Optimized Structure and Vibrational Properties by Error Affected Potential Energy Surfaces

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ABSTRACT: The precise theoretical determination of the geometrical parameters of molecules at the minima of their potential energy surface and of the corresponding vibrational properties are of fundamental importance for the interpretation of vibrational spectroscopy experiments. Quantum Monte Carlo techniques are correlated electronic structure methods promising for large molecules, where intrinsic errors are of fundamental importance for the interpretation of vibrational spectroscopy experiments. Quantum Monte Carlo techniques are correlated electronic structure methods promising for large molecules, which are intrinsically affected by stochastic errors on both energy and force calculations, making the mentioned calculations more challenging with respect to other more traditional quantum chemistry tools. To circumvent this drawback in the present work, we formulate the general problem of evaluating electronic correlation functions 30 limit the application of DFT to molecules where electronic correlation does not play a crucial role, leaving the correct treatment of such cases to more accurate post-Hartree-Fock methods. Unfortunately, the latter methods are computationally highly demanding as the number of electrons $N_e$ increases, since they scale as $O(N_e^{6\text{-}10})$. An alternative to the most traditional quantum chemical approaches is represented by quantum Monte Carlo (QMC),31−34 a name that stands for a collection of wavefunction-based methods for the stochastic solution of the electronic many-body Schrödinger equation. The simplest QMC approach is the variational Monte Carlo (VMC), which relies on the evaluation of the expectation value of the

PES around the configurational minimum up to the fourth order expansion to predict most of the spectroscopic parameters. An alternative approach, which directly includes finite temperature nuclear motions, is represented by ab initio molecular dynamics simulations,28,29 where IR and Raman spectra can be directly obtained from the Fourier transform of dipole and polarizability autocorrelation functions. Provided that we can proficiently account for the anharmonicity of the PES within a PT2 approach, at least for not too large-amplitude motions, the accuracy of the adopted electronic structure calculations becomes the leading ingredient for obtaining accurate results. A convenient balance between accuracy and computational cost is given by DFT, that scales as $O(N_e^{3\text{-}5})$, $N_e$ being the number of electrons in the system. However, well-known drawbacks of the available exchange-correlation functionals30 limit the application of DFT to molecules where electronic correlation does not play a crucial role, leaving the correct treatment of such cases to more accurate post-Hartree-Fock methods. Unfortunately, the latter methods are computationally highly demanding as the number of electrons $N_e$ increases, since they scale as $O(N_e^{6\text{-}10})$. Among these approximative methods, an approach based on second order vibrational perturbation theory (PT2) is successfully widely used,16−27 which takes the force fields of the
molecular Hamiltonian on a given parametrized wave function $\psi$. The parameters are chosen to minimize the variational energy which is calculated through stochastic integration. The choice of the wave function is crucial and is generally composed by an antisymmetrical determinantal part, aimed to describe static correlation effects, and a bosonic part, the Jastrow factor, which recovers most of the dynamical correlation effects. Recent wave function developments include the Jastrow-antisymmetrized geminal power (J-AGP), the Pfaffian, the backflow, and other multideterminant-Jastrow functions. Other QMC schemes beyond the variational ansatz are often based on projection methods. Although they are usually more accurate, they are computationally more demanding. For instance, the diffusion Monte Carlo (DMC) consists of an imaginary time evolution of a trial wave function that in this way is projected onto the ground state.

Within QMC, the energy of correlated molecular systems can be calculated with an accuracy in line with experimental data and with the most accurate traditional quantum chemistry methods but with a computational cost scaling with the size of the system as $O(N^3 \epsilon^{-4})$. These favorable scaling properties, the high accuracy of the results, and the fact that QMC algorithms are intrinsically parallel (demonstrated by an almost perfect scalability up to tens of thousands of processors at recent high performance computing facilities) should make QMC an optimal choice for accurate calculations of molecular geometries and vibrational properties. On the other hand, the underlying stochastic nature of the Monte Carlo algorithms is reflected in the fact that QMC energies and forces are affected by a stochastic error, which slowly decreases as the square root of the computational time. Due to this major drawback, structural optimizations of molecules using QMC methods have been efficiently implemented only in very recent years. To tackle this problem, in a recent work Sorella and co-workers proposed a combination of the reweighting method, the correlated sampling technique, and the space warp coordinate transformation. With the help of the algorithmic adjoint differentiation, all the components of the ionic VMC forces can be calculated with and without pseudopotentials in a computational time that is only about 4 times that of an ordinary energy calculation. Vibrational properties, which require the calculation of second derivatives of the energy, to the best of our knowledge have never been investigated with the help of preliminary DFT calculations where an uncertainty of the order of magnitude of typical QMC measures has been introduced. In the first section of the present paper, we review the calculation of vibrational properties from PT2, and in the second section, we address the problem of fitting the PES using error affected measures. Finally, we illustrate how to proceed in the choice of the grid points for the specific case of the water molecule, demonstrating the feasibility of the method for QMC calculations.

2. CALCULATION OF VIBRATIONAL PROPERTIES

We, as usual, separate the electronic and nuclear part of the Coulomb Hamiltonian following the Born–Oppenheimer approximation. We consider a set of configurations of the nuclear coordinates close to the equilibrium structure, and for each configuration we obtain the ground state energy and nuclear forces using a DFT or QMC calculation. The energy values can be fitted with a parametric function, in order to obtain the potential energy surface (PES) of the molecule in the neighborhood of the equilibrium structure. This parametric PES enters into the nuclear Hamiltonian, which describes the quantum mechanical motion of the nuclei, i.e., translations, rotations, and vibrations of the molecule.

In the following section, we will provide a short overview of the well established theory for the description of molecular vibrational properties, considering first the formulation of the nuclear problem in Cartesian and internal coordinates, followed by the harmonic approximation, and the second order perturbation theory (PT2) for the calculation of the anharmonic corrections. In the next section, we will consider the problem of fitting the PES from QMC calculations, which, differently from most quantum chemical methods for electronic structure calculation, is affected by stochastic errors.

2.1. Nuclear Hamiltonian. The nuclear Hamiltonian $\hat{H}$ is the summation of the kinetic $\hat{T}$ and potential $\hat{V}$ energy operators, the latter operator being given by the fitted PES. Given a molecule with $N$ atoms, the two terms can be easily written in terms of the 3N-dimensional Cartesian coordinates $R$:

$$\hat{H} = \hat{T} + \hat{V} = -\frac{1}{2} \sum_\xi \sum_\alpha \frac{1}{M_\alpha} \frac{1}{R_{\xi\alpha}^2} + V(R)$$

where $M_\alpha$ is the mass of the atom $\xi$ and $R_{\xi\alpha}$ is its Cartesian coordinate along the $\alpha$ axis. The use of atomic units is always implicitly assumed hereafter.

