000000000000000 3.7732656477725937
2 4311.9024599974764 4.1827037058586312
3 8488.9872790600057 3.5354457958151770
4 12523.812560924473 3.8730517993482945
5 16407.548813126232 3.9800419071796886
6 20129.470450786630 3.1403615181153479
```

3.2 Programs to fit experimental vibrational bending spectra

This part of the software has two different purposes depending on the input definitions, so, before explaining them, we present experimental energies and input files.

3.2.1 Program chi2_U3

2D Vibron Model Program Suite includes a chi2_U3 program to compute the χ^2 and use it in different minimization methods.

This part of the software has two different purposes depending on the input definitions, so, before explaining them, we present experimental energies and input files.

First of all we need to collect the experimental data into a file. In this example we use the hydrogen isocyanide (HNC) molecule [**MellauHNC**, **BunkerHNC**].

```
19
462.72 1.0 1 1
926.50 1.0 2 0
936.05 1.0 2 2
1398.56 1.0 3 1
1419.97 1.0 3 3
1867.05 1.0 4 0
1878.72 1.0 4 2
1913.87 1.0 4 4
2341.84 1.0 5 1
2366.83 1.0 5 3
2417.57 1.0 5 5
2809.29 1.0 6 0
2822.75 1.0 6 2
2863.11 1.0 6 4
2930.90 1.0 6 6
3281.50 1.0 7 1
3309.78 1.0 7 3
3367.37 1.0 7 5
3453.78 1.0 7 7
```

The first line must be the total number of experimental data. After that, the file is structured in 4 columns. The first column corresponds with the energy, the next one to the experimental error of data and, the last

two are the quantum numbers. If you want to include any data in the prediction but not at the fitting time, you can set the experimntal error to 0.0 . The energy's label must be in accordance with the nature of the molecule: linear, $U(2)$ basis $|n\ell\rangle$, or bent, $SO(3)$ $|v_b^\ell\rangle$.

Once finished, the next step is to construct the input file:

```
&INP0 BENT=.F., exp_data_file='exp_HNC_Danielle.dat' /
&INP1 N_val=40, LMAX=7, VMAX=7, EMINL=.F. /
#
# PARAMETERS THAT TUNE THE OUTPUT
#
&INP2 IPRINT=0, DIS_RES = *** /
&INP1b P11=1414.0D0 /
&INP2b P21=-29.837D0, P22=15.81D0, P23=-8.054D0 /
&INP3b P31=0.0D0, P32=4.9D-2, P33=0.0D0 /
&INP4b P41=0.0D0, P42=0.0D0, P43=0.0D0, P44=0.0D0, P45=0.0D0, P46=0.0D0, P47=0.0D0 /
```

- BENT: .T. (.F.) if the molecule is bent (linear).
- exp_data_file: The name of the experimental energies' file.
- N_val: Vibron number. Determines the Hilbert space size.
- LMAX: The highest angular momentum number ℓ in exp_data_file.
- VMAX: The highest harmonic number n for the linear case or the bending number v_b when BENT is true.
- EMINL: Fix the lowest level at the zero energy.
- Iprint: Integer variable. Controls the program's verbosity.
- DIS_RES: Match as true (.T.) to display residuals energies or false (.F.) to get only the χ^2 .
- P_{ij} : Initial value of the parameter P_{ij}

Running the program with DIS_RES=.T.:

```
$ echo fit.inp | ../../bin/chi2_U3_gfortran
```

```
7.0788432691415437E-003
-2.7776246656685544E-004
9.1877661752732820E-003
-3.3695279357061736E-002
-3.0352671044056478E-002
-8.4165727005256485E-002
-4.9249332600084017E-002
5.6619175268906474E-002
-3.3439091556601852E-002
-4.1176236554747447E-002
-5.0164535727162729E-002
7.7310508308073622E-002
0.16723874181343490
```

```

-2.9727267470207153E-002
 1.0623963672060199E-002
 5.5108955660216452E-002
-0.11479814853555581
 0.16486361585430132
 0.19601871556506012

```

On the other hand, if you set DIS_RES to false:

```

$ echo fit.inp | ../../bin/chi2_U3_gfortran

0.13699774310846663

```

3.2.2 Minimization using the Python-script

3.2.3 Minimization using Minuit-CERN package

In this section we are going to explain how to use this minimization method, where Minuit-CERN package [**minuit**] has been used. Most of the inputs items have been detailed at subsection **??**. The energies file have the same structure and the input file is similar changing the meaning of DIS_RES:

