

The **25th** International Conference on

# High Resolution Molecular Spectroscopy



**Bilbao** 2018 September 3rd–7th

Bizkaia Aretoa – UPV/EHU



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25th International Conference on High Resolution Molecular Spectroscopy – Abstract Book

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Varios Autores

Bilbao 2018

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# Scientific program

Monday, September 3, 2018

## Morning Sessions

8:00–9:45	Registration
Main Hall	
9:45–10:30	Opening Ceremony
Mitxelena Auditorium	<p><b>With the presence of institutions endorsing the conference:</b></p> <p><b>Prof. Adolfo Morais Ezquerro</b> Vice Counselor for Research and Universities of the Basque Government</p> <p><b>Prof. José Luis Martín González</b> Vice Rector for Research of the University of the Basque Country</p> <p><b>Prof. Fernando Plazaola Muguruza</b> Dean of the Faculty of Science and Technology of the University of the Basque Country</p> <p><b>Prof. Jens-Uwe Grabow</b> INTERNATIONAL STEERING COMMITTEE Chairman Institut für Physikalische Chemie und Elektrochemie Gottfried Wilhelm Leibniz Universität (LUH)</p> <p><b>Prof. Alberto Lesarri</b> EXECUTIVE COMMITTEE Department of Physical Chemistry and Inorganic Chemistry Faculty of Sciences University of Valladolid (UVa)</p> <p><b>Dr. Emilio J. Cocinero</b> LOCAL ORGANIZING COMMITTEE Department of Physical Chemistry Faculty of Science and Technology – University of the Basque Country (UPV / EHU) and Biofisika Institute (CSIC-UPV / EHU)</p>
10:30–11:10	Coffee Break
Main Hall	<b>Sponsors Stands will be placed at Main Hall</b>
	Plenary Session
	Chair: <b>Stepan Urban</b> , <i>University of Chemistry &amp; Technology Prague</i>
11:10–11:55	Plenary Lecture 1
Mitxelena Auditorium	PT1. Collision-induced spectra of N <sub>2</sub> and O <sub>2</sub> <b>Ad Van Der Avoird</b> , <i>Radboud University</i>
11:55–12:40	Plenary Lecture 2
Mitxelena Auditorium	PT2. Spectroscopy of molecular radicals in Helium droplets <b>Gary E. Douberly</b> , <i>University of Georgia</i>
13:00–14:30	Lunch Break
	<b>Iberdola tower (black tickets) or Deusto Library (white tickets)</b>

## Afternoon Sessions

Oral Sessions 1			
14:30–15:50	Oral Session A1:	Oral Session B1:	Oral Session C1:
	<b>Mixelena Auditorium</b>	<b>Baroja Auditorium</b>	<b>Oteiza Auditorium</b>
	Chair: <b>Vadym Ilyushyn</b> , <i>Institute of Radio Astronomy of NASU</i>	Chair: <b>Sonia Melandri</b> , <i>University of Bologna</i>	Chair: <b>Michel Mons</b> , <i>CEA Saclay</i>
	LARGE-AMPLITUDE MOTIONS	WATER CLUSTERS	FREQUENCY COMB AND FAST IR
<b>14:30–14:50</b>	A1.1. Microwave study of internal rotation in para Tolu-aldehyde: local versus global symmetry at the methyl-rotor site as an indicator of information transfer across the benzene ring	B1.1. Structures and dynamics of acrolein-(H <sub>2</sub> O) <sub>n</sub> clusters revealed by MW spectroscopy and ab initio calculation	C1.1. Fast scanning IR-spectrometer to measure transient molecules in a pulsed supersonic jet
	<b>Jon Hougen</b> , <i>NIST</i>	<b>Weixing Li</b> , <i>University of Bologna</i>	<b>Daniel Witsch</b> , <i>University of Kassel</i>
<b>14:50–15:10</b>	A1.2. Effective rotational hamiltonian for two-rotor systems with symmetric and asymmetric internal rotors (like Ethanol) applied to Ethylphosphine, CH <sub>3</sub> CH <sub>2</sub> PH <sub>2</sub>	B1.2. Discovering the conformations of formamide complexes and formamide-water clusters by microwave spectroscopy	C1.2. Wide bandwidth mid-IR spectroscopy with comb-referenced EC-QCL: application to the ν <sub>1</sub> fundamental band of <sup>14</sup> N <sub>2</sub> <sup>16</sup> O
	<b>Peter Groner</b> , <i>University of Missouri – Kansas City</i>	<b>Susana Blanco</b> , <i>Universidad de Valladolid</i>	<b>Davide Gatti</b> , <i>IFN/CNR &amp; Politecnico di Milano</i>
<b>15:10–15:30</b>	A1.3. Proton in a double-well potential: Acetylacetone and its derivatives by microwave spectroscopy	B1.3. Does the structure of the polycyclic aromatic hydrocarbon impact the aggregation of water on its surface? fluorene vs ace-naphthene	C1.3. Endless frequency-swept comb-calibrated spectrometer
	<b>Luca Evangelisti</b> , <i>University of Bologna</i>	<b>Amanda Steber</b> , <i>The Hamburg Centre for Ultrafast Imaging</i>	<b>Thomas Puppe</b> , <i>TOPTICA Photonics AG</i>
<b>15:30–15:50</b>	A1.4. Sensing the molecular structures of alkyl methyl ketones by internal rotation in the microwave spectrum	B1.4. Characterization of microsolvated 15C5 crown ether from broadband rotational spectroscopy	C1.4. Broadband complex gas spectroscopy with sub-kHz level resolution comb spectrometer
	<b>Maike Andresen</b> , <i>IPC RWTH Aachen University</i>	<b>Juan Carlos López</b> , <i>Universidad de Valladolid</i>	<b>Grzegorz Kowzan</b> , <i>Nicolaus Copernicus University in Torun</i>