The potential energy is generally assumed to be invariant under rotations and translations of the molecule as a whole (hypothesis of homogeneous and isotropic space). Therefore, it is independent of the position of the center of mass and on the orientation of the molecule (i.e., the 6 external degrees of freedom for a nonlinear molecule, or 5 in the case of a linear one), and the potential energy is often conveniently written in terms of $3N - 6$ (or $3N - 5$ for a linear molecule) internal coordinates. For simplicity of notation, hereafter we will consider always the case of a nonlinear molecule, so we will have $3N - 6$ internal degrees of freedom and six external ones. The internal coordinates describe the vibrational motion of the molecule and the external coordinates, the rotational and translational motion. The separation between an intramolecular term and a roto-translational term of motion, that is straightforward for the potential, is more complex for the kinetic part. In 1935, Carl Eckart showed, considering a body-
fixed reference frame which rotates and translates with the molecule, that the factorization of the nuclear wave function in a vibrational and rovibrational one is never exact, as there are always nonvanishing vibration–rotation coupling terms. However, there are some conditions, known as Eckart conditions, or Sayvetz conditions, which minimize these coupling terms. A nuclear kinetic operator that includes these coupling terms was obtained by Wilson and Howard in 1936 and refined by Darling and Dennison in 1940, until in 1968 Watson simplified it to obtain the form generally used nowadays, which is referred to as the Watson kinetic operator. In the following sections, we first review the harmonic approximation, where the coupling terms are completely neglected, and later a second order perturbation theory calculation, where coupling terms appear, coming from the simplified Watson form of the kinetic energy operator.

A molecule can usually be assumed to be semirigid; therefore its potential energy is proficiently written in terms of displacements from the equilibrium structure, corresponding to the (local) minimum of the PES. Given the equilibrium structure in Cartesian coordinates \( \mathbf{R}_0 \), the Cartesian displacement is \( \mathbf{X} = \mathbf{R} - \mathbf{R}_0 \).

Often it is more convenient to use internal coordinates \( \mathbf{S} \) (e.g., valence coordinates: bond lengths, angles, and dihedral angles) in place of the Cartesian ones. Indicating with \( \mathbf{S}_0 \) the internal coordinates for the equilibrium structure that correspond to \( \mathbf{R}_0 \) a generic displacement from equilibrium in internal coordinates is \( \mathbf{S} = \mathbf{S} - \mathbf{S}_0 \). A mapping between the Cartesian displacements \( \mathbf{X} \) and the internal coordinates \( \mathbf{S} \) exists, but in general this relation is linear only in the first approximation, since generally internal coordinates are curvilinear. Close to the equilibrium structure, we can Taylor expand \( \mathbf{S} \):

\[
S_i = B^a_i X_a + \frac{1}{2} B^{ab}_i X_a X_b + \frac{1}{3!} B^{abc}_i X_a X_b X_c + \ldots
\]  

(1)

where \( i = 1, \ldots, (3N - 6) \) indexes the internal coordinates; \( a, b, c = 1, \ldots, 3N \) index the Cartesian coordinates (i.e., components \( x, y, z \) for each atom). The Einstein summation notation of repeated indices is assumed hereafter, unless otherwise indicated or if in an expression summation terms are explicitly written. The coefficients \( B^a_i \) appearing in the linear term coincide with the well-known Wilson B matrix.

The potential energy function for the nuclei in the proximity of the equilibrium structure is well described as a Taylor expansion in terms of the internal displacement coordinates:

\[
V(\mathbf{S}) = F + F^H S + \frac{1}{2!} F^{Hi} S_i S = + \frac{1}{3!} F^{ijk} S_i S_j S_k + \ldots
\]

(2)

where the coefficients in the expansion are defined as \( F \equiv V, F^H \equiv \langle \partial V / \partial S \rangle, F^{Hi} \equiv \langle \partial^2 V / \partial S_i \partial S \rangle, \) etc., all calculated in \( S = 0 \). Clearly, if the configuration corresponding to \( S = 0 \), i.e., the reference structure, is a minimum for the potential energy, the coefficients \( F \) are all zero for the Euler conditions. The coefficient \( F \) represents the (local) minimum value of the potential energy, and it is an irrelevant offset for any vibrational calculation. For this reason, it is usually ignored, considering implicitly \( V(S) - F \) in place of \( V(S) \).

2.2. Harmonic Approximation and Normal Modes.

The vibrational normal modes calculation in harmonic approximation is a standard approach, well-known in the literature and implemented in most quantum chemistry codes. Here, we will summarize the main results, which we will use below.

If it is assumed that the reference structure is a minimum for the PES, the linear term in the Taylor expansion of \( V \) is zero. The main approximation in the harmonic approach is that only the leading terms, i.e., the quadratic (harmonic) ones, are considered in both the potential and kinetic energy, indicated hereafter by \( V_{har} \) and \( T_{har} \) and all remaining terms are neglected. In internal coordinates, the approximated potential and kinetic energy are respectively given by

\[
V_{har}(\mathbf{S}) = \frac{1}{2} F^{ij} S_i S_j
\]

(3)

\[
T_{har}(\mathbf{S}) = \frac{1}{2} (G^{-1})^{ij} \dot{S}_i \dot{S}_j
\]

(4)

where \( \dot{S}_i \) is the time derivative of \( S_i \) and the \((3N - 6) \times (3N - 6)\) matrix \( G \) is calculated, according to Wilson, as

\[
G^{ij} = \sum_{\xi} \sum_{\alpha} \frac{1}{M_\xi} B^{i\alpha}_\xi B^{j\alpha}_\xi
\]

the coefficients \( B^{i\alpha}_\xi \) being the same as those that appear in the linear term of eq 1, upon remapping of the indexes from \((\xi, \alpha)\) to \( a \).

Introducing the \((3N - 6)\) normal coordinates \( Q_r \) it is possible to express both the potential and kinetic energies in diagonal form:

\[
V_{har}(\mathbf{Q}) = \frac{1}{2} \sum_r \lambda_r (Q_r)^2
\]

(5)

\[
T_{har}(\mathbf{Q}) = \frac{1}{2} \sum_r (Q_r)^2
\]

(6)

where the coefficients \( \lambda_r = 4\pi^2 c^2 \omega_r^2 \) are the harmonic force constants that provide the harmonic frequencies of vibration \( \omega_r \). Once written in terms of the normal coordinates, the problem is straightforward both in the classical and in the quantum case, since it factorizes in \((3N - 6)\) independent harmonic oscillators (at least for nondegenerate frequencies, as it is supposed in this work).

Assuming that normal coordinates \( Q_r \) are linearly related with internal coordinates \( S_i \) by the relation \( S_i = L_r Q_r \), substituting in eq 3 and eq 4 and comparing with eq 5 and eq 6 yields the relations for the transformation \( L_r \) which written in matrix form are

\[
L^T \mathbf{F} \mathbf{L} = \Phi \text{ and } L^T \mathbf{G}^{-1} L = \mathbf{I}
\]

(7)

where \( \Phi \) is a diagonal matrix with elements of the harmonic force constants \( \lambda_r \), \( \mathbf{I} \) is the \((3N - 6)\) dimensional unit matrix, and superscript symbol + indicates the transpose matrix. The second equation implies that \( \mathbf{L}^T \mathbf{L} = \mathbf{G}^{-1} \), which substituted in the first condition yields \( \mathbf{GFL} = \mathbf{1} \Phi \). Clearly, if we are only interested in the harmonic frequencies, it is enough to solve the secular equation: \( \det(\mathbf{G} - \mathbf{A}) = 0 \).

2.3. Anharmonic Corrections by Second Order Perturbation Theory.

Using second order perturbation
theory (PT2)\textsuperscript{16,17,70–72} it is possible to go beyond the harmonic approximation and to take into account anharmonic effects, expanding the PES to the third and fourth order terms.

