Large prebiotic molecules in space: photo-physics of acetic acid and its isomers

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1 INTRODUCTION

The richness of interstellar chemistry has been growing steadily with the variety of objects and regions observed. The excitation and abundances of molecules contain key information on the physical structure and evolution of the host regions. Through molecules, we can trace the cycle of matter for interstellar space with the variety of objects and regions observed. The excitation of interstellar chemistry has been growing steadily with the variety of objects and regions observed. Among these molecules, acetic acid is of particular importance due to its structural proximity to glycine, the simplest amino acid. We compute electronic and vibrational properties of acetic acid and its isomers, methyl formate and glycolaldehyde, using density functional theory. From computed photo-absorption cross-sections, we obtain the corresponding photo-absorption rates for solar radiation at 1 AU and find them in good agreement with previous estimates. We also discuss glycolaldehyde diffuse emission in Sgr B2(N), as oppose to emissions from methyl formate and acetic acid that appear to be concentrate in the compact region Sgr B2(N-LMH).

Key words: astrochemistry - molecular processes - methods: numerical

ABSTRACT

An increasing number of large molecules have been positively identified in space. Many of these molecules are of biological interest and thus provide insight into prebiotic organic chemistry in the protoplanetary nebula. Among these molecules, acetic acid is of particular importance due to its structural proximity to glycine, the simplest amino acid. We compute electronic and vibrational properties of acetic acid and its isomers, methyl formate and glycolaldehyde, using density functional theory. From computed photo-absorption cross-sections, we obtain the corresponding photo-absorption rates for solar radiation at 1 AU and find them in good agreement with previous estimates. We also discuss glycolaldehyde diffuse emission in Sgr B2(N), as oppose to emissions from methyl formate and acetic acid that appear to be concentrate in the compact region Sgr B2(N-LMH).

Key words: astrochemistry - molecular processes - methods: numerical

The Sgr B2 molecular cloud complex is the prime target in the search for complex species, in particular in a hot core, the so-called Large Molecule Heimat, Sgr B2(N-LMH), within the more extended molecular cloud. In this compact source, smaller than the Oort cloud (~ 0.08 pc) with a mass of several thousands M⊙ (Miao & Snyder 1997), an extraordinary number of complex organics have been observed to exhibit very high column densities (e.g. amino acetonitrile, Belloche et al. 2008). Large partly hydrogen-saturated species challenge the completeness of the standard ion-neutral scheme in interstellar chemistry, suggesting that reactions on dust grains are involved in their formation (e.g. Bennett & Kaiser 2007).

Of the chemical species detected so far, particular attention has been paid to the formation of different isomer groups. In this work, we focus on C₃H₇O₂, i.e. acetic acid (CH₃COOH), glycolaldehyde (HCOCH(OH)), and methyl formate (HCOOCH₃), because of their potential role in the origin of life (e.g. Wächtershäuser 2000, Chyba & Hand 2003). Glycolaldehyde, the simplest of the monosaccharide sugars, has first been detected by Mehringer et al. (1976) towards Sgr B2(N-LMH), and its most recently determined column density in that source is 5.9 × 10¹³ cm⁻² (Hallen et al. 2006). It has been recently observed outside the galactic center by Beltérn et al. (2009) towards the hot molecular core G31.41+0.31, with the emission coming from the hot and dense region closest to the protostars. In addition, Crovisier et al. (2004) and Despois et al. (2005) presented an upper limit for glycolaldehyde abundance in the comet Hale-Bopp. Acetic acid shares the C=O backbone with glycine, from which it differs by an amino group (NH₂). First detected by Mehringer et al.
in Sgr B2(N-LMH), acetic acid shows a column density of $6.1 \times 10^{15}$ cm$^{-2}$ in that region (Remijan et al. 2002). Finally, methyl formate, discovered towards Sgr B2(N) by Brown (1975), has been also observed in other star forming regions of both high (MacDonald et al. 1996; GBB et al. 2001) and low (Remijan & Hollis 2006) mass, towards a proto-planetary nebula (Remijan et al. 2005), and in comets (Bockelée-Morvan et al. 2000; Despois et al. 2002; Remijan et al. 2006). The column density of methyl formate in Sgr B2(N-LMH) is $1.1 \times 10^{17}$ cm$^{-2}$ (Liu et al. 2001).

Sgr B2(N-LMH) is the only source where all three of these isomers have been observed. However, while acetic acid and methyl formate are concentrated in Sgr B2(N-LMH), glycolaldehyde isomers have been observed. However, while acetic acid and methyl formate are concentrated in Sgr B2(N-LMH), glycolaldehyde appears to be more diffusely spread through Sgr B2(N) (Hollis et al. 2001). This behaviour is generally shared by other aldehydes (Snider 2006). In comets only methyl formate has been observed so far.

