Rotationally Resolved Electronic Spectroscopy and Automatic Assignment Techniques using Evolutionary Algorithms

Michael Schmitt¹ and W. Leo Meerts²

¹Institut für Physikalische Chemie I, Heinrich-Heine-Universität, Düsseldorf, Germany ²Molecular and Biophysics Group, Institute for Molecules and Materials, Radboud University Nijmegen, Nijmegen, The Netherlands

1 INTRODUCTION

The classical assignment of high-resolution spectra is mostly based on pattern recognition by the eye. Over the years, a number of methods and computer programs have been developed that greatly helped the analysis of such spectra. An early attempt in the automated assignment of microwave and infrared spectra is the Loomis-Wood method (Loomis and Wood 1928), which gained wider applicability only after microcomputers became available. The group of Neusser (Helm et al. 1997) has developed the method of correlation automated rotational fitting (CARF), which directly fits the experimental data, without prior assignments. Meerts and Schmitt (2006) give a detailed overview of several of these techniques. Although the computer-aided analyses meant great progress, they still need large operator interference and, in general, are costly in time. Furthermore, quite some experience is required from the spectroscopist and, in most cases, the methods are limited to well-resolved spectra with not too much overlap between the transitions.

In order to be able to analyze and assign high-resolution spectra automatically in a routine manner, we have attacked this problem with the help of evolutionary algorithms (EAs). It turned out to be a very powerful technique that made it possible to assign rotationally resolved ultraviolet (UV) laser spectra with minimum operator interference. The main strength of EAs is their ability to perform global searches of the parameter space with a built-in learning mechanism. The algorithms are based on concepts copied from natural reproduction and selection processes. Inherent to the fact that the EAs are global optimizers, the spectra have to be calculated a large number of times in the evaluation phase. Fortunately, with the fast computers presently available and the possibility to perform vast parallel processing on large cluster machines, computing time is hardly a problem.

The process of automatic assignment of the spectra is actually based on pattern recognition guided by the EAs. Important, but not decisive, is the reliability of the experimental intensities and the presence of a model capable to predict the observed spectra. However, these side conditions can be waived significantly in some cases as has been shown by Meerts and Schmitt (2006).

Over the years, we have analyzed and assigned numerous spectra and it was found that well-resolved spectra with little overlap of spectral lines could be easily managed automatically, fast, and in a routine way. This makes it possible to analyze vast amounts of experimental data and concentrate on the results rather than on the assignments. It was further shown that in the cases where the spectra become more dense and even the overlap of spectral lines becomes very large, the automatic assignment procedure is still successful. Several examples are presented in this article (e.g., Section 5.2). This is of particular interest if large biologically relevant molecules or biomimetics are investigated. In

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the case of the investigation of isotopic species, it often happens that several isotopologs are simultaneously present in the sample, which generally leads to a very dense spectrum due to their small spectral shifts (Schmitt *et al.* 2005).

In general, with each type of spectrum, for which a sufficiently accurate model is available to describe the experimental data, the technique of EA fitting is applicable. Thus, the presented technique is not limited to rotationally resolved electronic spectroscopy, which is the main focus of this article, but can instead be also used for the interpretation of rovibrational spectra in the infrared (IR) (see Albert et al. 2011: High-resolution Fourier Transform Infrared Spectroscopy, Snels et al. 2011: High-resolution FTIR and Diode Laser Spectroscopy of Supersonic Jets, this handbook) or of rotational spectra in the microwave (MW) region (see Jäger and Xu 2011: Fourier Transform Microwave Spectroscopy of Doped Helium Clusters, Caminati 2011: Microwave Spectroscopy of Large Molecules and Molecular Complexes, this handbook). The only precondition in all these cases is the availability of a Hamiltonian, which is capable of a nearly perfect description of the spectra. This precondition is not fulfilled, e.g., for pure vibrational spectra, which in most cases cannot be described with the necessary accuracy, since anharmonic corrections to the calculated harmonic spectra are too involved, to be applied to the large molecules considered here. Also, the application of EAs to the interpretation of MW spectra is sometimes hampered, due to the nonreproducibility of intensities in modern Fourier transform microwave (FTMW) experiments of the Balle-Flygare type (see Jäger and Xu 2011: Fourier Transform Microwave Spectroscopy of Doped Helium Clusters, Grabow 2011: Fourier Transform Microwave Spectroscopy Measurement and Instrumentation, this handbook). Even though intensities are considered to be of minor importance in pattern recognition, they play an important role if the number of lines gets very large.

Toward the end of this article, we show the application of the EA-based fits to the interpretation of a completely different class of spectra, namely, nuclear magnetic resonance (NMR) spectra of solutes in liquid crystal solvents. Also, in this case, a sufficiently accurate Hamiltonian is at hand to reproduce the spectra and despite the large number of parameters to be determined, which make the manual assignment extremely tedious and time consuming, a straightforward solution on the basis of EA could be found.

2 THE CHOICE OF THE APPROPRIATE HAMILTONIAN

The essential starting point for any successful fit of a molecular spectrum is the determination of the correct Hamiltonian that is needed to describe the spectrum. However, even in cases where the complete Hamiltonian is not known or is too time consuming to be used, an overall fit of the spectrum can be performed if the perturbations introduced by the complete Hamiltonian act only on limited ranges of the spectra.

2.1 Rotational, Rovibrational, and Rovibronic Spectra

For most of the spectra presented here, a rigid asymmetric rotor Hamiltonian was employed (Allen and Cross 1963):

$$H_r = AJ_a^2 + BJ_b^2 + CJ_c^2$$
(1)

Here, $J_g(g = a, b, c)$ are the components of the body fixed angular momentum operator and A, B, and C are the three rotational constants. One set of rotational constants is needed for the simulation of microwave spectra, while for rovibrational or rovibronic spectra a second set of rotational constants for the (vibrationally or electronically) excited state is necessary.

Quartic and sextic centrifugal distortion constants may be included if necessary through the model of a distortable rotor in Watson's *A*-reduced form (Watson 1968).

The intensities of the rotational lines are calculated from the eigenvectors of the effective rotational Hamiltonian and the known direction cosine matrix elements (Gordy and Cook 1984). The only selection rule used is $\Delta J = 0, \pm 1$, which distinguishes *P*, *Q*, and *R* branches. The linestrength is proportional to

$$A_{r'r''} \propto |\mu_a \langle r' | \Phi_{Za} | r'' \rangle|^2 + |\mu_b \langle r' | \Phi_{Zb} | r'' \rangle|^2 + |\mu_c \langle r' | \Phi_{Zc} | r'' \rangle|^2$$
(2)

where the Z axis is the space-fixed axis along the direction of the polarization of light, $|r\rangle \equiv |J, K_a, K_c\rangle$ and

$$\mu_{a} = \mu \sin \phi \cos \theta$$

$$\mu_{b} = \mu \sin \phi \sin \theta$$

$$\mu_{c} = \mu \cos \phi$$
(3)

The double and single prime denote the ground and excited state, respectively, μ is the absolute value of the transition dipole moment (TDM), (ϕ , θ) are the spherical coordinate angles of the transition moment vector in the molecular fixed frame (a, b, c), and Φ_{Za} , Φ_{Zb} , and Φ_{Zc} are the direction cosines between the laboratory Z axis and the molecular a, b, and c axes.

In equation (3), θ is the angle between the projection vector of the transition moment on the *ab*-plane, and the

a-axis and ϕ the angle between the TM and the c-axis. In the general case (*abc*-type transition), θ is called the *azimuthal angle* and ϕ the polar angle and the pair is written as (ϕ , θ).

The total intensity is obtained by multiplying the line strength from equation (2) with the temperature-dependent population intensity. For simulation of gas phase spectra in a static cell, the system is in thermal equilibrium and a single temperature will be sufficient to calculate the distribution of rotational states using a Boltzman distribution. In molecular beams, the different J states relax differently depending on the size of J. This behavior can be simulated by a two-temperature model (Wu and Levy 1989):

$$n(T_1, T_2, w_T) = e^{-E/kT_1} + w_T e^{-E/kT_2}$$
(4)

Here E is the energy of the lower state, w_T is a weight factor, and T_1 and T_2 are the two temperatures.

Thus, for a relatively simple molecular beam spectrum with rotational resolution, the number of parameters that have to be determined in the course of the fit is already quite large: six rotational constants (three for each state) A, B, C, eventually 10 centrifugal distortion constants (five for each state), the center frequency v_0 , the two TDM angles θ and ϕ , the two temperatures T_1 and T_2 and a weight factor w_T from equation 4, and at least one line shape factor determining the width of the experimental spectral lines.

Any spectral perturbation which might arise from the coupling of angular momenta of internal motions of parts of the molecule with the overall angular momentum must be taken into account by using different Hamiltonians with a still larger number of parameters to be fit. Other angular momentum coupling schemes comprise Coriolis and Darling-Dennison couplings. Hamiltonians implemented in the fit program until now comprise perturbation from a single hindered three-fold rotation, (Remmers *et al.* 1998) from two three-fold rotors on a common torsional axis, (Tan *et al.* 1995) from a two-fold rotation, (Berden *et al.* 1996) and from axis reorientation (Berden *et al.* 1995). The latter effect is due to a rotation of the inertial axes when the structure of the molecule is distorted upon electronic excitation.