The first point in the approach is to consider the fourth order potential energy expansion written in terms of normal coordinates \( Q \):

\[
V(Q) = \frac{1}{2!} \Phi^{ii} Q_i Q_i + \frac{1}{3!} \Phi^{iij} Q_i Q_j Q_j + \frac{1}{4!} \Phi^{ijkl} Q_i Q_j Q_k Q_l
\]

(8)

where we recognize in \( \Phi^{ij} = \lambda \delta_{ij} \) the harmonic force constants already introduced in the previous section. The coefficients \( \Phi^{iij} \equiv \frac{\partial^2 V}{\partial Q_i \partial Q_i \partial Q_j} \) and \( \Phi^{ijkl} \equiv \frac{\partial^4 V}{\partial Q_i \partial Q_j \partial Q_k \partial Q_l} \) are the anharmonic force constants, respectively of third and fourth order.

Assuming that we have fitted the PES in terms of the internal displacement coordinates \( S \) around the minimum structure, obtaining the values of the tensors \( F \) that appear in eq 2 (the zeroth and first order terms are clearly irrelevant), we need to map these values into the tensors \( \Phi \) that appear in eq 8. To do this, we need to consider the transformation from internal curvilinear coordinates \( S \) to normal rectilinear \( Q \). Differently from what we did in the harmonic approach, here we must consider that this relation is not linear, since the internal coordinates are curvilinear. We can consider the following Taylor expansion of \( S \) in terms of \( Q \):

\[
S_i = L_i' Q_i + \frac{1}{2!} L_i'' Q_i Q_i + \frac{1}{3!} L_i''' Q_i Q_i Q_i + \ldots
\]

(9)

where the transformation elements are called \( L \) tensor elements.\textsuperscript{70} The linear terms \( L_i' \) coincide with the matrix \( L \) in eq 7, which solves the secular equation in the harmonic calculation. Substitution of eq 9 in eq 2 and comparison with eq 8 provides the relation between the \( \Phi \) coefficients and the \( F \) ones:

\[
\Phi^{ii} = F^{ii} L_i'
\]

(10)

\[
\Phi^{iij} = F^{iij} L_i' L_j' + F^{iij} (L_i'' L_j' + L_i' L_j'' + L_i''' L_j''')
\]

(11)

\[
\Phi^{ijkl} = F^{ijkl} L_i' L_j' L_k' + F^{ijkl} (L_i'' L_j' L_k'' + L_i'' L_j'' L_k' + L_i'' L_j''' L_k''') + F^{ijkl} (L_i''' L_j'' L_k'' + L_i''' L_j'' L_k''')
\]

(12)

in agreement with ref 70. The \( L \) tensor elements can be determined using the following relations:

\[
L_i' = \frac{\partial S_i}{\partial Q_i} = \frac{\partial S_j}{\partial Q_i} \frac{\partial Q_j}{\partial Q_i} = B_i^h H_i^h
\]

(13)

\[
L_i'' = \frac{\partial^2 S_i}{\partial Q_i \partial Q_i} = \frac{\partial^2 S_j}{\partial Q_i \partial Q_i} \frac{\partial Q_j}{\partial Q_i} = B_i^{ab} H_i^a H_i^b
\]

(14)

The second order perturbative calculation provides the coefficients in the expansion for the molecular vibrational energy:

\[
\hat{H} = \frac{1}{2} \sum_r \hat{h}_r^2 + \frac{\lambda}{2} \sum_r \hat{h}_r^2 + \frac{1}{6} \sum_{rsl} \Phi^{iij} \hat{Q}_i \hat{Q}_j \hat{Q}_l + \frac{1}{24} \sum_{rsstu} \Phi^{ijkl} \hat{Q}_i \hat{Q}_j \hat{Q}_k \hat{Q}_l + \sum_{\alpha} B_{\alpha \beta}^2
\]

(16)

where the coefficients \( H_i^\alpha \equiv (\partial V)/\partial (Q_i) \) can be calculated, using a matrix notation, as \( H = M'B(L^{-1})^\alpha \), and the tensors \( B_{\alpha \beta}^a \) and \( B_{\alpha \beta}^{abc} \) can be obtained from \( B \) by numerical differentiation.

Once all the coefficients of the expansion eq 8 are determined, we consider the Watson Hamilton operator up to the fourth order of expansion in normal coordinates \( Q \), that is\textsuperscript{15,71–72} (the Einstein convention is not used from here to the end of the section):

\[
\hat{H} = \frac{1}{2} \sum_r \hat{h}_r^2 + \frac{\lambda}{2} \sum_r \hat{h}_r^2 + \frac{1}{6} \sum_{rsl} \Phi^{iij} \hat{Q}_i \hat{Q}_j \hat{Q}_l + \frac{1}{24} \sum_{rsstu} \Phi^{ijkl} \hat{Q}_i \hat{Q}_j \hat{Q}_k \hat{Q}_l + \sum_{\alpha} B_{\alpha \beta}^2
\]

(17)

In these expressions, \( \hat{Q}_r \) is the operator relative to the \( r \)th normal coordinate, and \( \hat{p}_r \equiv -i \hbar \partial/\partial (Q_r) \) is the corresponding momentum operator, conjugate to \( Q_r \); \( \alpha = 1, 2, 3 \) represents the three rotational axes in the Eckart frame; \( B_{\alpha \beta} = 1/(2 \lambda_{\alpha \beta}) \) the corresponding equilibrium rotational constants, \( \lambda_{\alpha \beta} \) being the principal moments of inertia; \( \zeta_{\alpha \beta} \) represents the Coriolis coupling constants.

The secondorder perturbative calculation provides the coefficients in the expansion for the molecular vibrational energy:

\[
G(n) = \chi_0 + \sum_{\alpha} a_{\alpha} \left( n_{\alpha} + \frac{1}{2} \right) + \sum_{\alpha < \beta} \chi_{\alpha \beta} \left( n_{\alpha} + \frac{1}{2} \right) \left( n_{\beta} + \frac{1}{2} \right)
\]

(18)

where \( n = \{ n_{\alpha} \} \) are the quantum numbers associated with the \( \alpha \) vibration.

The anharmonic coefficients \( \chi_{\alpha \beta} \) are calculated as\textsuperscript{71}

\[
\chi_{\alpha \beta} = \frac{1}{16} a_{\alpha} a_{\beta} - \frac{1}{16} \sum_{r=1}^{3N-6} \Phi_{\alpha \beta r}^2 \left( a_{\alpha}^2 - a_{\beta}^2 - 3a_{\alpha}^2 \right)
\]

(19)

and for \( r \neq \beta \)

\[
\chi_{\alpha \beta} = \frac{1}{4} a_{\alpha} a_{\beta} \left[ \frac{1}{4} \sum_{r=1}^{3N-6} \Phi_{\alpha \beta r}^2 a_{\alpha}^2 - \frac{1}{2} \sum_{r=1}^{3N-6} \Phi_{\alpha \beta r}^2 \left( a_{\alpha}^2 + a_{\beta}^2 - a_{\alpha}^2 \right) \right]
\]

(20)
no degenerate harmonic frequencies and no quantum-mechanical resonances; for details on these specific cases, see for instance refs 16, 17, and 70.