An understanding of molecular structure, spectroscopy, and photo-absorption processes may be of critical importance in interpreting current observations. In particular, the lifetime of cometary molecules versus photo-destruction is a basic parameter for all cometary studies: as a matter of fact any error in the photodissociation rate translates linearly into an error on the abundance derived in the cometary nucleus. It is also needed for chemical modeling of planetary atmospheres. In this work, we derive electronic and vibrational properties of the isomer triplet C$\mathrm{H}_2\mathrm{O}_2$ described above using the Density Functional Theory (DFT). Photo-destruction rates for acetic acid and methyl formate were derived by Crowtshier (1994) using old laboratory absorption data published by Suto et al. (1988), while the rate for glycolaldehyde is just an estimate. In section 2, we present a brief outline of the method, together with a description of computational settings and results. Section 3 contains the application of these results to cometary photochemistry, while discussion and conclusions are in the last section.

2 THEORY AND RESULTS

We used DFT (e.g., Jones & Gunnarsson 1989) for the calculation of the equilibrium geometry of the electronic ground state and of the vibrational spectrum. We then applied the time-dependent extension of the theory (TD-DFT, Marques & Gross 2004) to compute the electronic excited states and the resulting photo-absorption spectrum for each molecule.

To obtain the ground–state optimised geometries we used the quantum chemistry program package TURBOMOLE (Ahrichs et al. 2007). Technical details about the specific choice of density functional and atomic basis set can be found in the Appendix.

After geometry optimisation, we performed the vibrational analysis obtaining energies and intensities of the normal modes of vibration in the harmonic approximation. Vibrational transitions for fundamental configurations of the isomer triplet are given in the Appendix.

Finally, keeping fixed the ground state geometries obtained above, we computed the photo-absorption cross-section for each molecule. We used two different implementations of TD–DFT in the linear response regime, in conjunction with different representations of the Kohn–Sham wavefunctions:

(i) the real–time propagation scheme using a grid in real space (Yabana & Bertsch 1999), as implemented in the OCTOPUS computer program (Marques et al. 2003).

(ii) the frequency–space implementation (Bauerschmitt & Alrichs 1996) based on the linear combination of localised orbitals, as given in TURBOMOLE.

In the real–time propagation scheme (i) the whole photo-absorption cross-section of the molecule, up to the far–UV, is obtained at once, which is particularly convenient for astrophysical applications. Technical details about the TD–DFT formalism implemented in the OCTOPUS program are reported in the Appendix; the resulting spectra are displayed in Fig. 1.

In the most widely used frequency–space implementation (ii) the poles of the linear response function correspond to vertical excitation energies and the pole strengths to the corresponding oscillator strengths. With this method computational costs scale steeply with the number of required transitions; electronic excitations are thus usually limited to the low–energy part of the spectrum. Table 1 shows that both combinations BP–TZVP and B3LYP–TZVP (see the Appendix for the nomenclature) provide a similarly good agreement between the computed transitions and the experimental results available for acetic acid at room temperature (Limão-Vieira et al. 2006), while BP–TZVP results in a much closer agreement with the experimental results for glycolaldehyde (Karanananand et al. 2007). We therefore chose to use the BP functional for all of our calculations. The resulting BP–TZVP absorption spectra for the three molecules, shown in Fig. 2, are obtained as a superposition of Gaussian functions with fixed arbitrary widths of 0.8 eV. The kind of calculations we performed only yield the positions and intensities of vertical, pure electronic transitions, and therefore give no information on band widths. However, in the available gas–phase spectra of acetic acid (Limão-Vieira et al. 2006) and glycolaldehyde (Karanananand et al. 2007) bands up to $\sim 10$ eV show broad profiles with a full width at half maximum of about 0.8 eV produced by the convolution of unresolved vibronic structure and the natural width of the transitions. The list of first 50 excited states and transition intensities is given in the Appendix.

The two TD–DFT implementations produce compatible results in the low–energy region, i.e. up to about 10 eV, while they tend to diverge significantly at higher energies. However, TD–DFT, as a method, is known to yield dependable results for individual transitions only for excitation energies up to the ionisation energy, which indeed is close to $\sim 10$ eV for all three isomers.