2.2 The NMR Hamiltonian in a Liquid Crystal

The Hamiltonian (in Hertz) of a molecule dissolved in a uniaxial nematic liquid crystal is given by

$$H = \sum_{\mu} \nu_{\mu} I_{Z;\mu} + \sum_{\mu < \nu} J_{\mu\nu} \mathbf{I}_{\mu} \cdot \mathbf{I}_{\nu}$$
$$+ \sum_{\mu < \nu} D_{\mu\nu} (3I_{Z;\mu} I_{Z;\nu} - \mathbf{I}_{\mu} \cdot \mathbf{I}_{\nu})$$
(5)

where ν_{μ} is the resonance frequency of nucleus μ and $J_{\mu\nu}$ and $D_{\mu\nu}$ are the indirect spin-spin and the direct dipole-dipole coupling between nuclei μ and ν . The chemical shielding term ν_{μ} can have both isotropic and anisotropic parts, whereas the $D_{\mu\nu}$ terms are completely anisotropic and hence do not contribute to the familiar spectra of isotropic liquids. We omit effects of the anisotropy in $J_{\mu\nu}$, which is to slightly modify the observed values of $D_{\mu\nu}$ and which is negligible for proton-proton couplings.

The dipolar coupling between nuclei μ and ν is

$$D_{\mu\nu} = -\frac{h\gamma_{\mu}\gamma_{\nu}}{4\pi^2} \left\langle r_{\mu\nu}^{-3} \left(\frac{3}{2} \cos^2 \theta_{\mu\nu,Z} - \frac{1}{2} \right) \right\rangle \tag{6}$$

where $r_{\mu\nu}$ is the internuclear separation and $\theta_{\mu\nu,Z}$ is the angle between the $\mu\nu$ and magnetic field directions, and the angle brackets denote an average over solute internal and reorientational motions.

3 THEORY OF EVOLUTIONARY ALGORITHMS

In the past decades, computational tools have been developed that are able to handle complex multiparameter optimizations intelligently and within an acceptable time interval. The use of EAs for solving such highly nonlinear and complex processes in science and engineering has become widespread. Although they are conceptually simple, their ability to avoid local optima and instead find the global optimum is remarkable and makes them suitable to handle complex optimization problems.

EAs represent a set of general purpose probabilistic search methods based on natural evolution. These algorithms mimic the concepts of natural reproduction and selection processes. The basic idea in EAs is to create an artificial environment which encodes the search problem into biology-like terms. With the proper definition of the genetic operators, the EAs let the artificial environment evolve and find the optimal solution.

Three main different EAs have been developed over the years: the genetic algorithm (GA) by Holland (1975), the evolution strategy (ES) by Rechenberg (1973) and the evolutionary programming by Schwefel (1993).

For the automatic assignment and analysis of the highresolution laser spectra, we have mainly used the GA (Meerts and Schmitt 2006). However, recently we implemented an ES in our procedures and the comparison between GA and ES is discussed in this article.

3.1 The Genetic Algorithm

A detailed description of the GA used in the automatic assignment and fitting of the spectra described in this article can be found in Meerts and Schmitt (2006).

The molecular parameters are encoded in binary or real type, each parameter to be optimized representing a gene. The vector of all genes, which contains all molecular parameters, is called a *chromosome*. In an initial step, the values of all parameters are set to random values between lower and upper limits, which are chosen by the user. The quality of the solutions is then evaluated by a fitness function. A flow chart of the procedure is shown in Figure 1.

One optimization cycle, including evaluation of the fitness of all solutions, is called a *generation*. Pairs of chromosomes are selected for reproduction and their information is combined via a crossover process. Since crossover combines information from the parent generations, it basically explores the fitness landscape. The value of a small number of bits is changed randomly by a mutation operator. Mutation can be viewed as exploration of the fitness surface. The best solutions within a generation are excluded from mutation. This elitism prevents already good solutions from being degraded. Mutation prevents the calculation from being trapped in local minima, as is often the case with more conventional fitting routines.

3.2 The Evolutionary Strategy

The evolution strategy algorithm starts with one or more parent(s). A parent is a trial solution that corresponds to a set of parameters like in the genetic algorithm. From this parent an offspring of multiple children is generated. The quality or performance of these children is checked and depending on the strategy the next parent is generated. There are several different strategies for the generation of the offspring as well as the generation of the next parent. For a more detailed description, see Rechenberg (1973).

The offspring is created from the parent(s) in a mutative step-size control. A drawback of the standard ES is the fact that the mutations of the decision and the strategy parameters, respectively, are subject to independent random processes. If, for example, an individual with a large step size undergoes only a very small change in the decision parameters and this small change turns out to yield a high fitness, the large step size will be inherited to the next generation. As a result, the fitness in the next mutations may be worse. This problem is resolved in derandomized (DR) algorithms, which make the random mutations in decision and strategy parameters dependent on each other. This idea was implemented for the first as DR1 and soon improved by the concept of accumulated information (Ostermeier et al. 1994), which will be called DR2. The history of the optimization is recorded and the evolution of the mutation ellipsoid is partially governed by past successful mutations.

A further improvement was achieved by Hansen and Ostermeier (Hansen and Ostermeier 2001) with the covariance matrix adaptation evolution strategy (CMA-ES). It has proved to be a particularly reliable and highly competitive evolutionary algorithm for local optimization and, surprisingly at first sight, also for global optimization (Hansen and Kern 2004). The CMA-ES does not leave the choice of strategy parameters open to the user—only the population size can be set. Finding good strategy parameters is considered to be part of the algorithm design.



Figure 1 Schematics of the GA-process.



Figure 2 The first four generations of an evolution strategy are as follows: (a) An initial population is generated, and the best offspring is used as the next parent. (b) The offspring is spread over a larger area in the second generation due to the relatively large step made in the previous generation. The vector from the parent to the best offspring (dashed line) is combined with the (shortened) mutation vector of the last generation (dotted line) to generate the new parent (solid line). (c) Owing to the correlation between the past two mutations the search range has been extended again in the general direction of both mutations while it has been limited in the perpendicular direction. The best offspring is now a local minimum. The memory effect of the evolutionary algorithm, which incorporates past mutation vectors into the calculation of the next parent, helps to overcome the local minimum and the next parent is still closer to the global minimum. (d) The barrier between the local and global minima has been overcome, and the optimization is progressing toward the global minimum. [Reproduced from Meerts *et al.* (2009) with permission from the American Institute of Physics.]

Figure 2 depicts the first four generations of an evolution strategy and demonstrates the effect of the chosen strategy. In general, the evolution strategies converge faster and are more rigid than the genetic algorithm.

3.3 The Fitness Function for the Analysis of Spectra

A proper choice of this fitness function is of vital importance for the success of the EA convergence. Meerts *et al.* (2004) have defined a fitness function F_{fg} as

$$F_{fg} = \frac{(f,g)}{\|f\| \|g\|} \tag{7}$$

Here f and g are the vector representations of the experimental and calculated spectrum, respectively. The inner product (f, g) is defined with the metric W, which has the matrix elements $W_{ij} = w(|j - i|) = w(r)$ as

$$(f,g) = f^T W g \tag{8}$$

and the norm of f as $||f|| = \sqrt{(f, f)}$; similar for g. For w(r), a triangle function was used (Hageman *et al.* 2000) with a width of the base of Δw :

$$w(r) = \begin{cases} 1 - |r| / \left(\frac{1}{2}\Delta w\right) & \text{for } |r| < \frac{1}{2}\Delta w \\ 0 & \text{otherwise.} \end{cases}$$
(9)

The above-defined fitness function is able to smoothen, in a controlled way, the fitness landscape and therefore allows the EA to locate the global minimum. The width of the function w(r) critically determines the ability of the EA to converge to the global minimum and also the speed of convergence. The smoothing of the fitness landscape allows to sense regions far from the minimum. The EA convergence is obtained in a well-defined procedure. At first the function w(r) should be chosen relatively broad; $\Delta w \approx$ 15 - 20 times the line widths of an individual transition in the spectrum. In this way, a first set of parameters is obtained, which still has to be refined. This is done by decreasing Δw and narrowing the limits of the parameter space to be searched in the fit. In a final calculation Δw is set to zero. Usually full EA convergence to the best set of parameters is achieved by narrowing Δw to zero in one or two steps.

3.4 Comparison between the Genetic and the Derandomized-ES DR2 Algorithms

The EAs contain a number of internal parameters, which determine the performance of the fit, and have to be optimized for a given problem themselves.

The speed and convergence of GA algorithms depend on a relatively large number of internal parameters among others on the encoding of the parameters (binary or real), the crossover type (one-point, two-point, uniform), the size of the starting population, the rate of elitism and the mutation probability. This has been discussed to be a major drawback of this method (Wehrens *et al.* 1999). The metaoptimization for some of the parameters is described by Meerts *et al.* (2004).

The number of optimizing parameters for the derandomized-ES DR2 and CMA-ES algorithms are quite a bit smaller than that of the genetic algorithm. In DR2 only a choice has to be made for the offspring size and the maximum age of an individual to stay in the population. In the CMA-ES only the size of the offspring can be changed.