Equation 18 yields the expressions for the fundamental frequency $\nu_i$:

$$\nu_i = \omega_i + \Delta_i = \omega_i + 2\chi_{rr} + \frac{1}{2} \sum_{s \neq r} \chi_{rs}$$

(21)

overtones:

$$[2\nu_i] = 2\omega_i + 6\chi_{rr} + \sum_{s \neq r} \chi_{rs} = 2\nu_i + 2\chi_{rr}$$

(22)

and combination bands:

$$[\nu_i + \nu_j] = \omega_i + \omega_j + 2\chi_{rr} + 2\chi_{rr} + \frac{1}{2} \sum_{k \neq r, s} (\chi_{rk} + \chi_{rk})$$

$$= \nu_i + \nu_j + \chi_{rs}$$

(23)

3. FIT OF THE PES BY ERROR AFFECTED MEASURES

3.1. Difficulties Due to Error Affected Measures. The methodology introduced in the previous sections for the calculation of vibrational frequencies requires tight (for the harmonic approach) or very tight (for PT2) geometry optimization criteria, namely the configuration at minimum should have a residual gradient smaller than $10^{-5}$ or $10^{-7}$ au, respectively. The consequence of an inaccurate knowledge of the structural minimum would indeed propagate to the frequencies, yielding unreliable results, especially for PT2 calculations. These tight optimization criteria are easily achieved using most common electronic structure methods like the self-consistent field (SCF) cycles used in DFT and post-Hartree–Fock methods.

Unfortunately, in QMC calculations the situation is drastically different, because every QMC measure is affected by a stochastic error $\sigma$ that is inversely proportional to the square root of the number of samplings $N_s$ that in turn is proportional to the computational time required for the calculation. This means that to reduce the error by a factor of 10, a computational effort 100 times larger would be necessary. Oppositely, in a SCF calculation this would just cost a few additional SCF iterations. For these reasons, QMC stochastic errors on energies are typically much larger than what is desirable for vibrational frequency calculations, being, for small molecules, on the order of $\sigma_E \sim 10^{-4}$ au for the energy evaluation and $\sigma_E \sim 10^{-3}$ au for each force component. It is important to point out that, thanks to recent developments in QMC obtained by Sorella and Capriotti, the calculation of all force components is only 4 times more computationally demanding than a single energy calculation.

Before introducing the formalism of the proposed fitting procedure, we discuss some general issues about the localization of a minimum using an error-affected PES. Let us suppose we would like to locate the minimum of the PES having information on energies affected by a stochastic error. From a structural point of view, if we want to find the distance $\Delta$ from the minimum in a potential that is approximatively $\propto \Delta^2$ using a single point estimation of the energy $E$ affected by an error $\sigma_E$, it leads to $\Delta \propto (2E/k)^{1/2}$ being affected by an error $\sigma_\Delta \sim \sigma_E/(k\Delta)$. As an example, considering the OH bond stretching force constant $k \sim 0.5$ au, with an error on the energy of $\sigma_E \sim 10^{-4}$ au, it is clear that it is not possible to estimate $\Delta$ with an accuracy larger than $\sim 1.4 \times 10^{-2}$ au. The situation is even worse for angles, since the bond bending force constants are typically 1 order of magnitude lower than the stretching constants. The computational effort to resolve the position difference $\Delta$ is therefore proportional to $1/(k^2\Delta^4)$, which clearly becomes prohibitive for a tight ($\Delta$) structural optimization.

A different situation comes out if the information on forces, and not only on energies, is known. Close to the minimum the force is approximatively $F = -k\Delta$; therefore, $\Delta$ can be calculated from a single point evaluation of the force as $\Delta = -F/k$, and the associated error is $\sigma_\Delta = \sigma_F/k$. Considering again the previous example of the OH bond stretching, with a typical $\sigma_F \sim 10^{-3}$ au, we have an accuracy on $\Delta$ of the order of $2 \times 10^{-3}$ au, not enough for a tight optimization but much better than the previous energy-based estimation. In this case, the computational effort to resolve a difference $\Delta$ using forces is therefore proportional to $1/(k^2\Delta^2)$, much more convenient than the case of energy.

3.2. Multidimensional Fit of PES. Taking into account the previous considerations, in our approach the equilibrium position and the other parameters of the PES near a minimum are calculated using a multidimensional fit of independent data points (energies or forces) calculated on a grid around an initial guess of the structural minimum of the molecule.

The chosen parametric function $V_0$ that describes the PES in the proximity of the minimum $S_0$ is the Taylor expansion of the potential, see eq 2, cut to the fourth order terms. Therefore, the parameters to fit are $K = \{S_0,F,F'=P_{0i},P_{0j},F_{ij}^{kl}\}$, where the first order coefficients $F'$ are not included, as they have to be zero in the minimum. As a result of the fitting procedure, we will have an estimate of the minimum position $S_0$, the Hessian matrix $F'$, and the third and fourth order coefficients for the PT2 calculation of the anharmonic corrections.

The fitting function $V_0$ presents a nonlinear dependence on the parameters of the structural minimum $S_0$. As will be clear later on, for technical reasons, it would be more convenient to use a linearized form of this function instead of all of its parameters. In the following, we therefore introduce a self-consistent approach, which is easier to implement but is completely equivalent. Let us consider a different fitting function $V$ that represents the fourth order expansion of the PES around a structure $S_0$ which is only a guess of the exact minimum. As a consequence of this, the linear coefficients $F'$ in the expansion cannot be assumed equal to zero any more; in fact, they will be as small as $S_0$ is close to $S_0$. Therefore, we obtain the equation:

$$V(S_0, k_G) = F_0 + F_0 S_G^0 + \frac{1}{2} F_0 S_G^0 S_G^0 + \frac{1}{6} F_0 S_G^0 S_G^0 S_G^0 + \frac{1}{24} F_0 S_G^0 S_G^0 S_G^0 S_G^0$$

(24)

that is linearly parametrized by $k_G = \{F_0,F_0,F_0,F_0,F_0,F_0,F_0\}$ and functionally depends on the displacement $S_G = S - S_0$ from the guess configuration.

The fitted coefficients $k_G$ are relative to the guess structure $S_0$ and can not be used directly for the harmonic and PT2 calculation. However, once we have fitted the function, it is easy to obtain a new guess $S_{G(i)}$ of the minimum configuration, just finding the configuration that has the gradient of the fitted potential equal to zero, and then to repeat the fitting process of $V(S_{G(i)}, k_{G(i)})$ with respect to this new guess structure $S_{G(i)}$. ...
obtaining a new set of coefficients $k_{G(i)}$. Since the fitting process is computationally very fast, we can easily iterate this procedure a sufficiently large number of times, converging at the end to the equilibrium configuration $\bar{S}_{G(i)} \rightarrow \bar{S}_f$ and obtaining the fitting parameters $k_{G(i)} \rightarrow k_f$ that represent the coefficients around the real minimum.

At each iteration $n$, we have to identify the configuration that minimizes the gradient, in order to obtain the new guess structure $\bar{S}_{G(i+1)}$. The exact solution is not so straightforward, since the gradient of the potential has terms of second and third order in the displacement; however, within the described iterative scheme the solution to the leading linear order is enough to ensure the above-mentioned convergence. Indicating with $H_{G(i)}$ the matrix formed by the elements of $F^T_{G(i)}$ and with $F_{G(i)}$ the vector given by $F_{G(i)}$, the approximated condition of zero gradient is written as $F_{G(i)} + H_{G(i)}(\bar{S}_{G(i+1)} - \bar{S}_{G(i)}) = 0$ which yields

$$\bar{S}_{G(i+1)} = \bar{S}_{G(i)} - H_{G(i)}^{-1}F_{G(i)}$$

$H_{G(i)}^{-1}$ being the Moore-Penrose pseudoinverse of $H_{G(i)}$. In our calculations, we have seen that a few ($n \sim 3$) iterations are enough to obtain a tight convergence (i.e., residual forces $F_{G(i)} < 10^{-10}$ au). In agreement with eq 25, zero residual forces means that we can confidently take $\bar{S}_{G(i)}$ as the minimum structure.