The real–time real–space implementation, on the other hand, has been demonstrated to yield good results for the overall density of electronic transitions even at high energies, with the caveat that single peaks of fine structure are meaningless (see e. g. Marques et al. 2003), at energies $\geq 10$ eV only the envelope of the spectra calculated by octopus is expected to be accurate, while the resolved fine structure is largely due to standing waves in the finite simulation box, is strongly dependent on the size of the box and is only partly quenched by the absorbing boundary conditions we adopted. Since we use these spectra at high energies to compute absorption rates in a continuous spectrum (see Sect 3), the effect of spurious fine structure averages out when integrating over ranges of many eVs, making these theoretical spectra quite adequate for their intended purpose.

3 PHOTOCHEMISTRY

Photo-absorption rates of the three isomers are computed by means of the relation

$$\beta (s^{-1}) = \int_{\Delta E} S_{\odot}(E) \sigma(E)dE$$

(1)
Table 1. Comparison between the computed electronic transitions as obtained using different exchange–correlation functionals and the experimental data reported by Limão-Vieira et al. (2006) for acetic acid and by Karunanandan et al. (2007) for glycolaldehyde.

|          | B-P/TZVP | B3LYP/TZVP | EXP  |
|----------|----------|------------|------|
| acetic acid | 5.60     | 5.84       | 6.09 |
|           | 6.93     | 7.49       | 7.22 |
|           | 8.20     | 8.30       | 8.15 |
|           | 8.25     | 8.50       | 8.35 |
|           | 9.20     | 9.25       | 8.82 |
|           | 10.07    | 10.54      | 10.29|
| Average relative error (%) | 3.4       | 3.2        |      |
| glycolaldehyde | 4.49     | 5.61       | 4.51 |
| Average relative error (%) | 0.4       | 24.4       |      |

where, for the sake of comparison, \( S_\alpha (E) \) is the solar spectrum at 1 AU provided by Huebner et al. (1992), and \( \sigma (E) \) the photo–absorption cross–section for a given molecule. The resulting photo–absorption rates are reported in Table 2 where we also show results for a radiation density expected in a photodissociation front near an OB star (Draine & Bertoldi 1996). We also list separately the contribution of the Lyα line in the solar spectrum to the absorption rates, which is of the order of \( \sim 15\% \) of the total, in agreement with previous estimates of Crovisier (1994). As evident in Fig 2 and Table 2 glycolaldehyde presents a relatively strong band near 4.5 eV, at an energy \( \sim 2 \) eV lower than the first transitions of comparable intensity in the other two members of the triplet (namely \( \sim 6.9 \) eV for acetic acid and \( \sim 7.5 \) eV for methyl formate). Since the solar spectral distribution decreases very steeply in this energy range, the estimated glycolaldehyde photo–absorption rate in the solar radiation field is by and large dominated by this single band, and results to be more than 100 times larger than those of acetic acid and methyl formate. This effect is not present for photo–absorption rates estimated for the acetic acid and by Crovisier et al. (2004). Photons absorbed in the lower energy bands of \( \text{CH}_2\text{OCHO} \) lead to a photodestruction rate \( \beta_{\text{CH}_2\text{OCHO}} \) \( \sim 3 \times 10^{-5} \) s\(^{-1}\) (cf. Table 2), much lower than the rate assumed in Crovisier et al. (2004).

In a recent analysis of ice composition in comet Hale–Bopp, Crovisier et al. (2004) quote a photodestruction rate (at 1 AU) for acetic acid \( \beta_{\text{CH}_3\text{COOH}} \) \( = 5.1 \times 10^{-5} \) s\(^{-1}\). This value was provided by Crovisier (1994), that also reported a rate \( \beta_{\text{CH}_3\text{OHCO}} = 4.7 \times 10^{-5} \) s\(^{-1}\) for methyl formate. The photodestruction rate of glycolaldehyde is unknown and assumed to be \( 1 \times 10^{-4} \) s\(^{-1}\) at 1 AU (Crovisier et al. 2004). Assuming a unit photodestruction yield and a factor of 2 indetermination in the calculations, our photodestruction rates are consistent with Crovisier (1994) values, that were based on old laboratory data provided by Suto et al. (1988). The case of glycolaldehyde is different, since most of photo–absorption is produced by the band at 4.5 eV, that appears to be too low in energy to provide a unit photodestruction yield. As a consequence, photodestruction channels are activated just via absorption in the high–energy bands, leading to a photo–absorption rate \( \beta_{\text{CH}_2\text{OCHO}} \) \( \sim 3 \times 10^{-5} \) s\(^{-1}\) (cf. Table 2), much lower than the rate assumed in Crovisier et al. (2004).

