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Valence and lowest Rydberg electronic states of phenol investigated by

synchrotron radiation and theoretical methods

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Abstract

We present experimental high-resolution vacuum ultraviolet (VUV) photoabsorption spectra of phenol covering for the first time the full 4.3–10.8 eV energy-range, with absolute cross-sections determined. Theoretical calculations on the vertical excitation energies and oscillator strengths were performed using time-dependent density functional theory (TD-DFT) at the equation-of-motion coupled cluster method restricted to single and double excitations (EOM-

CCSD) level. These have been used in the assignment of valence and Rydberg transitions of the phenol molecule. The VUV spectrum reveals several new features not previously reported in the literature, with particular reference to the 6.401 eV transition, which is here assigned to the $3s\sigma/\sigma^*(OH) \leftarrow 3\pi(3a'')$ transition. The measured absolute photoabsorption cross-sections have been used to calculate the photolysis lifetime of phenol in the Earth's atmosphere (0 – 50 km).

Keywords: phenol, C₆H₅OH, electronic excitation, vibrational excitation, Rydberg series, ab initio calculations, photoabsorption, synchrotron radiation

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1. INTRODUCTION

Phenol, C₆H₅OH, has attracted considerable attention from the international scientific community because of its relevance to industry but it also serves as a model system to biologically related molecules (e.g. amino acids). The conversion of renewable biomass into biofuels has shown to be a non-cooperative procedure (known as recalcitrant) which still imposes serious cost-effective processes in the de-polymerization of biomass into fermentable sugars.^{1,2} Phenol can also be used as a key model for understanding the UV photo-induced dynamics of some amino acids, where O-H bond rupture from the phenol moiety can be regarded as a proton release process (or proton tunnelling) from tyrosine residues which is known to be a prevalent mechanism in a wide range of natural enzymes.³ Another key aspect for studying phenol has been centred on probing non-radiative decay processes (from S_1 to S_0) in aromatic biomolecules,⁴ as well as exploring the role of $\pi\sigma^*$ excited states in photodissociation process allowing therefore to provide distinctively detailed insights into the fragmentation dynamics.⁵ This has been decisive to identify the importance of the strong vibronic coupling via a conical intersection between the S_0 and ${}^1\pi\sigma^*$ potential energy surfaces at longer O–H bond lengths.⁶⁻⁹

Phenol, an aromatic hydrocarbon, is considered an important atmospheric molecule related to urban air pollution as precursor for the formation of photo-oxidants and secondary organic aerosols. The products of the reactions of the photochemical degradation of benzene, toluene and xylene-isomers, with OH radicals in the terrestrial atmosphere also yield phenol production.¹⁰ Moreover, the contribution of aromatic hydrocarbons to ozone formation in Europe was estimated to be as high as 40%,¹¹ whilst Etzkorn et al.¹² have suggested that aromatics are the most important class of hydrocarbons for photochemical ozone formation.

Trost et al.¹³ have reported the room-temperature rate coefficient for reactions of the OH radical with phenol ($k_{OH} = 26.3 \times 10^{-12} \text{ cm}^3$ molecule⁻¹ s⁻¹), showing that this is likely to be

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the main sink mechanism for these molecules in the troposphere. Studies on NO₃-initiated oxidation of phenol at room temperature have been reported by Atkinson et al.¹⁰ with a rate constant value of $(2.1 \pm 0.5) \times 10^{-12}$ cm³ molecules⁻¹ s⁻¹, where reactions proceed via H-atom abstraction from the OH group. Night time reactivity of NO₃ radicals with phenol can be regarded as an important sink mechanism for these aromatic hydrocarbons as well as loss of NO₃, possibly contributing as a source of nitric acid, a key component of acid deposition.^{11,12}

Accurate absorption cross-section values, as well as a comprehensive description of the electronic states of aromatic hydrocarbon compounds, are needed as part of a wider research programme aimed at understanding the spectroscopy of these compounds and their role as trace gases in atmospheric chemical and physical reactions. A thorough literature survey reveals scarce data on the VUV electronic state spectroscopy of phenol in a wide energy range as is fully covered in the present contribution. Indeed, experimental information on phenol is mainly restricted to $244 < \lambda < 283$ nm (4.38 < E < 5.08 eV)¹³ and $200 < \lambda < 290$ nm (4.28 < E < 6.20 eV),¹⁴ with these studies including no assignments of the absorption bands. Kimura and Nagakura¹⁵ report the wavelength region 155–300 nm (4.13–8.00 eV) with four absorption bands assigned to $\pi \leftarrow \pi^*$ transitions.

The theoretical data of Lorentzon et al.¹⁶ covers the energy region (4.53–7.84 eV) with a few assignments of valence states and the first members (n = 3) of the Rydberg series converging to the two lowest ionisation energies. Miura and co-workers¹⁷ have reported oscillator strengths of the HOMO(π) \rightarrow LUMO(π^*) transition through time-dependent density functional theory, while electron energy-loss spectra have been reported by Ari et al.¹⁸ Multiphoton ionisation photoelectron spectroscopy of the phenol ¹B₂ state has been obtained to provide a consistent set of vibrational assignments,¹⁹ Kishimoto et al.²⁰ reported on Penning ionisation electron spectra, and Livingstone and co-workers²¹ have performed detailed studies to investigate the dynamical evolution of the initially prepared S₁ ($\pi\pi^*$)

excited state of phenol through a time-resolved photoelectron imaging technique. The fragmentation dynamics of gas phase phenol following excitation in the wavelength region $279.145 \ge \lambda \ge 260.000$ nm has been investigated by photofragment translational spectroscopy.²² Molecular geometries and vibrational spectra of phenol have been reported by experimental and quantum chemical data²³ as well as by FTIR experiments.²⁴ Rotationally resolved fluorescence excitation spectra of the 4.51 eV electronic transition of phenol in a supersonic jet has been measured.²⁵ Finally, we note comprehensive experimental and theoretical studies on the excited electronic states of phenol²⁶ providing the first investigation of all singlet and triplet states up to the first ionisation potential. This study prefaced substantive electron scattering investigations with phenol,²⁷⁻³¹ that illustrated the important role of phenol's excited and ionised states in describing its scattering behaviour.

Recently we have explored the valence and Rydberg electronic state spectroscopy of a related isoelectronic aromatic hydrocarbon, toluene $(C_6H_5CH_3)$,³² relevant to urban pollution where fossil fuel, industrial and biomass burning are the common sources of emissions. Thorough knowledge gained on the spectroscopic nature of toluene assists in the assignment of the absorption bands of phenol.