In this work, we report several results, namely the coefficients of the PES, the harmonic frequencies and the anharmonic corrections, that depend on the error affected measurements of energies and forces. The error propagation on the results has been obtained using a standard resampling method, namely the Jacknife variance estimator. In the following two subsections, we will see how the values of the fitting parameters are inferred using, respectively, error affected energy and force measures. For simplicity of notation, the subscript $G$ will not be reported, as it is clear that the explanation concerns a single step in the iteration, with displacements $S$ and fitting values $k$ relative to a particular guess structure.

### 3.3. Fit of PES Using Energy Measures

We want to show here how to determine the values of the parameters $k$ of the function $V(S,k)$ that fit the error-affect energy measures around the minimum. Consider a set of $N_f$ configurations ($S^a$), for $a = 1, ..., N_f$. For each configuration $S^a$, the calculations provide a measure of the potential energy $\bar{V}(S^a) \equiv \bar{V}_a$ and a corresponding stochastic error $\sigma_a(S^a) \equiv \sigma_a$

The likelihood that a particular $k$ fits the QMC energy measures $D_E = \{S^a, \bar{V}_a, \sigma_a\}_{a=1,...,N_f}$ is given by the function

$$L(k|D_E) = p(D_E | k) = \prod_a^{N_f} \frac{1}{\sqrt{2\pi \sigma_a^2}} \exp\left(-\frac{[V(S^a, k) - \bar{V}_a]^2}{2\sigma_a^2}\right)$$

where we have assumed that independent estimations of the energy at each configuration are normally distributed. Actually for VMC energy estimations, using the VMC standard sampling technique, this is true only in the limit of infinite sampling, while for finite sampling heavy-tails coming from the local energy singularities are observable as reported by Trail. The estimation of these leptokurtotic tails has unknown bias and should be evaluated for each individual case, and their magnitude can be reduced increasing the VMC sampling. We have explicitly verified that in our particular case the sampling we have used is enough to make negligible the deviations from a normal distribution. For the sake of completeness, it should be addressed that either if a reweighting sampling technique is used, the probability density distribution for the estimated energy would not be normal.

It is not necessary for our purposes to obtain the complete distribution of the likelihood function in terms of $k$, but it is enough to find the particular value of $k_M$ that maximizes the likelihood. It is convenient to consider the logarithm of eq 26, using the fact that the logarithm is a monotone transformation, and after dropping out some irrelevant terms it yields that $k_M$ is the minimum of the chi-squared function:

$$\chi^2(k) = \sum_a^{N_f} \left(\frac{V(S^a, k) - \bar{V}_a}{\sigma_a}\right)^2$$

A remarkable property of the used fitting parameters $k = \{F, F_i, F_{ip}, F_{iop}\}$ is that the fitting function depends linearly on them, therefore

$$V(S, k) \equiv \sum_p^N k_p V_p(S)$$

where $p$ is an index running over all the $N_f$ parameters $k$, and $V_p(S) \equiv (\partial V(S,k))/\partial k_p$ is not functionally dependent on $k$.

By substituting eq 28 in eq 26 it can be seen that the stationary condition $\partial V(S,k)/\partial k_p = 0$ is equivalent to solving $A_k M - B = 0$, where the elements of the $N_f$ dimensional square matrix $A$ are

$$A_{pq} = \sum_a^N \left(\sigma_a^2\right)^{-1} V_p(S^a) V_q(S^a)$$

and the $N_f$ dimensional vector $B$ is

$$B_p = \sum_a^N \left(\sigma_a^2\right)^{-1} V_p(S^a) \bar{V}_a$$

It follows that the fitting parameters are $k_M = A^{-1}B$, where $A^{-1}$ is the Moore-Penrose pseudoinverse of matrix $A$.

### 3.4. Fit of PES Using Force Measures

The methodology used for fitting the PES using force measures is very similar to the one used for the energy, but there is a little complication due to the fact that the forces come from the derivative of the potential with respect to the Cartesian coordinates $R$, not the curvilinear coordinates $S$ used for the fitting function $V(S,k)$.

There is a many-to-one mapping between the Cartesian coordinates $R$ and the corresponding curvilinear coordinates $S = S(R)$, because whereas, given $R$, the internal coordinates $S(R)$ are univocally defined, the opposite is not true due to the arbitrariness of the choice for $R(S)$ of the global orientation of the molecule. Since our QMC code works in Cartesian coordinates, we have decided to use $R$ also for this fitting. It follows that the Cartesian force fitting function of component $a$, with $a = 1, ..., 3N_i$, is

$$F^a(R, \tilde{k}) = -\frac{\partial V(S(R), \tilde{k})}{\partial R_a}$$

$$= -\sum_i^{3N-6} B^a_i \partial \bar{S}_i$$

where $B^a_i(R) \equiv (\partial S_i(R))/\partial R_a$ is given by the Wilson $B$ matrix, calculated for the reference position $R$. The fitting parameters $\tilde{k} = \{F, F_i, F_{ip}, F_{iop}\}$ differ from set $k$ in the energy fitting only for the absence of the $F_i$ that has been dropped out in the
derivative (see eq 31). The number of parameters $\hat{k}$ in the fit with the force is therefore $N_p - 1$, indexed with $p = 2, ..., N_p$. It is clear from eqs 28 and 31 that $\mathcal{F}_{x}(\mathbf{R}, \hat{k})$ is a linear function of $\hat{k}$; therefore, we can write

$$
\mathcal{F}_{x}(\mathbf{R}, \hat{k}) = \sum_{p=2}^{N_p} k^p \mathcal{F}_{x}^p(\mathbf{R})
$$

where the terms $\mathcal{F}_{x}^p(\mathbf{R}) \equiv (\partial \mathcal{F}_{x}^p(S(\mathbf{R}), \hat{k}))/\partial k^p$ are all independent from the parameters $k$, and by substituting in eq 31, these terms can be easily calculated as $\mathcal{F}_{x}^p(\mathbf{R}) = -B_{x}^{p}(\mathbf{R}) \partial V_x(S(\mathbf{R}))/\partial k^p$.

For the fitting, we consider a set of $N_m$ configurations $\{\mathbf{R}^a\}$, for $a = 1, ..., N_m$, and for each configuration we have an associated force measure $\tilde{F}_{x}(\mathbf{R}^a) \equiv \tilde{F}_{x}^a$ with an associated stochastic error $\sigma_{a}(\mathbf{R}^a) \equiv a^p, a = 1, ..., 3N$ being the component index. When the force components come from QMC calculations, they are usually correlated since they are derived by the same stochastic random walks. In this case also their $3N$-dimensional variance-covariance matrix $C_{a}$, which has on the $a$th diagonal elements the square of $\sigma_{a}$ can be calculated.

The likelihood function for the parameters $\hat{k}$ to fit the QMC measures $D_F = \{\mathbf{R}^a \tilde{F}_x^a, \mathbf{C}_a\}_{a=1,...,N_m}$ is given by the function

$$
I(\mathbf{k}|D_F) = p(D_F|\hat{k}) = \prod_{a} \frac{1}{(2\pi)^{3N/2} \sqrt{\text{det}(C_a)}} \exp\left\{-\frac{1}{2} \sum_{a,b} (C_a^{-1})_{ab} [\mathcal{F}^a(\mathbf{R}^a, \hat{k}) - \tilde{F}_x^a]^2 \right\}
$$

as follows from the definition of likelihood, having assumed a normal distribution for the QMC force estimates. In some cases, this assumption could be not exact, depending on the method adopted for the calculation of the forces and the finite VMC sampling. In particular for the calculations reported in this paper, we have verified that the deviations of the force estimations from a normal distribution are negligible.