Photons absorbed in the lower energy bands of \( \text{CH}_2\text{OHCHO} \)
may be resonantly scattered, or, if the molecule undergoes internal conversion, re-emitted in the IR range. Using an adapted version of a Monte Carlo model developed for emission by polycyclic aromatic hydrocarbons, which is indeed assumed to be pumped by complete internal conversion of the energy absorbed via electronic transitions in the visible and UV (Mulas 1998), we have constructed the expected IR emission by glycolaldehyde, powered by the solar flux at 1 AU, assuming all the energy absorbed in the band at 4.5 eV to be emitted in the IR. The emission coefficient (per molecule) is reported in Fig. 3.

We have thus far considered photo-absorption rates in the standard solar radiation field at 1 AU (for the “quiet” Sun), which are relevant for the photochemistry of glycolaldehyde, acetic acid and methyl formate in our present solar system. However, from the study of stellar proxies for the Sun it appears that young solar type stars emit high energy photons at a level three to four orders of magnitude higher than the present-day Sun, both during the pre-main sequence phase when the emission is dominated by intense daily or weekly flares (Favata et al. 2005), and during the first phases of the main sequence (Micela 2002). Therefore, chemical evolution can only be understood within the context of the evolving stellar radiation environment.

Without addressing the problem of molecular survival in a disk (e.g. Visser et al. 2007), we estimate the effect of the extreme UV emission (roughly the spectral range between 13 and 100 eV) from solar–type stars of different ages, exploiting the emission of six stars from the Sun in Time program (Guinan & Ribas 2002), whose fluxes are assumed to describe the evolution of the Sun’s emission (Ribas et al. 2005). Since Ribas et al. (2005) reported integrated fluxes (at 1 AU), the rate coefficients are approximated by

\[
\beta(t) \left( s^{-1} \right) = \frac{\langle \sigma \rangle_{\Delta E} S_E(t, \Delta E)}{\Delta E}
\]

where \( \langle \sigma \rangle_{\Delta E} \) is the average over the energy range \( \Delta E \) of the photo-absorption cross-sections shown in Fig. 1. \( \Delta E \) intervals are taken from Ribas et al. (2005). Table 4. Beyond 100 eV a collective description of the molecule is not anymore necessary, since X-ray absorption cross-sections can be closely approximated by adding the atomic cross-sections of individual atoms bound in the molecule (e.g., Cecchi-Pestellini et al. 2004). We therefore do not perform calculations in the X-ray energy range here. Results are reported in Fig. 3. It is evident that the high energy tail of stellar spectrum provides a significant enhancing of photo-absorption rates for acetic acid and methyl formate. Glycolaldehyde photon absorption rates are not changing too much, although the extreme UV contribution to photo-absorption is comparable to absorption in the near UV band at \( \sim 4.5 \) eV. In general, due to the increase in the stellar high energy component, the photo-destruction rates of the isomer triplet members increase roughly two order of magnitude in the environment of a young solar–like star.

4 DISCUSSION AND CONCLUSIONS

In this work we studied the photo-physics of the acetic acid and its isomers, glycolaldehyde and methyl formate. Computed photo-absorption rates are consistent with literature data for acetic acid and methyl formate. In the case of glycolaldehyde, for which the photo-absorption was completely missing up to the far–UV, our calculations indicate that photodestruction is slower than in

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Table 2. Photo-absorption rates (\( s^{-1} \)) for the isomer triplet \( \text{C}_2\text{H}_4\text{O}_3 \) in two radiation fields, assuming a band FWHM of \( \sim 0.8 \) eV based on available experimental data (Limão-Vieira et al. 2006; Karunanandan et al. 2007). The first column lists the computed absorption rates for the whole solar spectrum at 1 AU (Huebner et al. 1992), the second column in the solar Lyα line at 1 AU (Huebner et al. 1992), the third in a photodissociation front near an OB star (Draine & Bertoldi 1996).

| Species          | Solar flux at 1 AU (Huebner et al. 1992) | Solar Lyα at 1 AU (Huebner et al. 1992) | Unshielded OB field (Draine & Bertoldi 1996) |
|------------------|------------------------------------------|----------------------------------------|--------------------------------------------|
| Acetic acid      | CH₃COOH                                  | 6.7(-5)(1)                             | 9.7(-6)                                   | 2.9(-9)                                   |
| Glycolaldehyde(2)| HCOCH₂OH                                 | 9.6(-3)                                | —                                         | 1.3(-10)                                  |
| Methyl formate   | HCOOCH₃                                  | 2.8(-5)                                | 3.7(-6)                                   | 2.9(-9)                                   |

(1) \( 1.8(-5) = 1.8 \times 10^{-5} \).
(2) First row: photo-absorption rates due to the first two absorption bands (up to \( \sim 4.5 \) eV); second row: photo-absorption rates due to the remaining bands.