Oral Session 2			
16:00–18:00	Oral Session A2:	Oral Session B2:	Oral Session C2:
	<b>Mitxelena Auditorium</b>	<b>Baroja Auditorium</b>	<b>Oteiza Auditorium</b>
	Chair: <b>Isabelle Kleiner</b> , <i>LISA / CNRS</i>	Chair: <b>Wen-Bih Tzeng</b> , <i>Academia Sinica</i>	Chair: <b>Donald McNaughton</b> , <i>Monash University</i>
	PES AND SPECTRAL INTERPRETATION	RADICALS AND IONS	SMALL MOLECULES: IR
16:00–16:20	A2.1. Empirical potential energy surface and bending angle probability densities for the electronic ground state of HCO <sup>+</sup>	B2.1. FTMW spectroscopy of sulfur bearing free radicals, HCSC and CH <sub>3</sub> SS	C2.1. The water vapor absorption continuum in the atmospheric windows at 4.0, 2.1, 1.6 and 1.25 μm
	<b>Per Jensen</b> , <i>University of Wuppertal</i>	<b>Yasuki Endo</b> , <i>National Chiao Tung University</i>	<b>Alain Campargue</b> , <i>CNRS / Université de Grenoble</i>
16:20–16:40	A2.2. Torsion rotation program for nitromethane CH <sub>3</sub> NO <sub>2</sub>	B2.2. Millimeter-wave spectroscopy of the HDCCH radical	C2.2. Sub-MHz deuterium spectroscopy and comparison with ab initio calculations of the line-shape effects
	<b>Marek Kręglewski</b> , <i>Adam Mickiewicz University in Poznań</i>	<b>Keiichi Tanaka</b> , <i>Kyushu University</i>	<b>Mikolaj Zaborowski</b> , <i>Nicolaus Copernicus University in Toruń</i>
16:40–17:00	A2.3. Weak intramolecular interaction effects on the structure and torsional spectra of ethylene glycol isotopologues, an astrophysical species	B2.3. Accurate rotational fingerprints of radioactive radicals by mass-independent studies on AlO, TiO, and FeO	C2.3. The high-resolution spectrum of DC <sub>3</sub> N recorded in the infrared and millimeter-wave regions: a global analysis
	<b>María Luisa Senent</b> , <i>CSIC</i>	<b>Alexander Breier</b> , <i>Universität Kassel</i>	<b>Filippo Tamassia</b> , <i>University of Bologna</i>
17:00–17:20	A2.4. Modelling temperature dependent anharmonic spectra of pyrene (C <sub>16</sub> H <sub>10</sub> ): comparison of computational approaches	B2.4. Accurate sub-millimeter rest-frequencies for HCCO and DCCO radicals	C2.4. Sub-doppler metrology of HD
	<b>Shubhadip Chakraborty</b> , <i>Institut de Recherche en Astrophysique et Planetologie</i>	<b>Johanna Chantzou</b> , <i>Max Planck Institut for extraterrestrial Physics</i>	<b>Patrick Dupré</b> , <i>Laboratoire de Physico-Chimie de l'Atmosphère</i>
17:20–17:40	A2.5. Non-adiabatic coupling in the ozone molecule	B2.5. Terahertz spectroscopy of isotopologues of amidogen radical	C2.5. IR laser spectroscopy of the deuterated isotopologues of ammonia
	<b>Alexander Alijah</b> , <i>University of Reims</i>	<b>Mattia Melosso</b> , <i>Università di Bologna</i>	<b>Patrice Cacciani</b> , <i>Université Lille / CNRS</i>
17:40–18:00	A2.6. The long-range behavior of ab initio transition dipole moments and spin-orbit coupling matrix elements between the low-lying electronic states of alkali heterodimers	B2.6. Accurate rotational frequencies of deuterated Ammonium ions (d <sub>1</sub> -d <sub>3</sub> ) measured in a cryogenic ion trap.	C2.6. Photoacoustic spectroscopy of the oxygen a-band in support of OCO-2
	<b>Ekaterina Bormotova</b> , <i>Moscow State University</i>	<b>José Luis Doménech</b> , <i>Instituto de Estructura de la Materia CSIC</i>	<b>Elizabeth Lunny</b> , <i>California Institute of Technology</i>
18:00–20:00 Menchu Gal Terrace	<b>Welcome Cocktail</b> With representatives of the City Council of Bilbao		