In the present work, we did not estimate the full variance-covariance matrix but only the diagonal part. Therefore, afterwords we will neglect the correlation between the force components, and we will use only the standard deviations $\sigma_{a}$ of each force component.

As in the case of the energy fitting, we only need to minimize the likelihood function that is equivalent to finding the $\hat{k}$ that minimizes the $\chi^2$ function:

$$
\chi^2(\hat{k}) = \sum_{a} \sum_{a} \left( \frac{\mathcal{F}^a(\mathbf{R}^a, \hat{k}) - \tilde{F}_x^a}{\sigma_{a}} \right)^2
$$

Again the stationary configuration is given by the condition $(\partial \chi^2)/\partial k^p = 0$, which is equivalent to solving $A \hat{k} = \hat{B}$, where the elements of the $N_p$ - 1-dimensional square matrix $A$ are

$$
A_{pq} = \sum_{a} \sum_{a} (\sigma_{a})^{-2} \mathcal{F}_{x}^p(\mathbf{R}^a) \mathcal{F}_{x}^q(\mathbf{R}^a)
$$

and the $N_p$-1-dimensional vector $\hat{B}$ is

$$
\hat{B}_p = \sum_{a} \sum_{a} (\sigma_{a})^{-2} \mathcal{F}_{x}^p(\mathbf{R}^a) \tilde{F}_x^a
$$

for $p, q = 2, ..., N_p$. It follows that $\hat{k} = A^{-1} \hat{B}$, where $A^{-1}$ is the Moore-Penrose pseudoinverse of matrix $A$.

### 3.5. The Choice of the Fitting Grid

The choice of the grid points where energies and forces are evaluated is a crucial aspect for the proposed approach. Clearly the larger is the region of the PES explored by energy or force measurements, the lower is the influence of the stochastic errors on the parameters of the function fitting the PES, and ultimately on the equilibrium structure and the vibrational frequencies. On the other hand, the parametric fitting function is an expansion where we are neglecting all the terms beyond the fourth order expansion; therefore the larger the region of the PES expansion, the lower the accuracy of the fitting function and the larger the systematic error. This implies that on one hand we want to consider the larger mesh size in order to reduce the effect of the stochastic errors on the fitted parameters, but on the other hand the mesh cannot be too large; otherwise we would introduce a systematic error due to an inaccurate fitting function in the range where the QMC measures were obtained.

To study these effects for typical values of stochastic errors we studied the behavior of the systematic and stochastic errors using grids of different sizes and shapes. We considered as a model system for these tests the DFT PES of the water molecule where stochastic errors were introduced artificially on forces and energies. In this way, we can directly compare the parameters obtained with the fitting procedure with the exact parameters of the PES to directly estimate the effect of different grids. In the specific case of the water molecule, the grid points are chosen as follows:

$$
\begin{align*}
\Delta r_1 &= r_0 + \delta r \\
\Delta r_2 &= r_0 + \delta r \\
\Delta \phi &= \phi_0 + \delta \phi
\end{align*}
$$

where the center of the grid, given by $r_0$ and $\phi_0$ is our initial guess of the equilibrium structure. The integers $i$, $j$, and $k$ are so that $i | j | k \leq N$; $i \leq j$, and if $i = j$, then $i | j | k \leq N_2$. Otherwise, $i | j | k \leq N$. Therefore each mesh is specified by the parameters $N_1, N_2, N_3, \Delta r$, and $\Delta \phi$. The set of meshes considered in this paper are reported in Table 1.

<table>
<thead>
<tr>
<th>Mesh</th>
<th>$N_1$</th>
<th>$N_2$</th>
<th>$N_3$</th>
<th>$\Delta r$</th>
<th>$\Delta \phi$</th>
<th>Points</th>
</tr>
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<tr>
<td>mesh-1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>0.02</td>
<td>1°</td>
<td>59</td>
</tr>
<tr>
<td>mesh-2</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>0.03</td>
<td>3°</td>
<td>59</td>
</tr>
<tr>
<td>mesh-3</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>0.04</td>
<td>5°</td>
<td>59</td>
</tr>
<tr>
<td>mesh-4</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>0.08</td>
<td>10°</td>
<td>59</td>
</tr>
<tr>
<td>mesh-5</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td>0.20</td>
<td>20°</td>
<td>59</td>
</tr>
<tr>
<td>mesh-6</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>0.03</td>
<td>3°</td>
<td>77</td>
</tr>
<tr>
<td>mesh-7</td>
<td>4</td>
<td>4</td>
<td>5</td>
<td>0.03</td>
<td>3°</td>
<td>139</td>
</tr>
<tr>
<td>mesh-8</td>
<td>5</td>
<td>5</td>
<td>6</td>
<td>0.03</td>
<td>3°</td>
<td>208</td>
</tr>
</tbody>
</table>

$^a$Mesh types used in this paper that are univocally defined by the values of $N_1, N_2, N_3, \Delta r$, and $\Delta \phi$. For a description of the construction method, see section 3.5. $^b$In atomic units. $^c$In degrees
Table 2. Systematic Errors Induced by the Mesh Size

<table>
<thead>
<tr>
<th>ω</th>
<th>ORCA&lt;sup&gt;b&lt;/sup&gt;</th>
<th>mesh-1</th>
<th>mesh-2</th>
<th>mesh-3</th>
<th>mesh-4</th>
<th>mesh-5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0.02B/1°</td>
<td>0.03B/1°</td>
<td>0.04B/5°</td>
<td>0.08B/10°</td>
<td>0.20B/20°</td>
</tr>
<tr>
<td>ω&lt;sub&gt;2&lt;/sub&gt;</td>
<td>1626.3</td>
<td>1626.7</td>
<td>1626.6</td>
<td>1626.8</td>
<td>1628.9</td>
<td>1635.7</td>
</tr>
<tr>
<td>ω&lt;sub&gt;1&lt;/sub&gt;</td>
<td>3793.4</td>
<td>3793.7</td>
<td>3794.1</td>
<td>3794.5</td>
<td>3798.7</td>
<td>3853.5</td>
</tr>
<tr>
<td>ω&lt;sub&gt;3&lt;/sub&gt;</td>
<td>3896.1</td>
<td>3896.3</td>
<td>3896.7</td>
<td>3897.1</td>
<td>3900.3</td>
<td>3953.6</td>
</tr>
<tr>
<td>X&lt;sub&gt;22&lt;/sub&gt;</td>
<td>−19.53</td>
<td>−19.66</td>
<td>−19.68</td>
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<td>−19.64</td>
</tr>
<tr>
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<td>−13.27</td>
<td>−13.91</td>
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<td></td>
</tr>
<tr>
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<td>−16.75</td>
<td>−21.08</td>
<td></td>
</tr>
<tr>
<td>X&lt;sub&gt;11&lt;/sub&gt;</td>
<td>−44.25</td>
<td>−44.42</td>
<td>−44.83</td>
<td>−47.63</td>
<td>−70.39</td>
<td></td>
</tr>
<tr>
<td>X&lt;sub&gt;13&lt;/sub&gt;</td>
<td>−173.42</td>
<td>−173.96</td>
<td>−175.46</td>
<td>−186.21</td>
<td>−274.72</td>
<td></td>
</tr>
<tr>
<td>X&lt;sub&gt;13&lt;/sub&gt;</td>
<td>−50.31</td>
<td>−50.73</td>
<td>−51.07</td>
<td>−53.90</td>
<td>−77.42</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>Harmonic frequencies ω and anharmonic corrections X (both in cm<sup>−1</sup>) computed for a DFT calculation, for the B3LYP functional and aug-cc-pVTZ basis set. In the first column, the calculation has been performed directly using the ORCA package,<sup>80</sup> while in the remaining columns, we used the fitting of PES with forces in a grid with some different mesh types characterized by the Δr and ΔΦ reported below the mesh number (see details in the text and in Table 1).<sup>33</sup> Anharmonic coefficients are not reported because they are not provided by the ORCA package.