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Figure 3. IR emission spectrum of one glycolaldehyde molecule at 1 AU from the quiet sun, estimated by assuming that all the energy absorbed in electronic transitions is converted in vibrational excitation and subsequently reradiated (see text). Emission in each vibrational band is represented by a vertical bar whose abscissa corresponds to the wavelength of the vibrational mode and whose height equals the calculated emission intensity.
the other two members of the isomer triplet. However, the overall photo–absorption rate is much larger, and likely to produce either resonant scattering or IR emission powered by near UV solar photons.

Sgr B2 observations of glycolaldehyde show that, unlike acetic acid and methyl formate, its emission is extended in the surrounding molecular cloud. Such a behaviour, that has not been understood so far (e.g. Chengalur & Kanekar 2003), is part of a more general problem involving differentiation in isomers, such as e.g., isocyanide isomers (CH$_3$CN, CH$_2$NC). The behaviour of this isomer triplet has also recently been discussed by Lattelais et al. (2009) as a notable exception to what was called the “Minimum Energy Principle” (MEP) whereby whenever several isomers are possible for a given formula, their observed abundances are in order of binding energy. The authors made the educated guess that this anomaly is first created by differences in the chemical pathways leading to the formation of glycolaldehyde, acetic acid and methyl formate on the surfaces of dust grains, and then preserved due to large energy barriers for the conversion among them. Our results may help to shed some more light on this problem. Although both hot core and the embedding molecular cloud are dark regions, cosmic–rays provide a source of UV photons at high visual extinctions by exciting molecular hydrogen in the Lyman and Werner bands (Prasad & Tarafdar 1983). This locally generated photon flux typically has fluences lower than 10,000 photons cm$^{-2}$ s$^{-1}$ (Cecchi-Pestellini & Aiello 1992), and may produce important chemical effects (Gredel et al. 1989; Bennett & Kaiser 2007). However, the similarity in both the photo–absorption cross–sections and ionisation potentials of the three species makes chemical differentiation due to selective photo–destruction unlikely. As a possible explanation of the extended spatial scale of glycolaldehyde, we consider the possibility of a slow, selective isomerisation, i.e. the possibility that a species may convert itself into another member of the C$_2$H$_4$O$_2$ triplet by interacting with the radiation field. This would imply an isomerisation mechanism which operates on a timescale which is much longer than the typical lifetime of a hot core (few times 10$^5$ years, Williams et al. 2000), but still short enough to be effective on the timescale of the lifetime of a molecular cloud (10$^6$ years, Wilner et al. 2001). In this framework, hot cores would reflect the relative abundances among the isomers as created by their production mechanisms, whereas in the molecular cloud the isomers would “relax” to the most stable one, namely glycolaldehyde, in agreement with the MEP.

Isomerisation may be induced by the absorption of radiation essentially in two main ways; following the absorption of a UV–visible photon the molecule could

(i) move to an electronic state whose energy surface presents a minimum close to the equilibrium configuration of another isomer;

(ii) convert, via one or more non–radiative transitions, a substantial part of the electronic excitation energy into vibrational energy allowing the overcoming of the isomerisation potential barriers.

The present data do not allow a discrimination between the two cases. Moreover, the analysis of the first isomerisation channel would require a detailed study of the energy hypersurfaces in the excited states accessible with photons generated by the Prasad & Tarafdar (1983) mechanism. Therefore, we will discuss qualitatively the second isomerisation process. We assume that every absorption heats up the molecule in a time scale characteristic of electronic transitions ($\sim 10^{-8}$ s), that then decays via non–radiative transitions ($\lesssim 10^{-10}$ s). The electronic excitation energy can be uniquely released by a cascade of vibrational transitions. We also assume, as simplifying hypotheses, that (i) all the excitation energy is converted in vibrational excitation, (ii) that there exist only one isomerisation barrier, (iii) that the excitation energy is far greater than this barrier and (iv) that the isomerisation rate is high above the threshold and zero below it. Then every time a molecule absorbs an UV–visible photon, it will be vibrationally heated. If the energy of the absorbed photon is above the isomerisation threshold, the molecule will establish a statistical equilibrium, in which the probability to find it in one of the isomeric configurations will be proportional to the density of vibrational states at the given energy. The molecule will then cool down, in timescales of the order of a second, with a vibrational cascade (all vibrational modes are IR–active for all three isomers). Its energy will thus eventually fall below the barrier for the isomerisation.