Tuesday, September 4, 2018

Morning Sessions

Plenary Session			
Chair: <b>Marek Kręglewski</b> , Adam Mickiewicz University in Poznań			
9:00–9:45 Plenary Lecture 3			
Mitxelena Auditorium	PT3. Beryllium bonding probed by anion photodetachment spectroscopy		
Michael Heaven, Emory University			
9:45–10:30 Plenary Lecture 4			
Mitxelena Auditorium	PT4. Which spectroscopy to study astro-PAHs?		
Christine Joblin, IRAP – Université de Toulouse			
10:30–11:00 Coffee Break			
Main Hall	Sponsors Stands will be placed at Main Hall		
Oral Sessions 3			
11:00–12:40	Oral Session A3:	Oral Session B3:	Oral Session C3:
	Mitxelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Per Jensen</b> , University of Wuppertal	Chair: <b>José A. Fernández</b> , Universidad País Vasco UPV/EHU	Chair: <b>Robert Gamache</b> , University of Massachusetts Lowell
	LARGE-AMPLITUDE MOTIONS	LIF AND REMPI	INTERMEDIATES, CHIRALITY AND BIOMOLECULES
11:00–11:20	A3.1. Spin-torsion dominated hyperfine splittings in the first excited torsional state ( $v_t = 1$ ) of Methanol	B3.1. High resolution Rydberg spectroscopy of 3d metal sandwich compounds: ultrasensitive probing the electron density changes in organometallic molecules	C3.1. Local interactions of the asparagine and histidine aminoacids in a protein chain: gas phase conformer-selective IR spectroscopy of model molecules
	<b>Li-Hong Xu</b> , University of New Brunswick	<b>Sergey Ketkov</b> , G.A. Razuvaev Institute of Organometallic Chemistry RAS	<b>Michel Mons</b> , CEA Saclay
11:20–11:40	A3.2. Extension of the normal mode approach to nonrigid polyatomic molecules	B3.2. Two-color REMPI $[(1+1)+1]$ of the singlet oxygen $^1O_2$ arising in UV-photodissociation of van der Waals complex $C_5H_8-O_2$	C3.2. Water sculpts the distinctive shapes and dynamics of the Tn antigens: implications for their molecular recognition
	<b>Dominika Vigiłaska</b> , University of Reims	<b>Alexandr Bogomolov</b> , Voevodsky Institute of Chemical Kinetics and Combustion	<b>Aran Insausti</b> , Universidad del País Vasco (UPV/EHU)
11:40–12:00	A3.3. Essentially free internal rotation of the Propynyl Methyl group investigated by microwave spectroscopy	B3.3. The vibrational structure of the $NO_3 X^2A_2'$ state studied by SVL DF spectrum	C3.3. Testing the parity symmetry in cold chiral molecules using vibrational spectroscopy
	<b>Konrad Eibl</b> , IPC RWTH Aachen University	<b>Masaru Fukushima</b> , Hiroshima City University	<b>Anne Cournot</b> , Université Sorbonne Paris Cité