In this paper we report the results of an extensive study of the electronic state spectroscopy of phenol by high resolution VUV photoabsorption spectroscopy and *ab initio* theoretical calculations of the vertical excitation energies and oscillator strengths for the neutral electronic transitions. In the next section we provide a brief summary of the structure and properties of phenol. In addition to identifying the optical electronic transitions of phenol, the present work provides reliable photoabsorption cross-sections in the range 4.3–10.8 eV. In Section 3 we present the computational methods and in Section 4 a brief discussion of the experimental details. Section 5 is devoted to the results and discussion with a comparison with other absolute photoabsorption cross-sections. The absolute photoabsorption cross-

sections are used to calculate photolysis rates in the Earth's atmosphere up to the limit of the stratopause (50 km). Finally some conclusions that can be drawn from this study are given in Section 6.

2. BRIEF SUMMARY OF THE STRUCTURE AND PROPERTIES OF PHENOL

Phenol has symmetry C_S in the electronic ground state and the symmetry species available are A' and A". The calculated electron configuration of the \tilde{X} ¹A' ground state according to Jones el al.²⁶ is: (a) core orbitals $(1a')^2 (2a')^2 (3a')^2 (4a')^2 (5a')^2 (6a')^2 (7a')^2$; (b) valence orbitals $(8a')^2 (9a')^2 (10a')^2 (11a')^2 (12a')^2 (13a')^2 (14a')^2 (15a')^2 (16a')^2 (17a')^2 (18a')^2$ $(1a'')^2 (19a')^2 (20a')^2 (21a')^2 (2a'')^2 (3a'')^2 (4a'')^2$. The highest occupied molecular orbital (HOMO) 4a'', the second highest occupied molecular orbital (HOMO-1) 3a'' and the third highest occupied molecular orbital (HOMO-2) 2a'' have π character and may be labelled 4π , 3π and 2π , respectively. The lowest unoccupied molecular orbitals (LUMO) 5a'' and (LUMO+1) 6a'' are mainly of π^* antibonding character, with (LUMO+1) having a slightly σ^* character along the C–OH bond. Note that the $3s/\sigma^*(OH)$ is labelled (LUMO+1) in Ref. 26, whereas here the LUMO notation is reserved to pure valence MOs. Thus (LUMO+2) in Jones et al.²⁶ is now labelled (LUMO+1).

The earlier studies of Lorentzon et al.¹⁶ have predicted at the CASPT2 (complete active space self-consistent field with second –order perturbation correction) level of theory the valence and 3s, 3p and 3d Rydberg states whereas the recent experimental data on the electron-energy loss spectroscopy²⁶, at moderate electron energy resolution ($\Delta E \sim 70$ meV), did not allow for a more quantitative assignment of the Rydberg-like transitions. Bearing in mind the resemblance between phenol and toluene photoabsorption spectra, we may anticipate for the former an overlap of Rydberg states with valence states resulting in a complex intensity distribution in the electronic spectrum. Rydberg states may be identified

through knowledge of the ionisation energies and the application of quantum defect theory. The three lowest ionisation energies, which are required to calculate the quantum defects associated with transitions to Rydberg orbitals, have been identified experimentally to be at 8.64, 9.32 and $11.57 \text{ eV}.^{20}$

3. COMPUTATIONAL DETAILS

Electronic structure calculations were performed with two different methods (TD-DFT and EOM-CCSD) to determine the excitation energies of the neutral molecules (TABLES I and II) and the lowest ionisation energy. Density Functional Theory (DFT) using the LCωPBE long range corrected functional³³⁻³⁵ has been implemented with the Gaussian 09 code.³⁶ The geometry (FIG. 1) and harmonic frequencies were obtained using Dunning's cc-pVTZ basis set.³⁷ Then the properties of the excited states (energies and oscillator strengths in the length gauge) were obtained using TD-DFT for both triplet and singlet states. The basis set was augmented to aug-cc-pVTZ for all atoms and a set of diffuse (5s5p2d) orbitals, taken from Kaufmann et al.³⁸ was added to the centre of the benzenic ring (aug-cc-pVTZ+R basis set). The nature of the transitions was assessed by visual inspection of the natural transitions orbitals (NTOs),³⁹ computed for each calculated triplet and singlet transition.

Phenol geometry was also obtained at the CCSD(T) level with the Molpro 2012.1 code⁴⁰ with the cc-pVTZ basis set (FIG. 1). The lowest lying singlet transitions were then calculated at the EOM-CCSD (equation-of-motion coupled cluster method restricted to single and double excitations) level, using the aug-cc-pVTZ+R basis set. The nature of each transition was assessed by visual inspection of the corresponding EOM-CCSD natural orbitals for each transition. Finally, the lowest vertical ionization energy was also obtained at the RCCSD(T) level.⁴¹

4. EXPERIMENTAL DETAILS

4.1 PHENOL SAMPLE

The liquid sample used in the VUV measurements was purchased from Sigma-Aldrich, with a stated purity of \geq 99.5%. The sample was degassed by repeated freeze-pump-thaw cycles.

4.2 VUV PHOTOBASORPTION

The high-resolution VUV absorption spectrum of phenol (FIG. 2) was measured at the ASTRID2 storage ring at Aarhus University, Denmark, using the AU-UV beam line. The apparatus used for measurement of photoabsorption spectra has undergone several alterations since that originally described in Eden et al.42 with those modifications being described recently in Ref. 43. Briefly, the gas cell, which allows moderate (up to 80° C) heating of a sample during measurement, has a path length of 15.5 cm and is fitted with a heated 1 Torr Baratron capacitance manometer (Setra model 774). To ensure that the data is free of any saturation effects the absorption cross-sections were measured at an appropriate pressure in the range 0.02–0.43 mbar, with typical attenuations of less than 40%. The light exits the cell through a MgF₂ window, which sets the lower limit of detectable light to 115 nm. A small gap between the photo-multiplier tube (PMT) detector and the absorption cell is evacuated using a scroll pump for measurements below 200 nm. Above 220 nm, air is allowed into this gap to let oxygen absorb higher orders of light (at half the chosen wavelength) which may be passing through the cell. In this way, photoabsorption measurements can be performed with spectrally pure radiation from 115 nm to 340 nm, ensuring artefact-free spectra. Absolute photoabsorption cross-sections (σ) are obtained using the Beer-Lambert law $I_t = I_0 \exp(-n\sigma x)$, where I_t is the radiation intensity transmitted through the gas sample, I_0 is that through the evacuated cell, n is the molecular number density of the sample gas, σ is the absolute photoabsorption cross-section, and x is the absorption path length (15.5 cm). ASTRID2 is operated in a so-called top-up mode, keeping the stored electron beam current (and thus the intensity for a given wavelength) quasi-constant by adding small amounts of current to ASTRID2 to make up for the constant beam decay. The beam current thus varies about 3% during a scan, and this is taken into account by recording and normalizing to an accurately determined beam current.