Table 3. Total energies for the three species considered. The second column shows the energy difference $\delta$ in comparison with glycolaldehyde.

| Species        | Total energy (eV) | $\delta$ (eV) |
|----------------|------------------|---------------|
| Glycolaldehyde | -6231.83         | /             |
| Methyl formate | -6232.32         | 0.49          |
| Acetic acid    | -6233.04         | 1.21          |

Stein & Rabinovitch (1973), is shown in Fig. 5. The distances among the curves are due to the differences in the total energies shown in Table 3 and the differences in the frequencies of the vibrational modes.
The naive model drawn above would essentially turn all isomers into glycolaldehyde as soon as they absorb an UV photon. However, UV absorption, in the weak radiation field produced by cosmic-ray induced H$_2$ fluorescence, occurs on timescales of the order of $\gtrsim 50,000$ years, that are comparable to the lifetime of hot cores. As consequence, MEP should not operate effectively in the hot core phase.

Therefore, one of the simplistic assumption in the naive model must be incorrect, slowing down isomerisation. Either internal conversion has a low quantum yield for these molecules, meaning that they get vibrationally heated only a small fraction of the times they absorb an UV photon; or isomerisation is not fast whenever vibrational excitation is sufficient to overcome reaction barriers, due to the morphology of the molecular potential energy surface. The latter can easily be the case if the unimolecular reaction path(s) to isomerisation are narrow and complicated in terms of phase space of the ions.

Of course, conversion among isomers needs not to proceed only via unimolecular reactions induced by radiation: any conversion reaction including chemical reactions with abundant enough partners and without activation barriers could equally well operate on the right timescales to fulfil the MEP in molecular clouds but not in hot cores. A thorough study of the potential energy surface of these three isomers, including the reaction paths connecting them, is called for in order to further progress in the understanding of this observational puzzle.

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APPENDIX

We performed all calculations in the framework of Density Functional Theory, using the quantum chemistry packages TURBOMOLE (Ahlrichs et al. 2007) and OCTOPUS (Marques et al. 2003).

4.1 Geometry optimizations

To find the electronic ground–state geometry of the three molecules considered, we first tested different combinations of density functional and basis set exploiting TURBOMOLE, to identify the most suitable one for our purposes. To this aim, we compared the experimental ground state geometry of the acetic acid (Limão-Vieira et al. 2006) with the one we obtained using the combinations: BP–SV(P), BP–TZVP, B3LYP–SV(P), and B3LYP–TZVP (see the TURBOMOLE manual and references therein for precise definitions of basis sets and functionals). All combinations are in good agreement with experimental and previous theoretical data (Limão-Vieira et al. 2004), with the hybrid functional B3LYP showing a very slightly lower relative average error. This is not unexpected, since this functional is known to produce good results with other classes of organic molecules (e.g., Martin et al. 1996). Considering that both functionals show the same relative average error with both basis sets, we chose the larger TZVP basis set, since this is suggested by the TURBOMOLE manual as the default to get reliable quantitative results.

4.2 Vibrational properties

For our modelling purposes, we calculated the harmonic vibrational frequencies of acetic acid, glycolaldehyde, and methyl formate. All calculations were performed at the BP/TZVP level using the TURBOMOLE program package, and resulted compatible with previously published results (Limão-Vieira et al. 2006; Senent 2004; Senent et al. 2005). The absolute intensities $S$ of the IR–active modes are given in units of km mol$^{-1}$. For each vibrational mode of frequency $\tilde{\nu}$ expressed in cm$^{-1}$ the corresponding Einstein A coefficients for spontaneous emission can be computed as:

$$A (\text{s}^{-1}) = \frac{8\pi}{N_A c} \tilde{\nu}^2 S \simeq 1.2512 \times 10^{-7} \left( \frac{\tilde{\nu}}{\text{cm}^{-1}} \right)^2 \left( \frac{S}{\text{km mol}^{-1}} \right) \text{mol cm}^{-2} \text{km s}^{-1}.$$ 

$N_A$ being the Avogadro’s constant and $c$ the velocity of light.

4.3 Electronic spectra

In the real–time implementation of TD–DFT as given in OCTOPUS, the time–dependent Kohn–Sham equations are directly integrated in real time and the wavefunctions are represented by their discretised values on a spatial grid. The static Kohn–Sham wavefunctions are perturbed by an impulsive electric field and propagated for a given finite time interval. In this way, all of the frequencies of the system are excited. The whole absolute absorption cross–section $\sigma(E)$ then follows from the dynamical polarisability $\alpha(E)$, which is related to the Fourier transform of the time–dependent dipole moment of the molecule. The relation is:

$$\sigma(E) = \frac{8\pi^2 E}{hc} \Im \{\alpha(E)\},$$

\footnote{www.turbomole.com}
where $h$ is Planck’s constant, $\Im \{ \alpha(E) \}$ is the imaginary part of the dynamical polarisability, and $c$ the velocity of light in vacuum.