4. COMPUTATIONAL DETAILS

In this work, we have performed both DFT and QMC calculations: the first ones aimed to assess the optimal choice of the grid points around the configurational minimum to take under control both the systematic and the stochastic errors; the second ones aimed to test the validity of the methods when applied to QMC calculations.

DFT calculations were performed using the ORCA package,<sup>80</sup> with the B3LYP functional<sup>81–84</sup> and aug-cc-pVTZ basis set.<sup>85,86</sup> For each grid, we calculate energies and gradients. To simulate the presence of stochastic errors we added to the values of energies and forces random contributions normally distributed with the desired standard deviation σ<sub>n</sub> and σ<sub>f</sub>, respectively, for the energy and the force.

The QMC energy and force calculations have been carried out using the TurboRVB package developed by Sorella<sup>87</sup> that includes a complete suite of variational and diffusion quantum Monte Carlo programs for wave function and geometry optimization of molecules and solids. The wave function used is a Jastrow antisymmetrized geminals power<sup>80</sup> (J-AGP). Since in the present context the use of QMC is purely demonstrative of the feasibility of the PES fitting procedure, we used a minimal wave function, in order to minimize the computational costs. The orbital basis set for the AGP part is composed of [4s,4p,1d] Gaussian type orbitals (GTO) contracted in (2s,2p,1d) for the oxygen atom and of [4s,1p] GTO orbitals contracted in (2s,1p) for the hydrogen atoms. The exponent values were taken from the cc-pVQZ basis set, considering only values larger than 20 au. The Jastrow factor used here consists of several terms that account for the one-body, two-body, three-body, and four-body interaction between electrons and nuclei, as described in refs 36 and 61. For the three/four-body Jastrow part, we used a GTO basis set with (1s, 1p) uncontracted orbitals for oxygen and (1s) for hydrogen. An energy-consistent pseudopotential<sup>88</sup> was used for the oxygen atom. The wave function optimization was carried out, for each ionic configuration, on the AGP matrix parameters, on the contraction coefficients of the atomic AGP bases, and on all the Jastrow parameters, including exponents. All the QMC calculations here presented have been obtained inside a variational Monte Carlo (VMC) scheme, in which the VMC forces<sup>61</sup> can be computed very efficiently using the adjoint algorithmic differentiation recently developed and implemented by Sorella and Capriotti.<sup>36</sup>

The experimental configuration of the molecule was taken as the initial guess of the PES structural minimum. We calculated the QMC forces for the experimental configuration, using the setup described above, and we observed that a residual force of the order of 10<sup>−3</sup> au is present. This implies that the experimental configuration is close enough to the PES minimum to be taken as the center of the grid for the PES fitting that we are proposing here, but it would not be close enough for a single point evaluation of the Hessian matrix and harmonic frequencies.

5. RESULTS AND DISCUSSION

In this work, we considered as a test case for our approach the water molecule, and specifically the most common H<sub>2</sub>O molecule with <sup>1</sup>H and <sup>16</sup>O isotopes. Water counts for three internal coordinates, or nine Cartesian coordinates. The considered internal coordinates are the two distances r<sub>1</sub> and r<sub>2</sub> between the oxygen and the hydrogens, and the HOH angle is Φ. In the equilibrium configuration experiments<sup>89</sup> reported, r<sub>Exp</sub> = 0.95721(3) Å and Φ<sub>Exp</sub> = 104.522(5)<sup>°</sup>. The art computational results, obtained with coupled-cluster calculations, give equilibrium geometry quite close to the experimental one,<sup>90</sup> while an overview of the results that can be obtained by various quantum chemical computations are reported in ref 91. The numbers of independent parameters in a fourth order expansion of the potential in internal coordinates, taking into account the symmetry of the system, are 22 for the energy fit and 21 for the force fit. The number of parameters (of energies or forces) has to be larger than the number of parameters in order to have a well-defined fitting problem.

In the following, we will first investigate the problem of the systematics versus stochastic errors using DFT calculations, and finally we will report results obtained for a VMC calculation performed using a small wave function.

In order to investigate the systematic errors induced by the approximated parametrization of the PES, we consider here the grids from mesh-1 to mesh-5 defined in section 3.5 and Table 1. By fitting the forces provided by DFT, we obtained the results reported in Table 2. The comparison with the analytical DFT parameters in the first column shows how for small meshes the computed frequencies and anharmonic corrections are close to the analytical results, but as the mesh size increases they start to differ. This happens because the polynomial expansion is a good approximation of the potential in a not too big neighborhood of the equilibrium configuration. From this
calculation, we can therefore estimate up to which mesh size the systematic errors can be negligible with respect to other sources of error. In particular, for mesh-4 the systematic errors are less than \( \sim 6 \text{ cm}^{-1} \), i.e. \( \sim 0.2\% \), on the frequencies and less than \( \sim 13 \text{ cm}^{-1} \) on the anharmonic coefficients.

5.1. Stochastic Errors. To address the effects of the propagation of the stochastic error, we consider again the grids from mesh-1 to mesh-5. We use the DFT forces previously computed, and to simulate the stochastic errors of the QMC forces we add to each DFT force component a pseudorandom number normally distributed and with a variance \( \sigma^2 \). The results obtained fitting these forces are reported in Figures 1 and 2, au, Figures 1 and 2 and Table 2 suggest that mesh-4 has the best balance between stochastic and systematic errors.

Once it is established which is the optimal size of the grid, we can address here the importance of the way the points are distributed on the grid (the grid type) and of the influence of the total number of points on the grid. For this purpose, we consider the grids: mesh-6, mesh-7, and mesh-8, as defined in Table 1. The results of the fitting procedure of the forces are reported in Tables 3 and 4 respectively for the harmonic frequencies \( \omega \) and the anharmonic corrections \( X \). We may observe, by comparison with Table 2, that the systematic errors are under control in all cases, being \( \leq 2 \text{ cm}^{-1} \) for the harmonic frequencies and \( \leq 5 \text{ cm}^{-1} \) for the anharmonic corrections.

We would like now to compare different strategies for decreasing the stochastic errors in our computations, having a certain fixed amount of computer time. Let \( t \) be the computational time required to obtain certain stochastic errors of the harmonic frequencies \( \sigma_\omega N \) be the number of points in the grid, and \( n_s \) be the number of Monte Carlo samples in each single point calculation that provides forces with a stochastic error of \( \sigma_f \). We have that \( \sigma_\omega \propto \sigma_f / \sqrt{N} \) and \( \sigma_f \propto 1/\sqrt{n_s} \). The total computational time \( t \) is the sum of the time for the sampling \( t_s \propto N^n_s n_s \) and the time for the wave function optimization \( t_{opt} \propto N \). The more we sample the wave function, the more accurate has to be the optimization; therefore we can assume that the time for the optimization is a fraction \( x_{opt} \) of the time for the sampling. It follows that \( t \propto (1 + x_{opt})^n_s N \).