The accuracy of the cross-section is estimated to be better than \pm 5%. Only when absorption by the sample is very weak ($I_0 \approx I_l$), does the error increase as a percentage of the measured cross-section.

5. ELECTRONIC STATE SPECTROSCOPY: RESULTS AND DISCUSSION

The absolute high resolution VUV photoabsorption spectrum of phenol measured at room temperature is shown in FIG. 2, extending from 3.9 to 10.8 eV and FIGS. 3–7 show expanded views in five subsection energy regions. The major absorption bands can be classified as a mixture of Rydberg series and molecular valence transitions of $(\pi^* \leftarrow \pi)$ character. TABLE I compares the experimental results with the TD-DFT calculations and demonstrates reasonably good agreement. The calculations indicate that the electronic transitions for phenol have mixed valence-Rydberg character. The RCCSD(T) lowest vertical ionisation energy is 8.60 eV (TABLE II) and compares well with Lorentzon et al.¹⁶ (8.21 and 8.56 eV) and Kishimoto et al.²⁰ (8.64 eV). The measured lowest vertical IE of phenol (8.64 eV)²⁰ agrees well with the theoretical prediction.¹⁶ There exists several different numbering schemes of the vibrational modes of phenol in the literature, and we have followed the notation adopted in the studies of Anderson et al.¹⁹ and Nix et al.²² in the following sections.

5.1. VALENCE SPECTROSCOPY OF PHENOL

According to the calculations in TABLES I and II, the absorption bands centred at 4.605, 6.00(4) and 6.736 eV have been assigned to ($\pi^* \leftarrow \pi$) transitions. The calculated wave functions show a mixing between excitations from the HOMO (4π) and HOMO-1 (3π) to the LUMO (π^*) and LUMO+1 (π^*) MO's, and also a $3s\sigma$ (a") Rydberg character for the latter. An identical behaviour has been observed for toluene.²⁷ The calculated transition energies at the EOM-CCSD level (TABLE II) are generally overestimated by about 0.3 eV when compared to experiment, whereas for TD-DFT level of theory (TABLE I) such difference increases on average to 0.5 eV. Thus, in this case the EOM-CCSD results are more reliable than the TD-DFT ones, especially for Rydberg transitions. Nevertheless, the present TD-DFT/LC- ω PBE/aug-cc-pVTZ+R calculations appear to be closer to other theoretical results than the TD-DFT/B3LYP/aug-cc-pVDZ of Ref. 26. The assignments in the lowest absorption band are in reasonable agreement with Nix et al.²² reporting the band at 4.507 eV (275.113 nm), while Ari et al.'s¹⁸ electron energy loss data place the first three bands at 4.59, 5.93 and 6.68 eV, respectively and Kimura and Nagakura¹⁵ at 4.59, 5.82 and 6.70 eV.

The first band is reported here with a maximum absolute cross-section of 18.1 Mb (at 4.605 eV), whereas the second has a maximum of 19.1 Mb (at 6.00(4) eV) and the third at 192.5 Mb (at 6.736 eV) (see TABLES I and II). The feature at 6.401 eV, with a rather weak calculated oscillator strength, $f_L \approx 0.001$, is here tentatively assigned for the first time to $\sigma^*(OH) \leftarrow 3\pi(3a'')$ with the aid of the theoretical calculations (TABLE II). However, a careful look at TABLE II reveals that for this energy transition, other contributions from $3p\sigma/\sigma^*(OH) \leftarrow 4\pi$ and $3s\sigma/\sigma^*(OH) \leftarrow 3\pi$ may be present. The feature at 6.736 eV has been identified as an n = 3p member of a Rydberg series converging to the ionic electronic first excited state of phenol (see Section 5.2), the rather high calculated intensity ($f_L \approx 0.060$ in TABLE II) is primarily due to the pure valence π^* character of the MO. Pure Rydberg

transitions (TABLE VI) with considerable values of oscillator strengths in this energy range are discussed in Sec. 5.2.

VALENCE AND VIBRONIC EXCITATION IN THE RANGE 4.4–5.5 eV

The lowest-lying excited state of phenol $(2\pi^* \leftarrow 3\pi + 1\pi^* \leftarrow 4\pi)$, with a maximum at 4.605 eV, with a local cross-section value of 18.1 Mb (FIG. 3), has been assigned to the S_1 - S_0 $(\pi^* \leftarrow \pi)$ and shows extensive fine structure, which is much better resolved than in previous absolute cross section measurements²² and therefore new assignments are proposed. The vertical energy is in good agreement with the calculated value of 4.99 eV of Jones et al.²⁶ as well as from the present calculations in TABLES I and II, where the EOM-CCSD level of accuracy (4.947 eV, TABLE II) gives a better agreement with the experimental data. The adiabatic transition is observed at 4.508 eV (34.6 Mb) in very good agreement with the value of Nix et al.²² at 4.507 eV (275.113 nm). Vieuxmaire et al.⁴⁴ report an adiabatic excitation energy at 4.612 eV at CAS(10,9)/6-31G** level of theory against the experimental value of 4.507 eV according to Bist et al.⁴⁵ and Berden et al.⁴⁶, the latter being a comprehensive study on the structure and dynamics of internal rotation of phenol. We also note rotationally resolved fluorescence excitation spectra of phenol by Martinez et al.⁴⁷ place the (0-0) transition at 4.507 eV. This absorption band shows resolvable rovibronic structure, and assignments are presented in TABLE III, where they are revisited and compared with the detailed data of Nix et al.,²² with several new features reported here for the first time.

The present high-resolution spectrum reveals that the fine structure is due to several modes (see TABLE III for the proposed detailed assignments), with the main contributions from the oxygen stretching motion mode, v'_{6a} (a') and the O-H torsion mode, v'_{6b} (a'). These modes also appear coupled with other modes. Due to the complexity of such fine structure in the absorption band in FIG. 3, and in order to avoid congestion, we have represented a few

modes only. It is noteworthy that the normal vibrational description may lead to some Fermi resonances, making assignments particularly difficult. The features below the 0^{0}_{0} transition are tentatively assigned to hot-bands based on the previous work of Nix et al.²² The feature at 4.548 eV is tentatively assigned a sequence band of the OH torsion mode, further to Nix et al.²² assignments.