12:00–12:20	A3.4. Microwave spectroscopic and quantum chemical studies of the coupled large amplitude motions in S-Phenyl Thioacetate	B3.4. Line shape study of oxygen B-band with CRDS in Dual-Beam configuration	C3.4. Hydrated Lewis antigens in the gas phase: building up the environment or peeling off the solvent?
	<b>Lynn Ferres,</b> <i>IPC RWTH Aachen University</i>	<b>Katarzyna Bielska,</b> <i>Nicolaus Copernicus University in Torun</i>	<b>Pierre Carçabal,</b> <i>Institut des Sciences Moléculaires d'Orsay CNRS</i>
12:20–12:40	A3.5. Separately fitting the torsional symmetry species of molecules with one or multiple internal rotor(s)	B3.5. LIF Spectrum of a $^1\Sigma$ species containing Si: linear SiOSi?	C3.5. High-resolution ro-vibrational spectrum of the simplest Criegee intermediate CH <sub>2</sub> OO between 880 and 932 cm <sup>-1</sup>
	<b>Lam Nguyen,</b> <i>Laboratoire LISA</i>	<b>Masaru Fukushima,</b> <i>Hiroshima City University</i>	<b>Pei-Ling Luo,</b> <i>Institute of Atomic and Molecular Sciences Academia Sinica</i>
13:00–14:30	<b>Lunch Break</b>		
	<b>Iberdrola tower (black tickets) or Deusto Library (white tickets)</b>		

## Afternoon Sessions

Oral Session 4			
14:30–15:50	Oral Session A4:	Oral Session B4:	Oral Session C4:
	Mitxelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Filippo Tamassia,</b> <i>University of Bologna</i>	Chair: <b>Leonid Surin,</b> <i>Institute of Spectroscopy RAS</i>	Chair: <b>A. Robert McKellar,</b> <i>National Research Council of Canada</i>
	SMALL MOLECULES: IR	LARGE MOLECULES: MW	LINE SHAPES AND INTENSITIES
14:30–14:50	A4.1. Investigation of the $\nu_8$ and $\nu_{21}$ bands of propane CH <sub>3</sub> CH <sub>2</sub> CH <sub>3</sub> at 11.5 and 10.9 $\mu\text{m}$ : evidence of large amplitude tunnelling effects	B4.1. On not yet solved problems concerning some molecular systems investigated by rotational spectroscopy along my research life	C4.1. On the temperature dependence of half-widths and line shifts for molecular transitions in the microwave and infrared regions
	<b>Agnes Perrin,</b> <i>Laboratoire de Meteorologie Dynamique</i>	<b>Walther Caminati,</b> <i>University of Bologna</i>	<b>Robert Gamache,</b> <i>University of Massachusetts Lowell</i>
14:50–15:10	A4.2. High resolution FTIR study of the $\nu_6$ band of CH <sub>2</sub> F <sub>2</sub> in 3 $\mu\text{m}$ spectral region at low temperatures	B4.2. Rotational spectrum and molecular structure of succinic anhydride aided by computational calculations	C4.2. Calculation of half widths and line shifts of H <sub>2</sub> O–N <sub>2</sub> collision system using the modified complex Robert–Bonamy formalism
	<b>Chilukoti Ashok,</b> <i>Homi Bhabha National Institute</i>	<b>Donald McNaughton,</b> <i>Monash University</i>	<b>Bastien Vispoel,</b> <i>University of Massachusetts Lowell</i>
15:10–15:30	A4.3. The NH <sub>2</sub> scissors band of Methylamine	B4.3. Pursuing the rotational spectra of large molecular systems	C4.3. High accuracy ab initio DMS and extra high accuracy PES for sub-percent calculation of line intensities
	<b>Iwona Gulaczyk,</b> <i>Adam Mickiewicz University in Poznań</i>	<b>Iker León,</b> <i>Universidad de Valladolid</i>	<b>Oleg Polyansky,</b> <i>UCL</i>