A comparison has been made between the GA and DR2 algorithms using the same synthetic spectrum as in Meerts *et al.* (2004). The result of this comparison is shown in Figure 3. In these calculations the meta-parameter for the GA the parameters are *crossover probability* = 85%, *mutation probability* = 5% and *elitism* = 50%. This figure clearly shows that for this particular kind of spectrum

the DR2 is converging faster and yields parameter values that are closer to the *real* ones. Although the speed of convergence is an important criterion, the total computing time is a second one. Because of the elitism factor, not all individuals have to be (re)calculated in each generation in contrast to the ES. This makes the computing time for a full generation of a GA calculation typically a factor 2 faster than that of an ES calculation. However, since the ES tends to converge much faster than the GA, the convergence of an ES evolution needs typical a factor of 2 less computing time than that of a GA fit. One of the main advantages of ES over GA is the fewer number of internal strategy parameters that need to be determined in a meta-optimization.

3.5 Use of the GA for the Structure Determination

The GA is not only utilized for an automated assignment of the rovibronic spectra but also for the (nonlinear) fit of the molecular structure to the experimentally determined rotational constants. Ratzer *et al.* (2002) have presented a computer program to determine molecular structures from the rotational constants of a set of different isotopologs using a gradient-based χ^2 minimizer. This program was complemented with a GA-based global optimizer (Schmitt *et al.* 2006b). The global minimizer has been extended to allow for intermediate local minimization steps rather than simple cost function evaluations. This technique was originally described by Li and Scheraga as part of a simulated annealing (SA) exploration of the potential hypersurface of proteins (Li and Scheraga 1987). The algorithm, which



Figure 3 Comparison of three randomly started fits comparing the genetic (ga) and derandomized-ES DR2 (der) algorithms as function of the size of the population (ga) or offspring (der). In this figure, the *fitness* function equation (7) is shown, which has a maximum values of 100.

is also known as *basin-hopping* (Wales and Doye 1997), was successfully used in combination with GA and/or SA to determine minimum-energy structures of fullerenes and atomic clusters (Deaven and Ho 1995).

To increase the number of (independent) data points in the geometry determinations, both the rotational constants as well as the angles of the TDM of the respective isotopolog with the main inertial axes are taken into account. Both experimental data sets are completely independent from each other and can be extracted from the GA (or ES) fits of the spectra.

The cost function to be minimized in a fit of the structure to the rotational constants of k different isotopologs is the weighted sum of squared residuals (Brandt 1998):

$$\chi^2 = \chi^2_{\rm rot} = \Delta y^T W_{\rm rot} \Delta y \tag{10}$$

In this equation $y^T = (B_{a,1}^0, B_{b,1}^0, B_{c,1}^0, \ldots, B_{a,k}^0, B_{b,k}^0, B_{c,k}^0)$ represents the vector of n = 3k measured and calculated rotational constants y_{exp} and y_{calc} of the k isotopologs, respectively, $\Delta y = y_{exp} - y_{calc}$ is their corresponding vector of residuals and W_{rot} denotes a $n \times n$ positive-definite weight matrix, which is obtained from the covariance matrix of the *n* experimental rotational constants. The rotational constants directly depend on the Cartesian coordinates of the molecule, thus the mathematical relation between those *n* inertial values and the internal coordinates b_j , j = 1..m, of a hypothetical equilibrium structure is straightforward.

There exists no direct relation between the angles of the TDM with the main inertial axes, and the internal coordinates b_j of the molecule similar to that of the inertial parameters. However, we can deduce an indirect relation by recognizing that the rotation of the inertial frame of a given reference isotopolog into any other isotopolog also implies a corresponding change of the TDM vector. The TDM vector $\mu_{ref} = (\mu_{ref,a}, \mu_{ref,b}, \mu_{ref,c})^T$ of the reference isotopolog is transformed into the TDM vector μ_{other} of another isotopolog via

$$\mu_{\text{other}} = \boldsymbol{R}_{\text{other}}^T \boldsymbol{R}_{\text{ref}} \mu_{\text{ref}}$$
(11)

In this equation, \mathbf{R}_i denotes the 3 × 3 orthogonal rotation matrix, which transforms the initial geometry of the *i*th isotopolog into its own inertial system. Using both rotational constants and TDM orientations, we define the cost function as

$$\chi^{2} = \chi^{2}_{\rm rot} + \chi^{2}_{TM} = \Delta y^{T} W_{\rm rot} \Delta y + \Delta z^{T} W_{\rm TDM} \Delta z \quad (12)$$

where either $z^T = (\theta_1, \phi_1, \dots, \theta_r, \phi_r)$ or $z^T = (\lambda_{a,1}^2, \lambda_{b,1}^2, \dots, \lambda_{a,r}^2, \lambda_{b,r}^2)$ in the general *abc* case depending on the available *r* transition dipole moment data. In the

special case of *ab* hybrid spectra, the vector *z* simplifies to $z^T = (\theta_1, \ldots, \theta_r)$ or $z^T = (\lambda_{a,1}^2, \ldots, \lambda_{a,r}^2)$, respectively. Similarly to the weight matrix W_{rot} of the rotational constants, W_{TDM} is the weight matrix obtained by the covariance matrix of the TDM data.

4 ROTATIONALLY RESOLVED ELECTRONIC SPECTROSCOPY

Since most spectra that are dealt with in this chapter are rovibrationally resolved electronic spectra, we explain the experimental setup and some of the experimental implications in more detail.

The term *high-resolution electronic spectroscopy* is used in this chapter only for rotationally resolving techniques of electronic spectroscopy. Two prerequisites must be fulfilled in order to achieve this goal. First, a laser system is needed that emits in the visible (400-750 nm) and/or in the UV (400-250 nm) range, with a resolution of about 1 MHz or better. At a typical wavelength of 300 nm, a bandwidth of 1 MHz is equivalent to a resolution of 1 in 10^9 . Secondly, an experimental setup, in which the Doppler broadening is minimized and at the same time the temperature of the sample is reduced in order to decrease the number of populated rovibronic states. Especially for large molecules, this is an indispensable requirement, since the spectra get increasingly complicated with increasing number of transitions.

The problem of Doppler broadening in electronic spectra is outstandingly severe, since the Gaussian broadening due to the Doppler effect is proportional to the excitation frequency. Thus, while of minor importance in microwave spectroscopy (frequencies between 0.3 and 300 GHz), it turns a major problem in electronic spectroscopy (frequencies exceeding 1 PHz). Thus, the Doppler broadening in the UV is more than 10^6 times larger than in the microwave region. The reason for the Doppler broadening in gas phase experiments is the large velocity distribution of the molecules in the direction of the laser beam. Any technique aiming at a reduction of the Doppler width must deal with this velocity distribution. The application of seeded molecular beams (Ross 1966, Scoles 1988), which are crossed at right angles with the laser considerably reduces the Doppler broadening. By seeding, we refer to the technique of mixing a small amount of the molecules under investigation with a surplus of a rare gas, prior to expansion. A further reduction of the Doppler width can be achieved by confining the molecular beam to a small angular range about the beam axis using one or more skimmers. By means of this geometric limitation of the molecular beam in the direction of the laser beam, a reduction of the velocity distribution in a thermal ensemble by a factor of 200 can be achieved. Thus,



Figure 4 Molecular beam machine used for the experiments described in this article. (a) probe container; (b) nozzle; (c,d) beam skimmers; (e) imaging optics; (f) photomultiplier; (g) quadrupole mass spectrometer; (h) turbo-molecular pumps; (i) rotatory pumps, (j) oil diffusion pump; (k) roots blower pump.

the Doppler width, which in a thermal ensemble of Argon at 373 K at a central wavelength of 300 nm amounts to about 2 GHz, can be reduced to 10 MHz by pure geometric skimming.

Further reduction of the Doppler width can be achieved by geometric filtering of the central part of the emitted fluorescence. Figure 4 shows a typical molecular beam setup used for rotationally resolved electronic spectroscopy. The molecular beam, which is formed by expansion of the probe (a), seeded in Argon through the nozzle (b) is skimmed twice (c,d) for a maximum reduction of the Doppler width. The beam is crossed at right angles with the laser light and the fluorescence is collected at right angles through an imaging optics (e) and focussed onto a photomultiplier (f). The molecular beam position can be controlled by means of a quadrupole mass spectrometer (g).

Apart from cooling the seeded molecules in the molecular beam of noble gases to rotational temperatures below 2 K, condensation of the seed molecules takes place, making weakly bound complexes experimentally accessible. At typical vibrational temperatures of a few 10 K and rotational temperatures below 2 K, hydrogen-bound complexes and even van der Waals bound complexes are stable species and can be investigated spectroscopically in molecular beams. High stagnation pressures and large nozzle diameters are necessary prerequisites for cluster formation, since the rate of the cluster formation determining three body collisions scales with $P_r^2 D/T_r^2$ where P_r is the stagnation pressure in the reservoir, T_r is the reservoir temperature and Dis the nozzle diameter. From this it follows that high pumping speeds are necessary for observing larger clusters in molecular beams (Scoles 1988).

Several detection schemes are possible. The most common one is the use of laser-induced fluorescence (LIF), in which the emitted fluorescence acts as probe for the absorption of the laser light by the molecules under investigation (Majewski and Meerts 1984, Majewski *et al.* 1989, Schmitt *et al.* 2000).

Another possible detection scheme is the multiphoton ionization (MPI) (Sussmann *et al.* 1994), which has the advantage compared to LIF of yielding mass resolved spectra, which is especially advantageous if molecular complexes are investigated with similar absorption frequencies. On the other hand, the resolution in MPI experiments is limited by the necessary use of pulsed lasers with a temporal band width of a few nanoseconds to the Fourier limit of the pulse duration. Typical resolutions that can be achieved in these experiments are on the order of 70-100 MHz.