VALENCE AND VIBRONIC EXCITATION IN THE RANGE 5.3-7.0 eV

The lowest valence excitation band in this energy range (FIG. 4) has been assigned to the $(1\pi^* \leftarrow 3\pi + 2\pi^* \leftarrow 4\pi)$ transition in agreement with previous reports.^{16,26} The 00 transition is here proposed at 5.75(5) eV (see TABLE IV). The 1142 cm⁻¹ (0.142 eV) spacing reported by Anderson et al.¹⁹ for phenol excited state is in very good agreement with the mean value of 0.106 eV (FIG. 4 and TABLE IV), which has been assigned to excitation of ring CH in-plane deformation and CC stretching mode, v'_{9a} (a') with 1167 cm⁻¹ (0.145 eV) in the neutral ground state.^{19,22,23,48} The next band with the highest oscillator strength ($f_L \approx 0.61$ at TD-DFT level and ≈ 0.39 at EOM-CCSD), peaking at 6.736 eV, has been assigned to $(1\pi^* \leftarrow$ $3\pi + 2\pi^* \leftarrow 4\pi$) in TABLE I, whereas in the case of toluene it was reported at 6.786 eV and assigned to a mixed Rydberg/valence $(1\pi^* \leftarrow 3\pi + 3p\pi (a'') \leftarrow 2\pi)$.³² We note that the calculated vertical excitation energies are obtained more accurately from the calculations at the EOM-CCSD level of theory rather than the TD-DFT (LC-wPBE/aug-cc-pVTZ+R) level when compared to the experimental values (TABLE II). According to the calculations, the $3p\pi(a') \leftarrow 3\pi(3a'')$ Rydberg transition has a considerable oscillator strength (0.06) at the EOM-CCSD level and contributes to the feature observed in this energy range. A few quanta of oxygen stretching motion mode, ν'_{6a} (a'), are excited (see TABLE V) and the 0^0_0 transition is proposed at 6.461 eV, although we note a weak feature at 6.401 eV. Since the latter lies on an underlying state with a different slope from the rising left-hand side of the most intense absorption feature, we tentatively assign it to the $\sigma^*(OH) \leftarrow 3\pi(3a'')$ transition according to the calculations in TABLE II, although we do not discard the possibility of other transition, from $3p\sigma/\sigma^*(OH) \leftarrow 4\pi$ and $3s\sigma/\sigma^*(OH) \leftarrow 3\pi$, contributing to the spectrum. This energy region exhibits evidence for transitions to Rydberg states at 6.736, 6.91(1) and 7.12(8) eV (see Section 5.2 and Table VI), converging to the ionic electronic first excited and ionic ground states, respectively, and also shows some quanta excitation number of oxygen stretching motion mode, v'_{6a} (a') (TABLE V) with mean excitation energy of 0.056 eV. However, Anderson et al.¹⁹ pointed out that in the neutral excited states of phenol, the strong absorption band is dominated by two fundamental modes assigned to oxygen stretching modes v'_{6a} (a') and v'_{12} (a'), with excitation energies of 526 and 935 cm⁻¹. Nonetheless, the normal mode vibrational description may lead to Fermi resonances, making assignments particularly difficult. Finally we note in FIG. 4 a rather broad background contribution that can be attributed to a pre-dissociative state. A careful look at TABLES I and II shows a rather low oscillator strength (f_L = 0.0002 in TABLE II) transition, assigned to $\sigma^*(OH) \leftarrow 4\pi$ at ~5.75 eV (FIG. 3) and in good agreement with the S₂ ($^{1}\pi\sigma^{*}$) state of Livingstone et al.²¹ calculated at 5.55 eV at the EOM-CCSD level.

5.2 RYDBERG TRANSITIONS

The VUV photoabsorption spectrum above 6.0 eV consists of a few structures superimposed on a diffuse absorption feature extending to the lowest IEs, which are reported here for the first time. The experimental ionisation energy values of Kishimoto et al.²⁰ are used to determine the Rydberg series. The proposed Rydberg structures are labelled in FIGS. 4–7 and listed in TABLE VI. The peak positions have been tested using the Rydberg formula: $E_n = E_i - R / (n - \delta)^2$, where E_i is the ionisation energy, *n* is the principal quantum number of the Rydberg orbital of energy E_n , *R* is the Rydberg constant (13.61 eV), and δ is the quantum defect resulting from the penetration of the Rydberg orbital into the core. The experimental values for the lowest terms of the ns, np, and nd (n = 3) Rydberg series (TABE VI) are in good agreement with the calculations in TABLE II.

The lowest transition energy is assigned to $3s\sigma/\sigma^*(OH) \leftarrow 4\pi$ at 5.75(5) eV for the Rydberg series converging to the ionic electronic ground state, with a quantum defect $\delta = 0.83$ (TABLE VI), accompanied by vibronic structure, which is proposed to be mainly due to excitation of ring CH in-plane deformation and CC stretching mode, v'9 (a'), with an average value of 0.104 eV (FIG. 4 and TABLE IV) (and even of oxygen stretching motion mode, v'6a (a')), in very good agreement with 0.150 eV for the \tilde{X}^2A'' state from Anderson et al.²⁰ However, these authors note that this mode is a weakly active mode in the C₆H₅OH⁺ ion. The higher members of this Rydberg series are proposed to extend to n = 7. Because of the complexity of such fine structure in the absorption band in FIG. 6, and in order to avoid congestion, we have presented only a few modes.

The first members of the np and nd series are associated with peaks at 6.00(4) eV ($\delta = 0.73$) and 6.91(1) eV ($\delta = 0.19$), respectively (TABLE VI). The nd series shows an n = 3 vibrational excitation with two quanta of ring distortion mode, v'_{6a} (a'), with an average value of 0.053 eV (FIG. 6 and TABLE V), in very good agreement with 0.059 eV for the \tilde{X}^2A'' state from Anderson et al.²⁰ The higher members of these Rydberg series, for which the relative intensity decreases, are difficult to assign due to their overlap with other transitions and possible vibronic structure (see TABLE V). Close to the lowest ionic limit, features tentatively assigned to ring distortion mode, v'_{6a} (a') appear quite abundantly.