15:30–15:50	A4.4. Wrong distance – wrong line strength – wrong concentrations	B4.4. Exploring the generation of new species using laser ablation Fourier transform microwave spectroscopy techniques: the study of N-carbamoyl glycine	C4.4. Line shape parameters from first principles: investigation of the S and O branches in the H <sub>2</sub> –He system
	<b>Tom Moses Rubin</b> , <i>PTB</i>	<b>Lucie Kolesniková</b> , <i>Universidad de Valladolid</i>	<b>Hubert Józwiak</b> , <i>Nicolaus Copernicus University</i>
16:00–18:30 Axular Room and Chillida Room	<b>Poster Session 1 (P1.1. to P1.64.) and Coffee Break</b>		
18:30–20:30 Meeting point: Main Hall of Bizkaia Aretoa	<b>City Tour</b>		

## Poster presentations – Session 1

### **P1.1. DFT and TD DFT modeling of vibrational structures in high resolution MATI and REMPI spectra of chromium bisarene complexes**

Sergey Ketkov; Sheng-Yuan Tzeng; Elena Rychagova; Grigory Zhigulin; Wen-Bih Tzeng

### **P1.2. Vibronic emission spectroscopy of o-Ethynylbenzyl radical generated from corona discharge of o-Ethynyltoluene**

Sang Kuk Lee

### **P1.3. High resolution study of CH<sub>2</sub>=CD<sub>2</sub> molecule: line intensities and half-widths**

Kirill Berezkin; Elena Bekhtereva; Olga Gromova; Oleg Ulenikov; Claude Leroy; Christian Sydow; Sigurd Bauerecker

### **P1.4. The classification of lung cancers and their degree of malignancy by FTIR, PCA–LDA analysis, and a physics-based computational model**

Joanna Depciuch; Ewa Kaznowska; Kornelia Łach; Magdalena Kołodziej; Anna Koziorowska; Jitraporn Vongsivut; Izabela Zawlik; Marian Cholewa; Jozef Cebulski

### **P1.5. Spectroscopy group at the University of the Basque Country**

Camilla Calabrese, Aran Insausti; Mainer Parra; Cristóbal Pérez; Iciar Uriarte; Imanol Usabiaga; Emilio J. Cocinero; Francisco J. Basterretxea

### **P1.6. Spectroscopic characterization of the unsaturated Hydrocabons C<sub>3</sub>H and C<sub>5</sub>H using highly correlated ab initio methods**

Maria Luisa Senent; Sara Cherifa Bennedjai; Dalila Hammoutenne

### **P1.7. Broadening and shifting coefficients of rovibrational lines of HCl perturbed by He in the fundamental and the first overtone regions**

Ruslan Asfin; Alexandra V. Domanskaya; Gang Li; Volker Ebert

### **P1.8. Predissociation of the B state of S<sub>2</sub>: measurements and modeling**

Glenn Stark; Brenton Lewis; Stephen Gibson; Hannah Herde; Alan Heays; James Lyons; Nelson de Oliveira; Gillian Nave

### **P1.9. Resonant two-photon ionization and mass-analyzed threshold ionization spectroscopy of 2,4-Difluoroanisole**

Wen-Bih Tzeng; Shen-Yuan Tzeng

### **P1.10. High precision spectrum of the second overtone of <sup>12</sup>C<sup>16</sup>O**

Jin Wang; Yu Sun; Anwen Liu; Shuiming Hu



**P1.11. CO<sub>2</sub>-broadening and shift coefficients in the  $\nu_3$  and  $\nu_2+(\nu_4+\nu_5)^0$  bands of acetylene**

Valerii Perevalov; O.M. Lyulin; T.M. Petrova; A.M. Solodov; A.A. Solodov

**P1.12. The ab initio line-shape calculations for purely rotational transitions in the CO-N<sub>2</sub> system**

Hubert Jóźwiak; Hubert Cybulski; Franck Thibault; Nikodem Stolarczyk; Piotr Wcisło

**P1.13. Spectroscopic study of the  $7^1\Pi_u$  and  $7^1\Sigma_u^+$  states of Rb<sub>2</sub> molecule**

Pawel Kowalczyk; Włodzimierz Jastrzebski; Anna Grochola; Katarzyna Olkowska; Jacek Szczepkowski