Other detection schemes comprise sensitized phosphorescence as probe (Uijt de Haag and Meerts 1989), in cases were a strong intersystem crossing results in a weak or even completely absent fluorescent signal, or the direct measurement of absorption of radiation in the molecular beam via optothermal techniques (Becucci *et al.* 1997) in cases were radiationless processes dominate, which lead to highly vibrationally excited ground-state molecules.

Another complementary method for determination of rotational constants in different electronic states must be mentioned, when talking about rotationally resolved electronic spectroscopy. Felker and coworkers established the so-called rotational coherence spectroscopy (RCS) (Felker 1992), which was considerably improved by Riehn (2002) in the last few years. While the rotational resolution in the experiments described here, is achieved by limiting the spectral band width as described in the preceding article, RCS utilizes a completely different approach. Like the Fourier transform (FT) describes the mathematical relationship between time and frequency domain, this approach is utilized in RCS by measuring the time which a molecule needs for rotation per radian. For large molecules, these times are of the order of a few picoseconds. Thus, to measure the rotational recurrences of large molecules, exciting laser pulses with picosecond or even subpicosecond duration are necessary. Between spectrally and temporarily resolving spectroscopic techniques exists a complementary relationship, since the frequency distances of the rotational lines decrease with increasing moment of inertia, while the distance between two recurrences increases.

In the following it is shown, how the evolutionary methods can be utilized in the practical analysis of very complex spectra. We chose rotationally resolved electronic spectra in a molecular beam, Fourier transform infrared (FTIR) spectra in the gas phase, and liquid crystal NMR spectra to show the broad applicability of the method.

5 APPLICATION OF EA AND GA TO THE FIT OF ROTATIONALLY RESOLVED ELECTRONIC SPECTRA

5.1 Structure and Stabilization of Aromatic Water Complexes

Water complexes of polar aromatic molecules have found considerable interest as model systems for solvation processes, ground state and electronically excited state acidities, and dynamical processes, connecting the primarily excited state with other electronically excited states or with the electronic ground state. In all the examples that we present here, a splitting of the rovibronic spectra into two components by a tunneling motion of the water moiety is found. While for the smaller of these systems the rotational constants are so large that, despite the doubling of lines by the tunneling splitting a rotational resolution of the lines is achieved, larger systems like the p-cyanophenol(H_2O_1) or the tryptamine $(H_2O)_1$ cluster constitute a problem due to congestion of the spectra like in the case of the aromatic dimers. Another example of a water complex with strongly overlapping rovibronic lines is the p-cresol(H₂O)₁ cluster, in which a splitting due to the tunneling motion of the water moiety and an additional splitting from the internal rotation of the methyl group leads to four overlapping torsional subbands.

5.1.1 Phenol-water

Among the most studied hydrogen-bound clusters of aromatic molecules, the phenol(H_2O)₁ cluster certainly ranks at the top. From early days of laser spectroscopy in molecular beams on this cluster was subject by a multitude of experimental and theoretical investigations (Abe *et al.* 1982, Lipert and Colson 1988, Stanley and Castleman 1991, Schütz, *et al.* 1993, Berden *et al.* 1996).

The Spectra and their Assignment The rovibronic spectrum of the electronic origin of phenol(H2O)1 at $35\,996.47\,\mathrm{cm}^{-1}$ shown in Figure 5 is an example of a spectrum of a hydrogen-bonded cluster, that can be easily analyzed and fit using classical line position assignment techniques. Indeed, the original spectrum was completely analyzed manually (Berden et al. 1996). Owing to a hindered internal rotation of the water moiety in the cluster, the spectrum is split into two subtorsional components, $\sigma = 0$ and $\sigma = 1$ with a splitting of 25.4 GHz between the electronic origins of both subtorsional components. The separate simulations of each component is shown in Figure 5. The lowest trace, which shows a zoomed portion of the simulated spectrum along with the stick spectrum reveals that most of the lines in the spectrum are composed of single rovibronic transitions, making a line position assigned fit easy in this case.

Determination of the Structural Parameters The rotational constants of the phenol-water cluster, which are compiled in Table 1 show that all heavy atoms of $phenol(H_2O)_1$ must be in one plane. From the rotational constants, an inertial defect of $-2.4 \text{ u}\text{\AA}^2$ for the ground state and of $-2.8 \text{ u}\text{Å}^2$ for the electronically excited states is calculated. This is in agreement with a structure, in which the two hydrogen atoms of the water moiety are symmetrically out of the aromatic plane by about 1 Å. The fact that the rotational B and C constants of the $\sigma = 0$ and the $\sigma = 1$ subtorsional components are equal and the only difference is in the A rotational constant immediately leads to the conclusion that the axis of the internal rotation of the water moiety is parallel to the *a*-axis of the cluster. Using the rotational constants of the monomer and the phenol-water cluster, the position of the center of mass (CM) of the water moiety with respect to the position of the CM of phenol can be calculated. Since the position of the phenolic O-atom and the water O-atom with respect to each CM are known, the hydrogen bond length O-O can be calculated. The complete procedure is described in (Berden et al. 1996) and yields bond lengths of 293 pm for the electronic ground state and 289 pm for the electronically excited state. The above considerations lead to the structure of the phenol-water cluster as shown in Figure 6. The



Figure 5 Rovibronic spectrum of the electronic origin of phenol(H₂O)₁ at 35996.47 cm⁻¹. The traces show the experimental spectrum, the simulated spectrum, the simulations of the subtorsional components $\sigma = 0$ and $\sigma = 1$ and zoomed portion of the simulation together with the stick spectra of both components.



Figure 6 The structure of the phenol- $(H_2O)_1$ cluster.

barrier to torsion of the water moiety is determined to be 180 and 130 cm^{-1} for the S₀ and the S₁ states, respectively.

5.1.2 p-Cyanophenol-water

The rotationally resolved spectrum of the electronic origin of the p-cyanophenol(H₂O)₁ cluster has a considerably higher line density than the spectrum of phenol(H₂O)₁. Two main reasons exist for that. First, the *A*, *B*, and *C* rotational constants are smaller than in the phenol–water cluster, with the latter two being nearly half of the size as in phenol–water. Secondly, the changes of the rotational constants *B* and *C* upon electronic excitation (ΔB and ΔC) are much smaller in the p-cyanophenol–water cluster compared to phenol–water (cf Table 1). The Spectra and their Assignment The rovibronic spectrum of the electronic origin of p-cyanophenol(H_2O)₁ given in Figure 7 shows the same kind of torsional splitting due to the hindered water torsion as the phenol(H_2O)₁ spectrum, although the splitting between the subtorsional origins is only 8.1 GHz compared to 25.4 GHz (Jacoby *et al.* 2006). The smaller subtorsional splitting is caused by a much higher barrier to the water torsion compared to phenol(H_2O)₁, which can be traced back to an higher additional interaction of one of the water oxygen lone pairs, with the ortho CH bond, which is much more acidic than in any other hydroxybenzene derivative (cf the benzonitrile dimer in section 5.2.2).

The lowest trace in Figure 7, which shows the simulation along with the stick spectra of both subtorsional

	phenol(H ₂ O) ₁	p-cyanophenol(H ₂ O) ₁	p-methylphenol(H ₂ O) ₁
$A''(\sigma = 0)$ /MHz	4291.49(4)	3462.16(50)	3663.48(244)
$B''(\sigma = 0)/MHz$	1092.1445(2)	587.87(8)	765.60(59)
$C''(\sigma = 0)/MHz$	873.7271(2)	503.62(6)	637.00(48)
$A''(\sigma = 1)/MHz$	4281.76(1)	3459.95(6)	3652.19(53)
$B''(\sigma = 1)/MHz$	1092.3254(1)	588.00(4)	765.98(31)
$C''(\sigma = 1)/MHz$	873.9082(1)	503.75(2)	637.71(31)
V_{2}''/cm^{-1}	180	275	183
v_0^2/cm^{-1}	35995.62(1)	35304.46(2)	34971.891(10)
$\Delta \nu/\text{GHz}$	25.455(10)	8.1494(10)	29.428(7)
$\Delta A(\sigma = 0)/\mathrm{MHz}$	-102.7(6)	-48.69(24)	-72.61(345)
$\Delta B(\sigma = 0)/\mathrm{MHz}$	8.4(2)	-0.11(1)	9.18(83)
$\Delta C(\sigma = 0)/\mathrm{MHz}$	0.89(6)	-1.08(1)	3.76(68)
$\Delta A(\sigma = 1)/MHz$	-114.36(2)	-50.39(5)	-75.82(75)
$\Delta B(\sigma = 1)/MHz$	8.30(2)	-0.13(1)	8.37(44)
$\Delta C(\sigma = 1)/MHz$	0.94	-1.06(1)	3.90(44)
$\Delta V_2/\mathrm{cm}^{-1}$	-30	-92	-58

Table 1 Comparison of the molecular parameters of the electronic origin bands of the phenol(H_2O_{11} , the p-cyanophenol(H_2O_{11} , and the p-methylphenol(H_2O_{11} cluster.