The first members of the ns, np, and nd series converging to the ionic electronic first excited state of phenol (3a^{"-1}) are associated with the peaks at 5.89(3) eV ($\delta = 1.01$), 6.736 eV ($\delta = 0.70$), and 7.562 eV ($\delta = 0.22$), respectively (TABLE VI). The rather high quantum defect for the 3s Rydberg member is due to the contribution of the valence $\sigma^*(OH)$ state

(TABLE II). The Rydberg transitions (FIG. 6) are accompanied by vibronic structure, which is tentatively attributed to excitation of the v'_{6a} (a') mode (TABLE V), however, the normal mode configuration may also lead to Fermi resonances.

As far as members of the Rydberg series converging to the ionic electronic second excited state are concerned, n = 3 for ns, np and nd are proposed at 8.178, 9.00(4) and 9.70(9) eV, respectively, (TABLE VI) with quantum defects 1.00, 0.70 and 0.30, respectively. Some of the fine structure has been assigned to vibrational excitation involving ring distortion mode v'_{6a} (a'), as well as ring and OH bending character, v'_{8a} (a'), the latter in good agreement with the photoelectron spectrum of Anderson et al.,²⁰ although we were able to assign new features (see TABLE V and FIG. 7).

We have not made any attempt to identify higher members (n > 3) of the Rydberg series, due to the broad and structureless nature of the absorption bands. Note that the clear increase in the absorption with energy, in the range above 9.0 eV, may be related to low-lying pre-dissociative or dissociative excited neutral states.

5.3 ABSOLUTE PHOTOABSORPTION CROSS-SECTIONS AND ATMOSPHERIC PHOTOLYSIS

The present optical measurements were carried out in the pressure range 0.02-0.43 mbar and reveal no evidence for changes in absolute cross-sections or peak energies as a function of pressure, thus we believe the present spectra are free of any saturation effects. Previous absolute VUV photoabsorption cross-sections of phenol are available in the wavelength ranges 244–283 nm (4.38–5.08 eV),¹² 245–285 nm (4.35–5.06 eV),¹³ 200–290 nm (4.28–6.20 eV),¹⁴ and 160–240 nm (5.17–7.75 eV).¹⁵ Etzkorn et al.,¹² Trost et al.,¹³ Grosch et al.,¹⁴ and Kimura and Nagakura¹⁵ reported absolute cross-sections of magnitude 19.8 Mb (275.1 nm, 4.51 eV), ~37 Mb (275 nm, 4.51 eV), ~19.1 Mb (275 nm, 4.51 eV) and

36.1 Mb (6.82 eV), compared to the present value of 34.6 Mb at 4.508 eV. Such differences, as in the case of Etzkorn et al.¹² are mainly attributable to experimental resolution, where a value of 0.15 nm is reported, against the present value of 0.075 nm. We have noted over the last decade an excellent agreement of previous cross-sections measured on the UV1 beam line on ASTRID with the most precise data available in the literature (see Eden et al.⁴⁹ and references therein and more recently Ref. 50). This indicates that the present absolute phenol cross-sections are reliable over the energy range studied (4.3-10.8 eV).

The present absolute cross sections below 6.89 eV (above 180 nm) can be used in combination with solar actinic flux⁵¹ measurements from the literature to estimate the photolysis rate of phenol in the atmosphere from an altitude close to the ground to the stratopause at 50 km. Details of the calculation programme are presented in Ref. 52, in which the quantum yield for dissociation following absorption is assumed to be unity. The reciprocal of the photolysis rate at a given altitude corresponds to the local photolysis lifetime. Photolysis lifetimes of around 24 sunlit hours were calculated at ground level, with less than an hour above 30 km. This shows that phenol molecules can be broken up quite efficiently by UV absorption even at low altitudes. From the ground level up to the tropopause the lifetimes are considerably low, such that photolysis may be a sink mechanism. However, the rate coefficients obtained by Trost et al.¹³ for reactions between the OH radical and phenol have shown that this may provide a main reactive sink mechanism in the troposphere. Moreover, night time reactivity of NO₃ radicals with phenol can be also regarded as an important sink mechanism.¹⁰ Therefore, compared with radical reactions, UV photolysis is not expected to play a significant role in the tropospheric removal of phenol molecules.

6. CONCLUSIONS

The present work provides the first comprehensive study to date of the VUV electronic spectroscopy of phenol and provides the most reliable set of absolute photoabsorption cross-sections between 4.3 to 10.8 eV. The observed valence and Rydberg transitions have been assigned with the help of *ab initio* calculations on the vertical excitation energies and oscillator strengths at two different levels of accuracy. The theoretical results are in good agreement with the experiments, predicting significant mixing of Rydberg and π^* states, and allowed for novel assignments in the photon energy covered in this work which have not been previously reported in the literature. The state at 6.401 eV is tentatively assigned for the first time to the $\sigma^*(OH) \leftarrow 3\pi(3a'')$ transition. The analysis of the observed vibronic structure in the photoabsorption spectra is generally consistent with earlier data, although the higher resolution of the present experiments has enabled us to propose for the first time assignments for the valence and Rydberg excitations. The photolysis lifetimes of phenol have also been carefully derived for the Earth's troposphere and stratosphere and show that solar photolysis is expected to be a weak sink in the terrestrial atmosphere.

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Figure captions

FIG. 1. Calculated structure of phenol, C_6H_5OH , optimized at (a) LC- ω PBE/cc-pVTZ geometry and (b) CCSD(T)/cc-pVTZ geometry. Distances in Å.

FIG. 2. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 3.9–10.8 eV photon energy range.

FIG. 3. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 4.3–5.1 eV photon energy range. For the vibrational excitation assignments see TABLE III and text for details.

FIG. 4. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 5.3 – 6.3 eV photon energy range. For the vibrational excitation and Rydberg state assignments see TABLE IV, TABLE VI and text for details.

FIG. 5. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 6.2 – 7.6 eV photon energy range. For the vibrational excitation and Rydberg state assignments see TABLE V, TABLE VI and text for details.

FIG. 6. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 7.3 – 8.7 eV photon energy range. For the vibrational excitation and Rydberg state assignments see TABLE V, TABLE VI and text for details.

FIG. 7. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 8.6 – 10.8 eV photon energy range with Rydberg and vibrational excitation assignments. See TABLE V, TABLE VI and text for details.