```

&INP0 BENT=.F., exp_data_file='exp_HNC.dat', output_file_0="output_HNC" /
&INP1 N_val=40, LMAX=7, VMAX=7, EMINL=.F. /
#
# PARAMETERS THAT TUNE THE OUTPUT
#
&INP2 IPRINT=0, DIS_RES = .F. /
&INP1b P11=740.0D0 /
&INP2b P21=-3.597D0, P22=1.5D0, P23=-7.2D0 /
&INP3b P31=0.0D0, P32=0.002D0, P33=0.0D0 /
&INP4b P41=0.0D0, P42=0.0D0, P43=0.0D0, P44=0.0D0, P45=0.0D0, P46=0.0D0, P47=0.0D0 /
&fix_par fixed_par = 'fix 5 7 8 9 10 11 12 13 14' /

# P11 ---> 1
# P21 ---> 2
# P22 ---> 3
# P23 ---> 4
# P31 ---> 5
# P32 ---> 6
# P33 ---> 7
# P41 ---> 8
# P42 ---> 9
# P43 ---> 10
# P44 ---> 11
# P45 ---> 12
# P46 ---> 13
# P47 ---> 14

```

Most of definitions have been explained in previous sections, the others mean:

- BENT: .T. (.F.) if the molecule is bent (linear).
- exp_data_file: The name of the experimental energies' file.
- output_file_0: To choose the name of the file where the fit report will be saved.
- LMAX: The highest angular momentum number ℓ in exp_data_file.
- VMAX: The highest harmonic number n for the linear case or the bending number v_b when BENT is true.
- DIS_RES: Match as true (.T.) to display residuals energies.
- P_{ij} : Initial value of the parameter P_{ij}
- fixed_par: Minuit order. This string of characters consists of different integers that are associated with the P_{ij} parameters. If one of them appears, Minuit will keep it fixed.

If the program Min_minuit_gfortran is executed as mentioned:

```
$ echo fit.inp | ../bin/Min_minuit_gfortran
```

Note: The following floating-point exceptions are signalling: IEEE_INVALID_FLAG IEEE_DIVI

```
$ tail -n 43 output_HNC
```

EXT NO.	PARAMETER NAME	VALUE	APPROXIMATE ERROR	STEP SIZE	FIRST DERIVATIVE
1	P11	1413.8	0.37063	0.67417E-03	-0.14902
2	P21	-29.832	0.15233E-01	0.14225E-04	-1.8667
3	P22	15.807	0.10429	0.75373E-05	-0.14985
4	P23	-8.0528	0.28648E-02	0.38399E-05	-13.647
5	P31	0.0000	fixed		
6	P32	0.48757E-01	0.95612E-02	0.65872E-06	-1.3627
7	P33	0.0000	fixed		
8	P41	0.0000	fixed		
9	P42	0.0000	fixed		
10	P43	0.0000	fixed		
11	P44	0.0000	fixed		
12	P45	0.0000	fixed		
13	P46	0.0000	fixed		
14	P47	0.0000	fixed		

```
EXTERNAL ERROR MATRIX.      NDIM=  50      NPAR=  5      ERR DEF=  1.00
 0.137E+00-0.432E-02-0.926E-03-0.915E-03 0.200E-03
-0.432E-02 0.232E-03 0.563E-03 0.154E-04-0.537E-04
-0.926E-03 0.563E-03 0.109E-01-0.862E-04-0.978E-03
-0.915E-03 0.154E-04-0.862E-04 0.821E-05 0.628E-05
 0.200E-03-0.537E-04-0.978E-03 0.628E-05 0.914E-04
ERR MATRIX NOT POS-DEF
```

```
PARAMETER CORRELATION COEFFICIENTS
      NO.  GLOBAL      1      2      3      4      6
```



```

1  0.99666  1.000-0.766-0.024-0.861  0.056
2  0.98707 -0.766 1.000 0.355 0.353-0.369
3  0.99201 -0.024 0.355 1.000-0.288-0.980
4  0.99374 -0.861 0.353-0.288 1.000 0.229
6  0.99086  0.056-0.369-0.980 0.229 1.000
ERR MATRIX NOT POS-DEF
minuit> *****
**      4 **CALL      3.000
*****

-----
| rms =      7.9576384169698269E-002 |
|-----|
minuit> *****
**      5 **EXIT
*****
.....MINUIT TERMINATED BY MINUIT COMMAND: EXIT

$

```

3.3 Perl scripts