Figure 1. Harmonic frequencies obtained fitting DFT (B3LYP/aug-cc-pVTZ) forces, where random errors normally distributed have been added in order to simulate QMC stochastic errors for different mesh types (details in the text). Several values (labels) of the standard deviation for the force components were tried around the typical QMC ones. For each case, we reported the representative frequency with the corresponding error calculated using the Jackknife resampling method. The dashed line represents the reference results given by the ORCA package that can be considered free from any stochastic and grid error.

Figure 2. Same of the Figure 1, but with the anharmonic force constants reported.
Table 3. Harmonic Frequencies from Error Affected Measures

<table>
<thead>
<tr>
<th>$\sigma_f$</th>
<th>mesh-6</th>
<th>mesh-7</th>
<th>mesh-8</th>
</tr>
</thead>
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<tr>
<td>1.00e-10</td>
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<td>-20(0)</td>
<td>-20(0)</td>
</tr>
<tr>
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<td>-18(5)</td>
<td>-17(3)</td>
</tr>
<tr>
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<td>-11(9)</td>
<td>-18(5)</td>
</tr>
<tr>
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<td>-27(10)</td>
<td>-19(8)</td>
</tr>
<tr>
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<td>-13(0)</td>
<td>-13(0)</td>
</tr>
<tr>
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<td>-23(19)</td>
<td>-10(7)</td>
</tr>
<tr>
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<td>-15(25)</td>
<td>-31(10)</td>
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<td>-7(15)</td>
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Table 4. Anharmonic Constants from Error Affected Measures

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<td>1628(3)</td>
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The same amount of additional computer time can be distributed in different ways, either increasing the length of each stochastic sampling or increasing the number of sampling points. The first strategy is to decrease the stochastic errors by longer Monte Carlo sampling, without adding additional points to the grid. The only quantity that varies is therefore $n_f$. The second strategy is to add new points to the grid, namely changing $N$.

The numerical results in Table 1 are carried out using grids spanning the same configuration region but using a finer ($N = 208$ points), medium ($N = 139$ points), or coarser ($N = 77$ points) mesh. According to our previous considerations, if we take as fixed the $\sigma_f$ for each point calculation, the computational time for a QMC calculation is $t_c = c^*N_f$ when using mesh-6 and $t_c = c^*N_f$ when using mesh-8, $c$ being an opportune constant. Therefore, the expected increment in the computational time is $t_f/t_c \sim 2.7$. The reduction of the stochastic error obtained in this way according to our simulations is $\sim0.31(9)$ for the harmonic frequencies $\omega$, see Table 3, and $\sim0.21(5)$ for the anharmonic constants $\chi$, see Table 4. These results have to be compared with the reduction that we would obtain using the first strategy for the same increment of computational time, that is $(t_f/t_c)^{1/2} \sim 0.6$ both for $\omega$ and $\chi$. In conclusion, the calculations reveal that to reduce the stochastic error on frequencies and anharmonic constants, it is more effective to add additional mesh points rather than increase their accuracy.

To demonstrate the effectiveness of the method in a real albeit simple QMC case, we performed VMC calculations using minimal basis sets for the AGP and Jastrow part, on a mesh-4 grid centered around the experimental structural minimum, i.e., $\beta = 0.9572$ Å and $\phi = 104.52^\circ$. On this structure, we obtained a VMC energy of $-17.2144(6)$ au and residual force components $\leq0.003(1)$ au. The forces calculated in the grid points span from $-0.13$ au to 0.15 au and have an average stochastic error of $\sim0.0009$ au.

In Table 5 are reported the results of two different fits: one using the VMC energies and the other using the VMC forces. It is immediately clear how the quantities obtained from the energy fit have stochastic errors which are at least 1 order of magnitude larger than those obtained by the fit of the forces. This trend is observed both for the evaluation of the geometrical parameters at a minimum and for the harmonic and anharmonic constants. Since the calculations of QMC forces in our scheme requires only 4 times the computational effort of energy evaluations,\textsuperscript{95} it is clear that the force-fitting procedure is always extremely more convenient than the energy-fitting procedure.

Despite the variational wave function used being very small, the equilibrium configuration predicted using the force fit yields $r_{VMC} = 0.9552(2)$ Å and $\phi_{VMC} = 104.11(7)^\circ$, which are in good agreement with the experimental values and perform well in comparison with the other computational methods.\textsuperscript{91} On the other hand, the harmonic frequencies are overestimated if compared to the experimental values, whereas the estimation of the anharmonic constants are rather close. This discrepancy was not unexpected, because the basis sets for the AGP and Jastrow expansion are minimal. A complete basis set study requires much larger computational resources and will be the subject of further investigations. We point out that besides the discrepancy with respect to the experimental results, the statistical errors on the estimated harmonic frequencies are as small as 0.7%, demonstrating that the proposed method is able...
Table 5. QMC Results

<table>
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<th></th>
<th>energy fit</th>
<th>force fit</th>
<th>EXP</th>
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<td>0.95721(3)</td>
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"Equilibrium configuration (OH distance $r$ in Å and angle $\phi$ in degrees), harmonic frequencies $\omega$, anharmonic constants $\Delta \nu$, and fundamental frequencies $\nu$ theoretically computed by VMC energy fitting (first column), VMC force fitting (second column), and experimentally observed by Benedict et al.59 (third column). For all the computational results, the standard mesh $N_1 = 4, N_2 = 2, N_3 = 3$ was used, with $\Delta \rho = 0.08$ au and $\Delta \varphi = 10^\circ$ (see section 3.5). The energy/force was evaluated at each point of the mesh, and the potential expansion up to fourth order was fitted as described in section 3.3 [section 3.4]. The numbers in the brackets are the stochastic errors, estimated by the jackknife algorithm.

"to provide for small molecules a precise evaluation of vibrational frequencies.

6. CONCLUSIONS

In the present work, we proposed a theoretical framework for the calculation of equilibrium structures and vibrational properties through a potential energy surface affected by stochastic noise. The proposed procedure has as natural application the QMC electronic structure calculations, although it is not limited to these methods. We demonstrated that applying a multidimensional fitting procedure to the QMC energies or forces around the minimum, we can obtain accurate results on the structural parameters at the minimum and on the vibrational frequencies, despite the presence of the intrinsic stochastic error of the method. The theoretical developments in this work, as well as the numerical results, are aimed to minimize the computational time and mitigate the impact of the stochastic errors on the accuracy of the computed harmonic frequencies. In the QMC example, we applied our technique to the calculation of the equilibrium structure, harmonic frequencies, and anharmonic constants of the water molecule. The obtained results demonstrated how using information of forces instead of energies it is possible to reduce the statistical error on the vibrational frequencies by at least 1 order of magnitude, leading to an average error lower than 0.07% for the geometrical parameters and 0.7% for the harmonic frequencies. These results open the possibility to apply quantum Monte Carlo methods for the evaluation of vibrational spectroscopic parameters of molecules, although the scaling of the method with the number of atoms in the system limits the approach to relatively small molecules.

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Notes

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