Table captions

TABLE I. Calculated vertical excitation energies (TD-DFT level at the LC- ω PBE/aug-ccpVTZ+R) (eV), optical oscillator strengths (f_L), assignments and dominant configurations of phenol (C₆H₅OH) compared with experimental data and other work (details in text)

TABLE II. Calculated vertical excitation energies (EOM-CCSD level at the aug-cc-pVTZ+R basis set) (eV), optical oscillator strengths (f_L), assignments and dominant configurations (C₆H₅OH) compared with experimental data and other work (details in text)

TABLE III. Proposed vibrational assignments in the 4.3-5.1 eV absorption band of phenol, C₆H₅OH

TABLE IV. Proposed vibrational assignments in the 5.3–6.3 eV absorption band of phenol, C_6H_5OH

TABLE V. Proposed vibrational assignments in the 6.2-10.8 eV absorption region of phenol, C_5H_5OH

Table VI. Energy value (eV), quantum defect (δ) and assignment of the Rydberg series converging to the ionic electronic ground (4a^{"-1}), first (3a^{"-1}) and second (2a^{"-1}) excited states of phenol, C₆H₅OH

FIG. 1. Calculated structure of phenol, C_6H_5OH , optimized at (a) LC- ω PBE/cc-pVTZ geometry and (b) CCSD(T)/cc-pVTZ geometry. Distances in Å.



(a) LC-ωPBE/cc-pVTZ geometry



(b) CCSD(T)/cc-pVTZ geometry

FIG. 2. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 3.9–10.8 eV photon energy range.



FIG. 3. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 4.3–5.1 eV photon energy range. For the vibrational excitation assignments see TABLE III and text for details.



FIG. 4. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 5.3 – 6.3 eV photon energy range. For the vibrational excitation and Rydberg state assignments see TABLE IV, TABLE VI and text for details.



FIG. 5. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 6.2 – 7.6 eV photon energy range. For the vibrational excitation and Rydberg state assignments see TABLE V, TABLE VI and text for details.



FIG. 6. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 7.3 – 8.7 eV photon energy range. For the vibrational excitation and Rydberg state assignments see TABLE V, TABLE VI and text for details.



FIG. 7. High resolution VUV photoabsorption spectrum of phenol, C_6H_5OH , in the 8.6 – 10.8 eV photon energy range with Rydberg and vibrational excitation assignments. See TABLE V, TABLE VI and text for details.



TABLE I. Calculated vertical excitation energies (TD-DFT level at the LC- ω PBE/aug-cc-pVTZ+R) (eV), optical oscillator strengths (f_L), assignments and dominant configurations of phenol (C₆H₅OH) compared with experimental data and other work (details in text)

<u>Ctata</u>	$\mathbf{E}(\mathbf{J})$	£		E (-V) b	HOMO		Minud Character		Cross-sections
State	E (ev)	IL	E (ev) -	$E(ev)^{-1}$	номо	HOMO-1	Mixed Character	Exp. (ev)	(Mb)
					4π (4a")	3π (3a")			
\tilde{X} ¹ A'	_	_	_	_			_		
2 ¹ A'	5.310	0.0336	4.53	4.99			$\mathrm{HOMO} \rightarrow \mathrm{LUMO} + \mathrm{HOMO-1} \rightarrow \mathrm{LUMO+1}$	4.605	18.1
3 ¹ A'	6.122	0.0374	5.80	5.76			$\mathrm{HOMO-1} \rightarrow \mathrm{LUMO} + \mathrm{HOMO} \rightarrow \mathrm{LUMO+1}$	6.00(4)	19.1
1 ¹ A"	6.231	0.0001	5.76	5.13	3sσ/σ*(OH)			5.75(5)	
2 ¹ A"	6.744	0.0038	6.20	5.57	3pσ/σ*(OH)			6.00(4)	
3 ¹ A"	7.031	0.0006	6.41	5.92	3pσ/σ*(OH)				
4 ¹ A'	7.073	0.0576	6.31	7.01	3рπ				
5 ¹ A'	7.120	0.6075	6.50	6.71			$\mathrm{HOMO-1} \rightarrow \mathrm{LUMO} + \mathrm{HOMO} \rightarrow \mathrm{LUMO+1}$	6.736	192.5
6 ¹ A'	7.143	0.4208	6.56	6.66			$\mathrm{HOMO-1} \rightarrow \mathrm{LUMO+1+} \mathrm{HOMO} \rightarrow \mathrm{LUMO}$	6.736	192.5
4 ¹ A"	7.288	0.0010	6.42	5.98		3sσ/σ*(OH)			
5 ¹ A"	7.362	0.0021	6.75	6.31	3d o				
6 ¹ A"	7.435	0.0142	6.95	6.54	3d o			6.91(1)	

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7 ¹ A"	7.617	0.0010	7.10	6.93	3d o			
7 ¹ A'	7.698	0.0110	6.92		3dπ			
8 ¹ A"	7.722	0.0116	6.88	6.35			$\mathrm{HOMO-1} \to 3p\sigma + \mathrm{HOMO} \to 4s\sigma$	
8 ¹ A'	7.739	0.0061	6.95		3dπ			
9 ¹ A"	7.797	0.0062					$HOMO \rightarrow 4s\sigma + HOMO\text{-}1 \rightarrow 3p\sigma$	
10 ¹ A"	7.934	0.0031	7.07	6.66			$\mathrm{HOMO-1} \to 3\mathrm{p\sigma} + \mathrm{HOMO} \to 4\mathrm{p\sigma}$	
11 ¹ A"	7.952	0.0029					$\mathrm{HOMO-1} \rightarrow 3\mathrm{p\sigma} + \mathrm{HOMO} \rightarrow 4\mathrm{p\sigma}$	
9 ¹ A'	8.020	0.0010			4pπ			
12 ¹ A"	8.038	0.0001			4po			
10 ¹ A'	8.042	0.0064	6.85			Зрл		6.736
13 ¹ A"	8.081	0.0051		7.57			$\sigma_{ring}(CC) \rightarrow LUMO$	
14 ¹ A"	8.178	0.0012	7.84		4po		7.	.39(5)
15 ¹ A"	8.204	0.0014	7.57			Зрσ		
16 ¹ A"	8.231	0.0012			4d o			7.690
17 ¹ A"	8.305	0.0009			4d o			
18 ¹ A"	8.407	0.0009	7.62				$HOMO\text{-}1 \rightarrow 3d\sigma + HOMO \rightarrow 4d\sigma$	
19 ¹ A"	8.417	0.0021					$HOMO \rightarrow 3d\sigma + HOMO-1 \rightarrow 3d\sigma$	
11 ¹ A'	8.419	0.0009			Зрπ			

4do

^a Ref. 16 (CASPT2/ANO+R); ^b Ref. 26 (TD-DFT B3LYP/aug-cc-pVDZ); ^c the last decimal of the energy value is given in brackets for these less-resolved features

TABLE II. Calculated vertical excitation energies (EOM-CCSD level at the aug-cc-pVTZ+R basis set) (eV), optical oscillator strengths (f_L), assignments and dominant configurations (C₆H₅OH) compared with experimental data and other work (details in text)