**P1.14. The <sup>12</sup>CH<sub>4</sub> and <sup>13</sup>CH<sub>4</sub> absorption spectra at 296 K and 200 K in the range between 6600 and 12000 cm<sup>-1</sup>**

Leonid Sinitsa

**P1.15. Study of hydrates of verbenone by microwave Fourier transform spectroscopy and computational chemistry**

Pascal Dréan; Mhamad Chrayteh; Annunziata Savoia; Thérèse R. Huet

**P1.16. Study of the H<sub>2</sub>O-N<sub>2</sub> line broadening and shifting in the region of 16500–17000 cm<sup>-1</sup>**

Leonid Sinitsa; Victor Serdyukov; Nina Lavrentieva; Anna Dudaryonok

**P1.17. Rotational-predissociation double resonance spectroscopy of the He-HCO<sup>+</sup> complex**

Oskar Asvany; Thomas Salomon; Matthias Töpfer; Phillip Schreier; Stephan Schlemmer; Hiroshi Kohguchi; Leonid Surin

**P1.18. Combined PGOPHER analysis of bands in the 1<sup>st</sup> positive system of N<sub>2</sub>, 4500 – 15500 cm<sup>-1</sup>**

Amanda Ross; Patrick Crozet; Jérôme Morville; Colin Western; Dennis Tokaryk

**P1.19. Properties of HF@C<sub>60</sub> endofullerene from first principles**

Yulia Kalugina; Pierre-Nicholas Roy

**P1.20. High sensitivity Cavity Ring Down Spectroscopy spectroscopy of the  $\nu_1+4\nu_3$  band of NO<sub>2</sub> near 1.34  $\mu\text{m}$**

Anastasiia Lukashevskaya; Didier Mondelain; Alain Campargue; Valerii Perevalov

**P1.21. Laser absorption spectroscopy of methane at 1000 K between 1.75 and 1.3  $\mu\text{m}$**

Alain Campargue; Semyon Vasilchenko; Mélanie Ghysels; Didier Mondelain; Samir Kassi; Solenne Barry

**P1.22. Double resonance rotational spectroscopy of CH<sub>3</sub><sup>+</sup> – He**

Matthias Töpfer; Thomas Salomon; Stephan Schlemmer; Oskar Asvany; Otto Dopfer; Hiroshi Kohguchi; Koichi M. T.

**P1.23. The absorption band of nitrogen dioxide (<sup>14</sup>N<sup>16</sup>O<sub>2</sub>) by CRDS near 6000 cm<sup>-1</sup>**

Anastasiia Lukashevskaya; Olga Naumenko; Samir Kassi; Alain Campargue

**P1.24. Isolated small-amplitude fundamental embedded in a pure torsional bath: assignment and fit of the FIR and microwave spectra of the  $\nu_{10}$  vibrational state of acetaldehyde**

Vadym Ilyushyn; Eugene Alekseev; Olga Dorovskaya; Laurent Margulès; Roman Motiyenko; Manuel Goubet; Olivier Pirali; Sigurd Bauerecker; Christof Maul; Christian Sydow; Georg Mellau; Isabelle Kleiner; Jon T. Hougen

**P1.25. The high resolution spectrum of <sup>14</sup>ND<sub>3</sub> in the far-infrared**

Elisabetta Cane; Gianfranco Di Lonardo; Luciano Fusina; Adriana Predoi-Cross; Filippo Tamassia

**P1.26. Line positions and intensities for the  $\nu_3$  band of 5 isotopologues of germane for planetary applications**

Vincent Boudon; Tigran Grigoryan; Florian Philipot; Fridolin Kwabia Tchana; Laurent Manceron; Athéna Rizopoulos; Jean Vander Auwera; Thérèse Encrenaz

**P1.27. The third and fourth torsional states of acetic acid**

Vadym Ilyushyn; Olga Dorovskaya; Eugene Alekseev

**P1.28. First detection of H<sub>2</sub><sup>36</sup>S in the infrared region**

Valeriya A. Zamotaeva; Elena S. Bekhtereva; Olga V. Gromova; Oleg N. Ulenikov; Claude Leroy; Christian Sydow; Sigurd Bauerecker