We performed the OCTOPUS calculations using Becke (1988) exchange and Perdew (1986) correlation functionals. The ionic potentials are replaced by norm-conserving pseudo-potentials (Trouillet & Martins, 1991). We used a grid spacing of 0.12 Å and a box size of 6.8 Å, determined by convergence tests on ground state properties and on the photo-absorption spectrum at energies $\lesssim 10$ eV. This box size ensures that each atom is at least 4 Å away from its edges. We furthermore added a 1 Å thick absorbing boundary, which partially quenches spurious resonances due to standing waves in the finite simulation box used to confine the molecules (Yabana & Bertsch, 1993; Marques et al., 2003). We used a total time integration length $T = 20 \, \hbar / c$, corresponding to an energy resolution of $\hbar / T = 0.05$ eV. For the numerical integration of the time evolution we used a time step of $0.0008 \, \hbar / c$, which ensured energy conservation with good accuracy, within numerical noise.

In the most widely used frequency-space TD-DFT implementation, the poles of the linear response function correspond to vertical excitation energies and the pole strengths to the corresponding oscillator strengths. With this method computational costs scale steeply with the number of required transitions and electronic excitations are thus usually limited to the low-energy part of the spectrum. The frequency-space TD-DFT calculations with TURBOMOLE were performed at the BP/TZVP level of theory, since this showed the best agreement with experimental data. We report in following Table 4 the first 60 singlet–singlet electronic transitions of the three molecules under study with the corresponding oscillator strengths.

### Table 4. Vertical electronic transitions (eV) and corresponding oscillator strengths of the three molecules considered, as computed at the BP/TZVP level using the TURBOMOLE program package.

| Acetic acid | Glycolaldehyde | Methyl formate |
|------------|----------------|----------------|
| 5.70       | 7.3(-4)        | 4.46           |
| 6.94       | 5.4(-2)        | 4.49           |
| 7.74       | 4.7(-3)        | 6.43           |
| 8.25       | 2.6(-3)        | 7.34           |
| 8.37       | 1.3(-1)        | 7.35           |
| 8.82       | 1.9(-4)        | 5.77           |
| 9.05       | 3.3(-3)        | 7.61           |
| 9.08       | 1.4(-4)        | 8.48           |
| 9.16       | 8.1(-4)        | 8.50           |
| 9.27       | 2.9(-3)        | 8.63           |
| 9.33       | 1.2(-1)        | 8.83           |
| 9.88       | 1.1(-1)        | 8.90           |