State	E (eV)	f_L	$< r^2 > a$	E (eV) ^b	E (eV) °	НОМО	HOMO-1	НОМО-2	Mixed Character	Exp. (eV) ^d	Cross-sections
											(Mb)
						4π (4a")	3π (3a")	2π (2a")			
$ ilde{X}$ $^{1}\mathrm{A'}$	_	_	88						_		
2 ¹ A'	4.947	0.0210	91	4.53	4.99	LUMO (5a", 1π*)				4.605	18.1
1 ¹ A"	5.699	0.0002	108	5.76	5.13	3sσ/σ*(OH)				5.75(5)	
^d . ∑ 3 ¹ A′	6.162	0.0337	99	5.80	5.76	LUMO+1 (6a", 2 π *)				6.00(4)	19.1
⁰ ⊕ 2 ¹ A"	6.184	0.0043	168	6.20	5.57	3pσ/σ*(OH)				6.00(4)	
^{SO} 3 ¹ A"	6.491	0.0004	160	6.41	5.92	Зро					
⊑0_1 4 ¹A"	6.634	0.0011	138	6.42	5.98		$3s\sigma/\sigma^*(OH)$			5.89(3)/6.401	
00 4 ¹ A'	6.642	0.0183	161	6.31	7.01	Зрл					
⁰ 5 ¹ A"	6.876	0.0053	181	6.75	6.31	3d o					
[○]	6.906	0.0089	178	6.95	6.54	3d o				6.91(1)	
	7.011	0.3889	113	6.50	6.66				$\mathrm{HOMO} \rightarrow \mathrm{LUMO} + \mathrm{HOMO-1} \rightarrow \mathrm{LUMO+1}$	6.736	192.5
¹ A"	7.079	0.0117	153	6.88	6.35		Зрσ				
≤ ► ¹ A′	7.101	0.5172	123	6.56	6.71				HOMO-1 \rightarrow LUMO + HOMO \rightarrow 3d π	6.736	192.5
SO											
Σ											37
$\langle $											

8	$^{1}A''$	7.113	0.0001	194	7.10	6.93	3d o		
9	¹ A"	7.276	0.0130	164	7.07	6.66		3ро	
7	$^{1}A'$	7.280	0.0840	202	6.92		3dπ		
10	¹ A"	7.325	< 0.0001	298			4sσ		7.12(8)
8	¹ A′	7.337	0.1376	192	6.95		3dπ		
11	¹ A"	7.494	0.0012	425			4pσ		7.39(5)
9	¹ A′	7.555	0.0603	178	6.85			Зря	6.736
12	$^{1}A''$	7.581	0.0002	436			4pσ		
10	$^{1}A'$	7.592	0.0069	434			4рπ		
d. /28	¹ A"	7.654	0.0007	163	7.57	7.08		3do	
t≷ie	¹ A"	7.732	0.0002	272	7.62				$\mathrm{HOMO-1} \rightarrow 3\mathrm{d}\sigma + \mathrm{HOMO} \rightarrow 4\mathrm{d}\sigma$
SOUL 15	¹ A"	7.758	0.0004	261	7.84				$HOMO \rightarrow 4d\sigma + HOMO-1 \rightarrow 3d\sigma$
uouj 16	¹ A"	7.798	0.0016	426			4d o		7.690
17 ogge	¹ A"	7.836	0.0016	312			4do e		7.690
0 18	¹ A"	7.879	0.0002	675			5sσ		7.802
Ď	¹ A"	7.903	0.0002	362			4d o		
DZ	¹ A"	7.972	0.0001	243				3do	
ΥIE	¹ A"	7.989	0.0022	818			5ро		7.885
	$^{1}A'$	7.992	0.0027	974			4dπ		
105									
2									

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22 ¹ A"	8.007	0.0009	1013		5ро				
23 ¹ A"	8.023	0.0018	443		5d o				
12 ¹ A'	8.043	0.0040	210			3dπ			
13 ¹ A'	8.069	0.0093	369			3dπ			7.562
14 ¹ A'	8.120	0.0024	304		5dπ				
24 ¹ A"	8.124	0.0059	865		5d o				8.040
15 ¹ A'	8.146	0.0046	206		5dπ				
25 ¹ A"	8.165	0.0046	1046		5d o				
26 ¹ A"	8.170	0.0002	355			4sσ			7.803
^d . ∑27 ¹ A"	8.228	0.0052	104	7.57				$\sigma_{ring} \rightarrow LUMO$	
00 28 ¹ A"	8.246	0.0010	208		5d o				
29 ¹ A"	8.281	0.0005	1005		5ро				
16 ¹ A'	8.344	0.0007	978		5рπ				
⁰⁰ 30 ¹ A"	8.347	0.0018	433			4ро			
0 17 ¹ A'	8.420	0.0007	442			4pπ			8.072
\mathbf{A}'	8.712	0.0745	167				LUMO (5a",1π)		
X) ¹ A'	8.829	0.0036	872			$4d\pi$			8.400
) ¹ A'	8.858	0.0003	372			$4d\pi$			
N Ion	8.598				RCCSD(T)				

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^a Mean value of r^2 (electronic radial spatial extents); ^b Ref. 16 (CASPT2/ANO+R); ^c Ref. 26 (TD-DFT B3LYP/aug-cc-pVDZ); ^d the last decimal of the energy value is given in brackets for these less-resolved features; ^e this MO has a f-shape character as in the $\sigma^*(CC)$ MO of cyclopropane, see discussion in Refs. 53 and 54.