Figure 7 Rovibronic spectrum of the electronic origin of p-cyanophenol(H₂O)₁ at 35304.46 cm⁻¹. The traces show the experimental spectrum, the simulated spectrum, the simulations of the subtorsional components $\sigma = 0$ and $\sigma = 1$ and zoomed portion of the simulation together with the stick spectra of both components.

components reveal impressively why the use of automated assignment techniques is mandatory in this case. Up to 50 single rovibronic transitions contribute to each feature in the spectrum, which appears to be a single "line".

Determination of the Structural Parameters The same considerations, as made for the phenol $(H_2O)_1$ cluster, also hold for the p-cyanophenol $(H_2O)_1$ cluster. The orientation

of the water moiety in the cluster can be described as *trans* linear. The fit of the hydrogen bond lengths in the ground and the electronically excited state reveal that both hydrogen bonds are shorter $(r_{OO}(S_0) = 283 \text{ pm}; r_{OO}(S_1) = 269 \text{ pm})$ than in the case of phenol–water. Also, the decrease of the hydrogen bond length, upon electronic excitation, is larger than in phenol–water. Both facts can be traced back to the higher acidity of p-cyanophenol.

While phenol is a moderate acid in its electronic ground state ($pK_a = 9.86$), its acidity increases upon electronic excitation by nearly four orders of magnitude to a pK_a of 6 (Granucci *et al.* 2000). p-cyanophenol is a much stronger acid in the ground state ($pK_a = 7.74$) and simultaneously, the increase of acidity upon electronic excitation to a pK_a of 3.33 larger than in phenol.

Also, the acidity of the *ortho* CH position is considerably increased compared to phenol. With the strong reduction of the O–O hydrogen bond length, both oxygen lone pairs can overlap to form two hydrogen bonds. The stronger bond is formed to the hydroxy group, the weaker one to the CH group (cf Figure 8).

5.1.3 p-Methylphenol-water

The interpretation of the methylphenol $(H_2O)_1$ water cluster spectrum bears the highest difficulties among the hitherto described hydroxybenzene-water clusters. It is not just that the rotational constants are smaller than in phenol-water but there is also an additional complication that arises from the internal torsional motion of the methyl group. While in phenol-water and p-cyanophenol-water the spectrum was split into two subtorsional components ($\sigma = 0$ and $\sigma = 1$), in the p-cresol-water complex four subtorsional components have to be considered. Each of the $\sigma = 0$ and $\sigma = 1$ components is further split by the threefold methyl rotation in a set of A and doubly degenerate E levels (Myszkiewicz et al. 2005). The two subtorsional bands of the water torsion and the respective subtorsional bands of the methyl torsion could be fitted independently, showing that the two motions are practically decoupled.

The Spectra and their Assignment Figure 9 shows the rovibronic spectrum of the electronic origin of pmethylphenol(H₂O)₁ at 34971.89 cm⁻¹. At first sight the higher line density compared to phenol(H₂O)₁ and to pcyanophenol(H₂O)₁ cluster is obvious. The origins of the $\sigma = 0$ and $\sigma = 1$ are marked in the simulation trace of



Figure 8 The structure of the p-cyanophenol- $(H_2O)_1$ cluster.

Figure 9, together with the A - E splitting of each component. The origins of the $\sigma = 0$ and $\sigma = 1$ components are 29.4 GHz apart, compared to 25.4 GHz for phenol-water and 8.1 GHz for p-cyanophenol-water, alluding similar torsional barriers for the water torsion in p-methylphenol-water and phenol-water. The molecular parameters from the GA fitp of p-methylphenol(H₂O)₁ are compiled in Table 1 and are compared to those of phenol(H₂O)₁ and of p-cyanophenol(H₂O)₁.

Determination of the Structural Parameters Again, as for phenol(H_2O)₁, and p-cyanophenol(H_2O)₁ a *trans* linear hydrogen bond arrangement is found with a O–O hydrogen bond length of 290 pm in the electronic ground state and 285 pm in the electronically excited state, close to the value of unsubstituted phenol and considerably longer than in pcyanophenol(H_2O)₁. The barrier in the electronic ground state is determined to be 183 cm^{-1} , close to the value of phenol–water.

5.1.4 Tryptamine-water

The case of the tryptamine(H_2O)₁ cluster is insofar exceptional, that the tryptamine monomer moiety exists in seven different conformational forms in the gas phase (Nguyen *et al.* 2005). First Park *et al.* (1986) and later Connell *et al.* (1990) and Schmitt *et al.* (2005) showed how this structural diversity collapses into a single conformer upon cluster formation with one water molecule.

The Spectra and their Assignment Contrary to the phenol-water clusters described in the preceding sections, the vibronic spectrum of the origin of the tryptamine $(H_2O)_1$ cluster does not show a torsional splitting. Figure 10 presents the experimental spectrum with the simulation using the best parameters from a GA fit. As in the water clusters of the phenol derivatives, a multitude of lines contribute to each feature observed in the experimental spectrum, which is what made the use of GA for the automated spectral assignment and determination of the molecular parameters given in Table 2 indispensable.

Determination of the Structural Parameters Extensive ab initio calculations are necessary to determine the binding position of the water molecule to the tryptamine moiety in the cluster. Several positions are possible: First, the water might bind to the pyrrolic NH group via a hydrogen bond, with the indole acting as proton donor, similar to the indole–water cluster. Secondly, the water can bind to the ethylamino group via a hydrogen bond. Three cases must be distinguished here: (i) the amino group might act as proton acceptor, the water as proton donor; (ii) the amino group is the proton donor via the H_a atom; (iii) the amino group



Figure 9 Rovibronic spectrum of the electronic origin of p-methylphenol(H₂O)₁ at 34971.89 cm⁻¹. The traces show the experimental spectrum, the simulated spectrum, the simulations of the subtorsional components $\sigma = 0$ and $\sigma = 1$ and zoomed portion of the simulation together with the stick spectra of both components.

is the proton donor via the H_b atom. From comparison of the rotational constants and the stabilization energies, only the structure, shown in Figure 11 is considered, which immediately gives the answer to the question, why one out of seven tryptamine conformers is frozen out, upon water complexation. In the structure shown, water acts as proton donor to the amino group, but at the same time is the proton acceptor to the CH group of the pyrrole ring. Since the Gpy(out) conformer is the only one, in which the distance of the CH group and the amino group is small enough to be bridged by a single water molecule, this explains the exclusive appearance of only one conformer.

Here, as in other complexes of indole derivatives with polar solvent molecules, the question arises about the electronic nature of the electronically excited state. Two close-lying electronically excited states, which are historically labeled L_a and L_b , might change their energetic ordering upon cluster formation with polar molecules, since the L_a state is highly polar, whereas the polarity of the L_b state is much smaller. In the present case, comparison of the structure from ab initio calculations, shown in Table 2 with the experimental rotational constants clearly shows that the experimentally observed electronic origin of the water cluster remains of L_b type, as the monomer.

5.2 Global Versus Local Electronic Excitation in Symmetric and Antisymmetric Aromatic Dimers

Many publications have raised the question, if the electronic excitation in symmetrically bound homodimers of aromatic compounds is localized in one of the monomer moieties or is completely delocalized over both monomer moieties. Prototypes of these symmetrical dimers are the benzoic acid dimer investigated by Poeltl and McVey (1984) and by Remmers et al. (2000), the salicylic acid dimer (Yahagi et al. 2001), the 2-pyridone dimer (Held and Pratt 1992), (Müller et al. 2002), the o-cyanophenol dimer (Lahmani et al. 2002) and the benzonitrile dimer (Schmitt et al. 2006a). While for the 7-azaindole dimer and the 2-pyridone dimer delocalized excitation is found, the benzoic acid dimer, the fluorobenzoic acid dimer, and the o-cyanophenol dimer show a local excitation in only one of the moieties. The strength of electronic coupling between two or more chromophores is crucial in exciton formation for energy harvesting or energy funneling devices in nature.

5.2.1 The Phenol Dimer

Among others, the phenol dimer is an ideal model system to study the very sensitive equilibrium between hydrogen bonding and van der Waals interactions. The spectroscopy



Figure 10 Rovibronic spectrum of the electronic origin of the water cluster of the Gpy(out) conformer of tryptamine together with the best simulation, a zoomed portion of the experimental spectrum and the zoomed simulation together with the stick spectrum.

Table 2 Molecular Parameters for the tryptamine–water cluster determined from GA fits, as described in the text. The MP2 calculations were performed with the G03 program package Frisch *et al.* 2003 using the 6-311G(d,p) basis set. RICC2 calculations were performed with Turbomole V5.8 using the Ahlrichs-type valence triple-zeta basis sets with polarization functions (VTZP) from the turbomole library Ahlrichs *et al.* 1989 for both the ground and first electronically excited $\pi\pi^*$ states.

	Experiment	RIMP2	RICC2 (S ₀ and L_b)	RICC2 (S ₀ and L_a)
A''(MHz)	1465.78	1469	1464	1464
B''(MHz)	483.43	471	498	498
C''(MHz)	397.99	387	410	410
v_0/cm^{-1}	34957.11	_		32549
$\Delta A(MHz)$	-6.82			-76
$\Delta B(MHz)$	-4.71	_		+39
$\Delta C(MHz)$	-3.72	—		+26

of complexes of phenol with different solvent molecules has been the subject of a plethora of publications in the last decades. In most of these complexes phenol acts as a proton donor with respect to the solvent molecule.