REFERENCES

Ahrlich R., Furch F., Hättig C., Klopper W., Sierka M., Weigend. F., 2007, TURBOMOLE, Version 5.9.1
Bauerschmitt R., Ahrlich R., 1996, Chem. Phys. Lett., 256, 454
Buckelée-Morvan D., Lis D. C., Wink J. E., Despois D., Crovisier J., Bachiller R., Benford D. J., Biver N., Colom P., Davies J. K., Gérard E., Germain B., Houde M., Mehringer D., Moreno R., Paubert G., 2000, A&A, 353, 1101
Brown R. D., Crofts J. G., Godfrey P. D., Gardner F. F., Robinson B. J., Whiteoak J. B., 1975, ApJ, 197, L29
Cecchi-Pestellini C., Aiello S., 1992, MNRAS, 258, 125
Chengalur J. N., Kanekar N., 2003, A&A, 403, L43
Chyba C. F., Hand K. P., 2005, ARA&A, 43, 31
Crovisier J., 1994, J. Geophys. Res., 99, 3777
Crovisier J., Buckelée-Morvan D., Colom P., Biver N., Despois D., Lis D. C., the Team for target-of-opportunity radio observations of comets 2004, A&A, 418, 1141
Despois D., Biver N., Buckelée-Morvan D., Crovisier J., 2005, in Lis D. C., Blake G. A., Herbst E., eds, Astrochemistry: Recent Successes and Current Challenges Vol. 231 of IAU Symposium, Observations of Molecules in Comets, p. 469
Draine B. T., Bertoldi F., 1996, ApJ, 468, 269
Favata F., Flaccomio E., Reale F., Micela G., Sciortino S., Shang H., Stassun K. G., Feigelson E. D., 2005, ApJS, 160, 469
Gibb E., Nummelin A., Irvine W. M., Whittet D. C. B., Bergman F., 2000, ApJ, 545, 309
Gredel R., Lepp S., Dalgarno A., Herbst E., 1989, ApJ, 347, 289
Guinan E. F., Ribas I., 2002, in Montesinos B., Gimenez A., Guinan E. F., eds, The Evolving Sun and its Influence on Planetary Environments Vol. 269 of Astronomical Society of the Pacific Conference Series, Our Changing Sun: The Role of Solar Nuclear Evolution and Magnetic Activity on Earth’s Atmosphere and Climate. p. 85
Halfen D. T., Apponi A. J., Woolf N., Poll R., Ziurys L. M., 2006, ApJ, 639, 237
Hartquist T. W., Williams D. A., eds, 1998, The Molecular Astrophysics of Stars and Galaxies. Clarendon Press, Oxford
Hollis J. M., Lovas F. J., Jewell P. R., 2000, ApJ, 540, L107
Hollis J. M., Vogel S. N., Snyder L. E., Jewell P. R., Lovas F. J., 2001, ApJ, 554, L81
Huebner W. F., Keady J. L., Lyon S. P., 1992, Astrophys. Spa. Sci., 195, 1
Jones R. O., Gunnarsson O., 1989, Reviews of Modern Physics, 61, 689
Karunanandan R., Hölscher D., Dillon T. J., Horowitz A., Crowley J. N., Vereecken L., Peeters J., 2007, J. Phys. Chem. A, 111, 897
Lattelais M., Pauzat F., Ellinger Y., Ceccarelli C., 2009, ApJ, 696, L133
Limão-Vieira P., Giuliani A., Delwiche J., Parafita R., Mota R., Duflot D., Flament J.-P., Drage E., Cahillane P., Mason N., Hoffmann S., Hubin-Franskin M.-J., 2006, Chem. Phys., p. 339
Liu S.-Y., Mehringer D. M., Snyder L. E., 2001, ApJ, 552, 654
MacDonald G. H., Gibb A. G., Habing R. J., Millar T. J., 1996, A&AS, 119, 333
Marques M. A. L., Castro A., Bertsch G. F., Rubio A., 2003, Comp. Phys. Comm., 151, 60
Marques M. A. L., Gross E. K. U., 2004, Ann. Rev. Phys. Chem., 55, 427
Martin J. M. L., El-Yazal J., Francois J., 1996, J. Phys. Chem., 100, 15358
Mehringer D. M., Snyder L. E., Miao Y., Lovas F. J., 1997, ApJ, 480, L71
Miao Y., Snyder L. E., 1997, ApJ, 480, L67
Micela G., 2002, in Montesinos B., Gimenez A., Guinan E. F., eds, The Evolving Sun and its Influence on Planetary Environments Vol. 269 of Astronomical Society of the Pacific Conference Series, Evolution of Stellar Coronal Activity on the Main Sequence. p. 107
Mulas G., 1998, A&A, 338, 243
Perdew J. P., 1986, Phys. Rev. B, 33, 8822
Prasad S. S., Tarafdar S. P., 1983, ApJ, 267, 603
Remijan A., Snyder L. E., Liu S.-Y., Mehringer D., Kuan Y.-J., 2002, ApJ, 576, 264
Remijan A. J., Friedel D. N., de Pater I., Hogerheijde M. R., Snyder L. E., A’Hearn M. F., Blake G. A., Dickel H. R., Forster J. R., Kraybill C., Looney L. W., Palmer P., Wright M. C. H., 2006, ApJ, 643, 567
Remijan A. J., Hollis J. M., 2006, ApJ, 640, 842
Remijan A. J., Wyrowski F., Friedel D. N., Meier D. S., Snyder L. E., 2005, ApJ, 626, 233
Ribas I., Guinan E. F., Güdel M., Audard M., 2005, ApJ, 622, 680
Senent M. L., 2004, J. Phys. Chem. A, 108, 6286
Senent M. L., Villa M., Meléndez F. J., Domínguez-Gómez R., 2005, ApJ, 627, 567
Snyder L. E., 2006, Proc. Natl. Ac. Sci., 103, 12243
Stein S. E., Rabinovitch B. S., 1973, J. Chem. Phys., 58, 2438
Suto M., Wang X., Lee L. C., 1988, J. Phys. Chem., 92, 3764
Trouiller N., Martins J. L., 1991, Phys. Rev. B, 43, 1993
Visser R., Geers V. C., Dullemond C. P., Augereau J., Pontoppidan K. M., van Dishoeck E. F., 2007, A&A, 466, 229
Wächtershäuser G., 2000, Science, 289, 1307
Williams J. P., Blitz L., McKee C. F., 2000, Protostars and Planets IV, p. 97
Wilner D. J., De Pree C. G., Welch W. J., Goss W. M., 2001, ApJ, 550, L81
Yabana K., Bertsch G. F., 1999, Int. J. Q. Chem., 75, 55