TABLE III. Proposed vibrational assignments in the 4.3-5.1 eV absorption band of phenol, C₆H₅OH

Energy (eV)	Assignment	Energy (eV)	Assignment
This	s work [*]	Nix et a	l. ²²
4.355	$7a_1^0$	_	_
4.380	18 ⁰ ₁	_	_
4.383	$6a_2^0$	_	_
4.406	12 ⁰	_	_
4.416	$18b_{2}^{0}$	_	_
4.440	$6b_1^0$	_	_
4.443	$6a_1^0$	4.442	$6a_1^0$
4.458	$18b_{1}^{0}$	_	_
4.501	$16b_1^1$ a	_	_
4.508	000	4.507	000
_	_	4.512	$4^0_15^1_0$
4.513	?	4.513	$4^1_0\tau 0 H^0_1$
4.548	$ au OH_1^1$	4.548	$\tau 0 H_1^1$
_	_	4.554	$16a_0^2$
4.555	$18b_0^1$	4.556	$18b_{0}^{1}$
4.567	$6a_0^1$	4.567	$6a_0^1$
4.577	$6b_0^1$	_	_
4.60(1)(s)	$18b_0^2$	_	_
4.605	12 ¹ ₀	_	_
4.624	$6a_0^2$	_	_
4.626	$18a_0^1$	4.626	$18a_{0}^{1}$
4.636	$9a_0^1$	4.629	$9a_0^1$
4.649	$6b_0^2$	-	_
4.65(4)(w)	$18b_0^3$	_	_
4.666	$7a_0^1$	_	_
4.683	$6a_0^3/6a_0^112_0^2$	_	-
4.702	12_0^2	_	-
4.713	$6b_0^3$	_	-
4.721	$6a_0^17a_0^1$	_	_
4.741	$6a_0^2 12_0^2$		
4.763	$9a_0^2$	_	_
4.781	$1_0^17a_0^1/6a_0^27a_0^1$	4.781	$1_0^17a_0^1$
4.798	$12_0^3/6a_0^312_0^2$	_	_
4.818	$1_0^1 12_0^2$	4.817	$1^1_0 12^2_0$
4.822	$7a_0^2$	_	_
4.838	$6a_0^37a_0^1$	_	_
4.877	$6a_0^17a_0^2$	_	_
4.896	124	_	-

(s) shoulder feature. (w) weak feature. (b) broad structure (the last decimal of the energy value is given in brackets for these less-resolved features). ? feature remains unassigned.

* Vibrational modes notation adopted from Refs. 19 and 22; ^a From Ref. 45.

TABLE IV. Proposed vibrational assignments in the 5.3–6.3 eV absorption band of phenol, C_6H_5OH

Energy (eV)	Assignment	$\Delta E (eV)$
5.75(5)(b)	000	
5.85(9)(b)	9^{1}_{0}	0.104
5.89(3)(b)	_	_
6.00(4)(b)	9^{1}_{0}	0.111
6.10(2)(b,w)	9 ₀ ²	0.098

(b) broad structure; (w) weak feature (the last decimal of the energy value is given in brackets for these less-resolved features)

Energy (eV)) Assignment	$\Delta E (eV)$	Energy (eV)	Assignment	$\Delta E (eV)$
6.461	000	_	7.690	4d	_
6.52(2)(s)	$6a_0^1$	0.061	7.741	$4d + 6a_0^1$	0.051
	_	_	7.799	$4d + 6a_0^2$	0.058
6.634	$6a_0^3$	_			
6.68(4)(b)	$6a_0^4$	0.050	7.803	5s / 4s	_
6.735	$6a_0^5$	0.051	7.858	$5s / 4s + 6a_0^1$	0.055
6.790	$6a_0^6$	0.055			
6.84(8)(s)	$6a_0^7$	0.058	7.885	5p	_
			7.951	$5p + 6a_0^1$	0.066
6.91(1)(s)	3d	_	8.001	$5p + 6a_0^2$	0.050
6.96(2)(s,w)	$3d + 6a_0^1$	0.051	8.05(6)(w)	$5p + 6a_0^3$	0.055
7.01(7)(s,w)	$3d + 6a_0^2$	0.055			
			7.915(w)	_	_
7.39(5)(s)	4p	_	7.962	$6a_0^1$	0.047
7.44(4)(s)	$4p + 6a_0^1$	0.049	8.007	$6a_0^2$	0.045
7.496	$4p + 6a_0^2$	0.052			
7.562	$4p + 6a_0^3 / 3d$	0.066	8.093	6s	_
7.622	$3d + 6a_0^1$	0.060	8.15(2)(s)	$6p / 6s + 6a_0^1$	0.059
7.675	$3d + 6a_0^2$	0.053	8.197	$6p / 6s + 6a_0^2$	0.045
7.735	$3d + 6a_0^3$	0.060			

TABLE V. Proposed vibrational assignments in the 6.2-10.8 eV absorption region of phenol, C_5H_5OH

			8.429	9s	_
7.60(6)(s)	_	_	8.50(1)(b)	$9s + 6a_0^1$	0.072
7.66(0)(w)	$6a_{0}^{1}$	0.054	8.565	$9s + 6a_0^2 / 5p$	0.064
7.711	$6a_0^2$	0.051			
7.76(9)(s)	$6a_0^3$	0.058	8.472	5s	_
7.823(w)	$6a_0^4$	0.054	8.52(1)(s)	$5s + 6a_0^1$	0.049
7.880	$6a_0^5$	0.057			
			8.84(3)(b)	бр	_
			9.00(4)(b)	$6p + 8a_0^1 / 7p / 3p$	0.161
			9.17(0)(b)	$6p + 8a_0^2 / 7p + 8a_0^1 / 3p$	0.166
				$+ 8a_0^1$	

(s) shoulder structure. (b) broad structure. (w) weak structure (the last decimal of the energy value is given in brackets for these less-resolved features).

Table VI. Energy value (eV), quantum defect (δ) and assignment of the Rydberg series converging to the ionic electronic ground (4a"⁻¹), first (3a"⁻¹) and second (2a"⁻¹) excited states of phenol, C₆H₅OH

En	δ	Assignment	En	δ	Assignment
$IE_1 = 8.64 \text{ eV}$			$IE_2 = 9.32 \text{ eV}$		
5.75(5) (b)	0.83	3s	5.89(3) (s)	1.01	3s
7.12(8) (b,w)	1.00	4s	7.803	1.00	4s
7.803	0.97	5s	8.472	0.99	5s
8.093	1.01	6s			
8.263	0.99	7s	6.736	0.70	3p
8.363	0.99	8s	8.072	0.70	4p
8.429	0.97	9s	8.565	0.75	5p
			8.84(3) (b)	0.66	6р
6.00(4)	0.73	3p	9.00(4) (b)	0.44	7p
7.39(5) (s)	0.69	4p			
7.885	0.75	5p	7.562	0.22	3d
8.15(2) (s)	0.72	6р	8.400	0.15	4d
8.299	0.68	7p	8.74(7) (s,w)	0.13	5d
6.91(1) (s)	0.19	3d	$\overline{IE_3 = 11.57 \text{ eV}}$	_	
7.690	0.21	4d	8.178	1.00	3s
8.040	0.24	5d			
8.233	0.22	6d	9.00(4) (b)	0.70	3p
8.346	0.20	7d			

9.70(9) (b) 0.30 3d

(b) broad structure. (w) weak structure. (s) shoulder structure (the last decimal of the energy value is given in brackets for these less-resolved features).