The phenol dimer takes a special position, since one of the cluster constituents acts as proton donor, the other as proton acceptor, both of which are accessible spectroscopically as was shown by Fuke and Kaya (1983). Van der Waals interactions between the π -systems of the aromatic rings are most likely to play an important role for mutual arrangement of the monomer moieties, while for most of the other phenol clusters the structure of the cluster is mainly governed by dipole–dipole interactions. *The Spectra and their Assignment* Figure 12 shows the rovibronic spectrum of the electronic origin of the phenol dimer together with the best GA fit using the molecular parameters from Table 3.

The lowest trace of Figure 12 shows a zoomed part of the convoluted simulation along with the unconvoluted stick spectrum. Most lines in the spectrum are superpositions of up to 15 single rovibronic transitions that contribute with similar intensities. Such heavy congestion of lines can, in general, only be handled using an EA-based fitting strategy. The large density of rovibronic lines avoids a classical assignment procedure or at least makes it an incredibly tedious task.



Figure 11 The structure of the tryptamine- $(H_2O)_1$ cluster.



Figure 12 Rovibronic spectrum of the electronic origin of the $h_6 - h_6$ isotopolog of the phenol dimer at 36044.70 cm⁻¹. The following traces show the simulated spectrum using the rotational constants from Table 3, a zoomed portion of the experimental spectrum and the zoomed simulation together with the stick spectrum.

The spectrum is fit using a rigid rotor Hamiltonian. It consists of about 13000 lines in a range of less than 2 cm^{-1} . An additional challenge lies in the fact that the spectrum is composed of nearly equal *a*-, *b*-, and *c*-type fractions. At a rotational temperature of about 5 K, more than 100 J states are populated with an intensity of at least 0.5 % of the strongest transition in the spectrum. The molecular parameters obtained from the fit are given in Table 3. Other isotopologs, which are indispensable for the structure determination have been measured and are gathered in Table 3.

Determination of the Structural Parameters The intermolecular parameters were determined using the GA-based fit of the structural parameters to the rotational constants and to the TDM orientations of the five isotopologs given in Table 3 as described in Section 3.5. Table 4 and Figure 13 show the bond lengths, angles, and dihedral angles, which determine the relative orientation of the two monomer moieties with respect to each other for the ground and the first excited state.

The geometries shown in Table 4 and Figure 13 give clear evidence, that the hydrogen bond strength of the phenol dimer increases upon electronic excitation, while the dispersion interaction between the monomer moieties decreases.

5.2.2 The Benzonitrile Dimer

The dimer of benzonitrile displays an exception from the other cases in this article, since no $O-H \cdots O$ hydrogen bond is formed. Nevertheless, the monomer moieties are bound

	$h_{6} - h_{6}$	$d_1 - d_1$	$d_1 - h_6$	$d_{6} - d_{6}$	¹³ C- ¹³ C
<i>A''</i>	1416.99(39)	1376.23(10)	1399.78(124)	1239.33(12)	1413.22(15)
B''	313.51(1)	312.84(3)	312.72(6)	287.25(1)	311.84(2)
C''	288.11(1)	286.44(2)	287.16(6)	264.63(1)	286.70(2)
ϕ	58.2(30)	55.2(100)	66.6(23)	58.3(20)	57.4(30)
θ	29.5(30)	35.7(100)	37.6(16)	40.0(35)	35.9(60)
ΔA	10.71(1)	4.55(1)	7.81(10)	4.28(2)	10.60(1)
ΔB	-5.31(1)	-3.95(1)	-5.63(1)	-3.93(1)	-5.25(1)
ΔC	-5.82(1)	-4.76(1)	-5.99(1)	-4.90(1)	-5.76(1)

Table 3 Molecular parameters of the electronic origin band of five isotopologs of the phenol dimer obtained from genetic algorithm fits. Rotational constants and their differences in the electronically excited state are given in megahertz, the angles of the transition dipole moment ϕ and θ with the inertial axes are given in degrees.



Figure 13 (a) Atomic numbering of the phenol dimer, used in the definitions of Table 4. (b) Two different views of the S_0 (r_s) and the S_1 (r_s) geometry of the phenol dimer from Table 4.

Table 4 Experimentally determined pseudo-Kraitchman (r_s) geometry parameters of the phenol dimer in the S₀ and in the S₁-state. All distances are given in picometers, angles and dihedral angles in degrees. For the fit of the intermolecular structure of the dimer in the S₁ state we used a locally excited donor moiety.

	$S_0(r_s)$	$S_1(r_s)$
$r(H_{7a}O_8)$	236.9(54)	174.4(16)
$a(O_7H_{7a}O_8)$	150.5(20)	179.9(1)
$a(C_9O_8H_{7a})$	139.8(14)	135.7(9)
$d(O_8H_{7a}O_7C_1)$	9.4(47)	63.4(9)
$d(C_9O_8H_{7a}O_7)$	62.8(46)	-114.8(9)
$d(C_{10}C_9O_8H_{7a})$	-178.2(22)	-101.5(14)

by a hydrogen bond, which is a (weaker) $C-H\cdots N$ bond in this case. Since the C-H groups in ortho position to the nitrile group are quite acidic, a quite strong hydrogen bond can be formed between the nitrile group acting as proton acceptor and the CH group acting as proton donor. Of main concern is the symmetry and the structure of the dimer in the ground and excited states.

The Spectra and their Assignment Figure 14 shows the rovibronic spectrum of the electronic origin of the benzonitrile dimer together with the best GA fit using the molecular parameters from Table 5. The lowest trace shows a zoomed part of the convoluted simulation along with the calculated stick spectrum.

The spectrum is fit using a rigid rotor Hamiltonian. It is of a mixed *abc*-type and consists of about 18 000 lines in a range of 55 GHz. At a rotational temperature of about 2.8 K more than 95 J states are populated with an intensity of at least 0.5 % of the strongest transition in the spectrum. The rotational constants for both electronic states, the origin frequencies, the polar angles θ and ϕ , defining the



Figure 14 Rovibronic spectrum of the electronic origin of the benzonitrile dimer at 36420.10 cm^{-1} . The traces show the experimental spectrum, the simulated spectrum using the rotational constants from Table 5, a zoomed portion of the experimental spectrum and the zoomed simulation together with the stick spectrum.

Table 5 Molecular parameters of the electronic origin band of the benzonitrile dimer as obtained from the genetic algorithm fit. Standard deviations are given in parentheses. The MP2 calculations were performed with the G03 program package Frisch *et al.* 2003 using the 6-311G(d,p) basis set. RICC2 calculations were performed with Turbomole V5.8 using the VTZP basis set from the turbomole library Ahlrichs *et al.* 1989 for both the ground and first electronically excited $\pi\pi^*$ states.

	Experiment	MP2	RICC2
A"/MHz	1610.99(17)	1654	1645
<i>B"/</i> MHz	187.48(7)	185	186
C"/MHz	168.12(7)	167	167
$\Delta I''/\mathrm{u}\mathrm{\AA}^2$	-3.2727	0	0
ν_0/cm^{-1}	36420.10(1)		37851
$\Delta A/\mathrm{MHz}$	-28.08(10)		-28
$\Delta B/\mathrm{MHz}$	-0.01(10)		0
$\Delta C/MHz$	-0.31(10)		0
$\Delta \Delta I/\mathrm{u} \mathrm{\AA}^2$	-0.1667	0	0
	-0.1007	0	0

orientation of the transition dipole moment in the inertial frame, and the Lorentzian contribution to a Voigt profile with 15 MHz Gaussian contribution are given in Table 5.

Comparison of the stick spectrum with its convolution using the experimental line widths immediately reveals, why a "classical" fit with line position assignments is impossible for a molecular system of that size. Each "line" in the experimental spectrum is composed of up to 40 single rovibronic transitions, in many cases of very similar intensities. Figure 14 shows an expanded



Figure 15 The structure of the benzonitrile dimer. The hydrogen bond lengths for the electronic ground state and the excited state (in parentheses) are given as obtained from the structure fit to the rotational constants. The asterisk denominates the benzonitrile moiety, which is electronically excited.

view of the simulated stick spectrum along with the convoluted spectrum using a Gaussian width of 15 MHz and a Lorentzian contribution of 10 MHz.

Determination of the Structural Parameters The structure of the benzonitrile dimer has been determined from the rotational constants of a single isotopolog. Therefore, quite a few model assumptions have to be made. For the electronic ground state, the two monomer moieties have been fit to the microwave transitions from (Casado *et al.* 1971). Subsequently, the intermolecular parameters of the dimer are fit, assuming unchanged monomer geometries upon cluster formation. The result is shown in Figure 15. The S₁ structure for the dimer is fitted assuming local excitation in one of the monomer moieties. The geometry of one monomer was fitted to the rotational constants of the excited state of benzonitrile of Borst *et al.* (Borst *et al.* 2001), while the other was kept fixed at the Kraitchman geometry of the ground state. For the fit of the excited state of the monomer all C–C bond lengths in the aromatic ring are treated equally and are fit, furthermore the C₁C₇ distance is fit, the C₇N₈ distance is kept fixed at the ground-state value, and all CH bond lengths are kept fixed at a value of 107.1 pm, which is a typical value for excited state aromatic ring determined in this way is 142.8 pm, the C₁C₇ distance was determined to be 142.7 pm.

In this way we could fit the excited-state structure of the dimer, resulting in one considerably shorter N_8H_{10a} bond length in the electronically excited state compared to the ground state and one longer one.

The benzonitrile dimer exhibits a local excitation of one of the cluster moieties like the benzoic acid dimer. Obviously, the electronic coupling between the two monomer moieties is not strong enough to allow a delocalized electronic excitation.

5.2.3 The Benzoic Acid Dimer

One of the most well-known symmetric aromatic homodimers is the benzoic acid dimer. Since the C_1C_7 axes of the monomer moieties are nearly on a straight line, the A rotational constant of the cluster will be larger than in the case of the benzonitrile dimer. Nevertheless, the cluster is extended considerably more in the direction of the A axis compared to the benzonitrile dimer, leading to much smaller B and C rotational constants. As discussed in the following, the need for an automated assignment tool like ES is even larger in this case, than in any other, described hitherto.

The Spectra and their Assignment As in the case of the phenol dimer several isotopologs have been investigated. In the following, we show the results for only one of them, exemplarily. The investigated isotopologs are the undeuterated (d_0d_0) , two singly deuterated $(d_1d_0 \text{ and } d_0d_1)$, one doubly deuterated (d₁d₁), and the fully deuterated (d_5d_5) isotopolog. Single deuteration (d_1) in the above examples of isotopologs takes place at the hydrogen of the carboxy group. Figure 16 shows the rotationally resolved electronic origin of the d_0d_0 isotopolog of the benzoic acid dimer. The lowest trace shows a zoomed part of the convoluted simulation along with the calculated stick spectrum. The red and the blue sticks represent transitions within the two sublevels, caused by the tunneling of the two carboxy protons in the dimer. In such a dense spectrum one cannot talk about rovibronic lines; this is clearly a limit where the term rovibronic contour is more appropriate, despite the high experimental resolution. Each



Figure 16 Rovibronic spectrum of the electronic origin of the d_0d_0 isotopolog of the benzoic acid dimer at 35723.79 cm⁻¹. The traces show the experimental spectrum, the simulated spectrum using the rotational constants from Table 6, a zoomed portion of the experimental spectrum and the zoomed simulation together with the stick spectrum.

Table 6 Molecular parameters of the electronic origin band of the benzoic acid dimer as obtained from the EA fit. Standard deviations are given in parentheses. The MP2 calculations were performed with the G03 program package Frisch *et al.* 2003 using the 6-311G(d,p) basis set. RICC2 calculations were performed with Turbomole V5.8 using the VTZP basis set from the turbomole library Ahlrichs *et al.* 1989 for both the ground and first electronically excited $\pi\pi^*$ states.

	Experiment	MP2	RICC2
A"/MHz	1924.99(20)	1919	1925
<i>B"/</i> MHz	128.06(14)	127	128
C"/MHz	120.15(14)	119	120
$\Delta I''/uÅ^2$	-2.5139	0	0
v_0/cm^{-1}	35723.786(1)		36789
$\phi/^{\circ}$	73.18	_	89.9
θI°	79.00		90.0
$\Delta A/MHz$	-16.07(9)		-16
$\Delta B/MHz$	-0.52(12)		-1
$\Delta C/MHz$	-0.51(12)	_	0
$\Delta I'/uÅ^2$	-2.8210	—	0

feature in this contour consists of a few hundred single rovibronic transitions, clearly showing why the utilization of an evolution strategy is indispensable in this case. The benzoic acid dimer spectra were the first to be fit using the derandomized evolution strategy (ES DR2), cf section 3.2.

Determination of the Structural Parameters The main concern about the structure of the benzoic acid dimer is the symmetry of the cluster in the ground and electronically excited states. While all experimental investigations for the ground state agree in a C_{2h} symmetry, i.e., identical monomer moieties, the symmetries of the electronically excited state and the transition states for both ground and excited states are in question. The investigations described here show that the symmetry of benzoic acid dimer changes to C_s upon electronic excitation. Structurally, this can be explained by a tilting of the two monomer moieties, caused by the fact that they are no longer equivalent in the excited state since electronic excitation is associated with a change of acidity, as in the case of water clusters of several phenol derivatives (cf section 5.1.2).

5.3 Determination of a Large Number of Molecular Parameters

Evolution strategies are especially successful in cases were the parameter space is very large and/or unstructured. In the following two examples, we present results of spectral fits, in which a large number of parameters have to be fit simultaneously.

5.3.1 The $2_0^1 4_0^3$ Rovibrational Band of the $\widetilde{A}^1 A_2 \leftarrow \widetilde{X}^1 A_1$ Electronic Transition of Formaldehyde

Motsch et al. (2008) have presented the rotationally resolved room temperature absorption spectrum of the $2_0^1 4_0^3$ and the $2_0^2 4_0^1$ the rovibronic bands of formaldehyde in the $\widetilde{A}^1 A_2 \leftarrow \widetilde{X}^1 A_1$ electronic transition. Owing to the high temperature, high J-states are populated and sextic centrifugal distortion terms in the fit have shown to be necessary. The appropriate nuclear spin statistics (Ka even levels = 1; K_a odd levels = 3) has also been taken into account. In this case, the number of parameters to be fit amounts to 6 rotational constants (3 for each state), 10 quartic centrifugal distortion constants (5 for each state), 14 sextic centrifugal distortion constants (7 for each state), and one central frequency = 31 parameters. This is a typical case for the application of evolution strategies, since the advantage of these techniques over classical (local) optimizers is increasing with increasing parameter space.

5.3.2 The Case of Many Overlapping Isotopologues: Tryptamine

Another case of a large number of parameters which have to be fit occurs, when several isotopologues of one molecular species absorb with overlapping rovibronic bands. A case that has been studied extensively by us is the tryptamine system. Addition of D_2O to tryptamine results in a mixture of eight different readily forming isotopologues due to three acidic positions (the pyrrolic NH and the two amino H atoms) with the possibility of $H \leftrightarrow D$ exchange: one undeuterated, three singly deuterated, three doubly deuterated, and one triply deuterated isotopologues (Schmitt et al. 2005). All eight rovibronic spectra completely overlap, and a total of 61 parameters have to be fit for the whole spectrum (six rotational constants and a central frequency for each of the isotopologues and additionally two temperatures, a temperature weight factor (cf equation 4) and two line shape parameters). Regarding the high line density due to the small rotational constants and the large number of parameters to be fit, no other method as evolution strategies seem to be able to tackle this problem.

6 GA FIT OF ROVIBRONIC OR ROVIBRATIONAL CONTOURS

The examples shown above of very large molecular complexes pose the question as to whether the fitting technique is still applicable when the information from individual transitions or from groups of transitions is completely lost due to heavy spectral overlap. In the following, we show that the GA-based fitting strategy can also be used in cases where the spectral resolution is so low that only rotational contours can be resolved. This might occur in electronic spectroscopy, when the lifetime of the excited state is considerably smaller than 500 ps, leading to large Lorentzian broadening. Other limitations might be a low experimental resolution due to limited bandwidth of the exciting laser, as encountered frequently in pulsed laser experiments, or due to limited spectral resolution of the detection scheme as found, for example, in low resolution FTIR spectroscopy.

6.1 The Contours of Adenine Rovibronic Spectra

The medium resolved (the spectral resolution in these pulsed laser experiments is 0.04 cm^{-1} corresponding to 1200 MHz) LIF spectra of selected vibronic bands of adenine, shown in Figures 17 and 19, are an example of insufficient rotational resolution in electronic spectra due to a very short excited-state lifetime (Lee *et al.* 2006). The upper trace in Figure 17 shows the experimental LIF spectrum of the C-band of adenine, which has been subject to a long lasting controversy about its nature. Although, no single rovibronic feature is observed, a general structure with P, Q, and R branches is visible. The second trace shows the GA fit of the band. In the cases discussed so far, the unresolved rovibrionic contours are a consequence of very high line density. The stick spectrum, which is shown

in a zoomed central part of the simulated spectrum, clearly shows that for the adenine the rovibronic contour is caused by a broadening of each individual line by the short life time of the excited state. The lowest trace shows a Voigt convolution of the stick spectrum with a Gaussian and a Lorentzian width of 25 MHz, respectively. This would be equivalent to a (hypothetical) excited state lifetime of 6 ns, which is a typical value for indolelike chromophores. Clearly, a beautifully resolved spectrum could be expected with such a long lifetime.

The fit of a rovibronic contour like the one shown in Figure 17 is not as straightforward as the GA fits described in the previous sections. Many molecular parameters have to be fixed since the fit problem is severely underdetermined. Fortunately, the microwave spectrum of adenine was measured, so that the ground-state rotational constants are known. The structure shown in Figure 18 is deduced from these constants. Thus, there were only a limited number of parameters to fit: three changes of rotational constants upon electronic excitation, the direction of the TDM, the rotational temperature, and the Lorentzian linewidth. The most important parameters are those that determine the direction of the TDM, because they give important indications to the nature of the electronically excited state.

Other bands that appear in the low-resolution spectrum of adenine are the D and the E band. Both bands have rovibronic contours that are completely different from that



Figure 17 Rovibronic spectrum of the C-band of adenine at 36061.8 cm^{-1} along with the simulation using the best parameters from the GA fit. The subsequent trace shows a zoomed part of the spectrum with the stick spectrum overlayed. The lowest trace shows a convolution of the complete stick with a Voigt profile using Gaussian and Lorentzian widths of 25 MHz, respectively.



Figure 18 The structure of adenine.

of the adenine C band. Figure 19 shows the experimental spectrum and the simulation of the E band, which is very similar to the contour of the D band.

The main contribution to the intensity of the C band arises from *c*-type transitions, indicating an out-of-plane transition moment, while for the E and D bands mainly inplane transition moment components (*a* and *b*) dominate. Using this information, the C band can be identified as the electronic origin of the (out-of-plane) polarized $n\pi^*$ state, the D band as the origin of the (in-plane) polarized $\pi\pi^*$ state, and the E band as an out-of-plane vibrational band of the $n\pi^*$ state. This out-of-plane vibration is forbidden, but gains intensity via a vibronic coupling (Herzberg 1966) of the $n\pi^*$ and the $\pi\pi^*$ state. The large intensity of the electronically forbidden E band points to a large vibronic coupling between these two states.

6.2 The Fourier Transform Infrared Spectrum of Benzotriazole

Another example of a spectrum for which only the rotational contour of a vibrational band in spite of a fully resolved rovibrational spectrum was obtained is the FTIR spectrum of benzotriazole in the gas phase, which was examined by Roth et al. (1999). Two tautomers contribute to the IR absorption in the observed region, namely, the 1Hand the 2H-tautomer, which are shown in Figure 20. The reasons for the lack of rotational resolution in this example are the high temperature (400 K) and connected therewith the very high line density and Doppler width along with the limited resolution (0.5 cm^{-1}) of the FTIR spectrometer. Figure 21 shows the FTIR spectrum, which has been taken between 3450 and $3550 \,\mathrm{cm}^{-1}$ in the region of the NH stretching vibration. The NH stretching vibrations of the two tautomers have different absorption frequencies, and also different band types. The directions of the changes of the dipole moments are shown in the inertial axis system for each tautomer in Figure 20. Obviously, in the 2H tautomer a pure a-type band will be observed, while for 1*H*-benzotriazole an *ab*-hybrid type can be expected.

Also in this case, the GA fit is simplified by the fact that the ground-state rotational constants are known from microwave spectroscopy (for the 1H-tautomer) and from rotationally resolved electronic spectroscopy (for the 2H-tautomer). The NH-stretching band of the 2H-tautomer



Figure 19 Rovibronic spectrum of the E-band of adenine at 36247.7 cm^{-1} along with the simulation using the best parameters from the GA fit. The subsequent trace shows a zoomed part of the spectrum with the stick spectrum overlayed. The lowest trace shows a convolution of the complete stick with a Voigt profile using Gaussian and Lorentzian widths of 25 MHz, respectively.



Figure 20 Structure of 1H-benzotriazole (right) and 2H-benzotriazole (left). The arrow shows the direction of the dipole moment change upon NH stretching vibration.

must be a pure *a*-type band from symmetry considerations (cf Figure 20). The hybrid type of 1H-benzotriazole, the changes of the rotational constants upon vibrational excitation, the center frequencies of both bands, and the relative intensity of both bands have been fit. The lowest trace in Figure 21 shows a small portion of the simulated stick spectrum. The blue stick represent transitions of the 2H-tautomer, the red sticks of the 1H-tautomer. Clearly, it is the high temperature of the sample, which makes it impossible to obtain rovibrational resolution in this case.

6.3 Concluding Remarks to the Fit of Rotational Contours

The above examples clearly show that the technique of automated fits using the GA also is successful for spectra, which lack full rotational resolution. Of course, the more spectral features are resolved, the more accurate is the determination of the molecular parameters and the less parameters have to be fixed in the fit. High temperatures are sometimes advantageous when the resolution is limited by the experimental setup, since the large number of rotational levels, which are occupied at high temperatures, makes spectral features, which are caused by band heads more easily visible. In this context, the reader is referred to the pioneering work of Hollas on the analysis of rotational band contour analysis of vibronic transitions at elevated temperatures (Hollas 1982).

7 ANALYSIS OF LIQUID CRYSTAL NMR SPECTRA

As a further example of the use of EA, we now discuss a completely different type of high-resolution spectra: nuclear magnetic resonance (NMR) spectra of molecules that are partially oriented in an anisotropic liquid-crystalline environment. The NMR spectroscopy of orientationally ordered solutes in anisotropic liquids yields a wealth of information about both the solute and the intermolecular interactions between the solute and the solvent (Burnell and de Lange 2003). The extension to solutes that can exist in several symmetry-unrelated conformers is both interesting and challenging. The conformer distribution in such flexible molecules is governed by the total potential, which, in the



Figure 21 FTIR Spectrum of benzotriazole, taken at 400 K, along with the individual simulations of both tautomers. The lowest trace shows the stick spectra of both tautomers in a zoomed 10-GHz window.



Figure 22 ¹H NMR spectrum of *p*-bromo-biphenyl dissolved in "magic mixture". The experimental spectrum is compared with the final fit, a zoomed portion of the experimental spectrum and the zoomed simulation together with the stick spectrum. The horizontal scales are in hertz. [Reproduced from Meerts *et al.* (2007) with permission from Elsevier Publishers.]

condensed phase, includes both intra and intermolecular terms. This distribution can also be affected by orientational ordering. The extent of both isotropic and anisotropic effects in the solvent is difficult to measure. Because dipolar couplings between nuclei on different rigid subunits of a flexible molecule are very sensitive to conformer geometry and orientational order, they offer an excellent route to gaining insight into this difficult problem. Up to now, the measurement of such dipolar couplings has been hampered by the difficulty of analysis of the complicated spectra that are obtained.

The number of independent intramolecular pair interactions determines the complexity of assignment of these NMR spectra. This number is small for molecules with a limited number of spins (≤ 6) and high symmetry. For solutes with more spins and less symmetry, the number of different dipole-dipole interactions increases rapidly, thus making the analysis of complex NMR spectra notoriously difficult. This in spite of the fact that in most cases the resolution is sufficiently high that little overlap in the transitions occurs. The origin of the difficulties in the analysis lies in the nature of the NMR Hamiltonian: the spectra usually lack easy recognizable regularities. Many of such spectra have therefore remained intractable for decades.

The dominating terms in the Hamiltonian of a molecule dissolved in a uniaxial nematic liquid crystal are given in equation 5. The number of dipolar interactions $D_{\mu\nu}$ grows

strongly with the number of spins in the molecule and their values can easily differ by an order of magnitude. In an attempt to reduce the number of parameters and to estimate the strength of the dipole–dipole interactions Meerts *et al.* (2007) have used the relations between the Saupe order matrix S_{kj} and the direct dipole–dipole interactions:

$$D_{\mu\nu} = -\frac{h\gamma_{\mu}\gamma_{\nu}}{4\pi^2} \langle r_{\mu\nu}^{-3} \rangle \sum_{k,l} S_{kl} \cos\theta_k \cos\theta_l \qquad (13)$$

where θ_k is the angle between the k molecule-fixed Cartesian axis and the vector connecting nuclei μ and ν at internuclear distance $r_{\mu\nu}$. The angular brackets denote averaging overall internal motions.

The order matrix S_{kl} is defined as averages over reorientational motion of the second-order Legendre polynomials:

$$S_{kl} = \frac{1}{2} \langle 3\cos\theta_{Z,k}\cos\theta_{Z,l} - \delta_{kl} \rangle \tag{14}$$

where $\cos \theta_{Z,k}$ is the angle between the molecule-fixed k axis and the space-fixed Z axis, which lies along the liquidcrystal director. The S tensor is traceless and symmetric.

In this way, it is possible to make a crude prediction of the dipole–dipole interaction from an estimated structure of the molecule.

Meerts *et al.* (2007) tackled the NMR spectrum of the nine-spin system *p*-bromo-biphenyl dissolved in a nematic

crystal by a two-step process. In the first step, a relatively crude geometrical structure of the solute is assumed and two independent order parameters, chemical shielding parameters and the dihedral angle between the two rings are varied in a GA-fitting procedure until the best correspondence with the experimental spectrum is obtained. Owing to the approximations made in this step, complete agreement cannot be obtained. However, in practice, the choice of geometry does not have to be perfect in order to obtain sufficient common features between calculated and experimental spectra. Approximate dipolar couplings are calculated in this step. In a second GA-calculation procedure, these quantities are used as starting values for a fit in which all dipole-dipole couplings and chemical shielding parameters are varied until convergence is reached. Exceptionally good agreement is obtained in step two because no constraints are placed on the dipolar coupling and chemical shielding parameters. A typical result is presented in Figure 22.

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ABBREVIATIONS AND ACRONYMS

VTZP	Ahlrichs-type valence triple-zeta basis sets
	with polarization functions
СМ	center of mass
CARF	correlation automated rotational fitting
CMA-ES	covariance matrix adaptation evolution strategy
DR	derandomized
EAs	evolutionary algorithms
ES	evolution strategy
FTIR	Fourier transform infrared
FTMW	Fourier transform microwave

FT	Fourier transform	
GA	genetic algorithm	
IR	infrared	
LIF	laser-induced fluorescence	
MW	microwave	
MPI	multiphoton ionization	
NMR	nuclear magnetic resonance	
RCS	rotational coherence spectroscopy	
SA	simulated annealing	
TDM	transition dipole moment	
UV	ultraviolet	

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