**P1.29. New electronic states in the spectra of TaH and TaD**

[Thomas Varberg](#); Samuel Gleason; Dalir Kellett; Paul Reischmann

**P1.30. Accurate first principles global calculations and isotopic effects on infrared spectra for phosphine PH<sub>3</sub> and hydrocarbons (CH<sub>4</sub>, C<sub>2</sub>H<sub>4</sub>)**

Vladimir Tyuterev; [Dominika Viglaska](#); Michael Rey; Andrey Nikitin; Eveniya Starikova

**P1.31. Vibrational dependence and prediction of line shape parameters for the H<sub>2</sub>O–H<sub>2</sub> collisional system**

[Robert Gamache](#); Bastien Vispoel

**P1.32. Ab initio reduced matrix elements for H<sub>2</sub><sup>16</sup>O: application to H<sub>2</sub>O–H<sub>2</sub> line shape calculations via the modified complex Robert–Bonamy method**

Bastien Vispoel; Robert Gamache

**P1.33. Classical trajectory simulation of collision-induced absorption spectra**

Daniil N. Chistikov; [Artem A. Finenko](#); Yulia N. Kalugina; Sergei E. Lokshtanov; Sergey V. Petrov; Andrey A. Vigasin

**P1.34. Atlas of experimental and theoretical high-temperature methane cross-sections from T=296 K to 1000 K in the mid-infrared range**

[Vladimir Tyuterev](#); Andy Wong; Peter Bernath; Michael Rey; Andrey Nikitin

**P1.35. High-resolution laser spectroscopy of nitrogen dioxide in the region of 14500–16800 cm<sup>-1</sup>**

[Shunji Kasahara](#); Takumi Yoshizawa; Michihiro Hirata; Kohei Tada

**P1.36. Modified complex Robert–Bonamy calculations including line coupling on the H<sub>2</sub>O–N<sub>2</sub> and CO<sub>2</sub>–N<sub>2</sub> molecular systems**

[Bastien Vispoel](#); Richard Lynch; Robert Gamache

**P1.37. Temperature dependence of line broadening coefficient of chloromethane diluted in carbon dioxide**

[Sylvain Léonis](#); Muriel Lepere

**P1.38. A room-temperature HCN line list with a new dipole moment surface**

[Vladimir Makhnev](#); Aleksandra Kyuberis; Oleg Polyansky; Irina Mizus; Jonathan Tennyson; Nikolay Zobov

**P1.39. Quantum chemical characterisation of difluorine peroxide FOOF**

[Olfa Ferchichi](#)

**P1.40. On the consistency of absolute ozone intensities in MW, 10 and 5 micron ranges from ab initio predictions and laboratory observations**

[Vladimir Tyuterev](#); Alain Barbe; Semen Mikhailenko; Evgeniya Starikova

**P1.41. Rotational and vibrational dependences of line half-widths for CO confined in silica aerogel**

[Alexander A. Solodov](#); Tatyana M. Petrova; Yuriy N. Ponomarev; Alexander M. Solodov

**P1.42. Measurements and calculations of H<sub>2</sub>– broadening and shift parameters of water vapor transitions in 6700–9000 cm<sup>-1</sup> spectral region**

Tatyana M. Petrova; Alexander M. Solodov; [Alexander A. Solodov](#); Vladimir. M. Deichuli; Vitaly. I. Starikov

**P1.43. The use of spectral moments to simulate rototranslational collision-induced bandshapes**

[Sergei Lokshtanov](#); Daniil Chistikov; Artem Finenko; Yulia Kalugina; Sergey Petrov; Andrey Vigasin

**P1.44. The high resolution far infrared spectrum of <sup>18</sup>O enriched water vapor at the SOLEIL synchrotron**

[Mikhail Tretyakov](#); Semen Mikhailenko; Tatiana Odintsova; Olivier Pirali; Pascale Roy; Alain Campargue

**P1.45. Global deperturbation treatment of 5<sup>1</sup>Σ<sub>u</sub><sup>+</sup> and 5<sup>1</sup>Π<sub>u</sub> states in Rb<sub>2</sub>**

[Asen Pashov](#); Anna Grochola; Jacek Szczepkowski; Pawel Kowalczyk; Włodzimierz Jastrzebski

**P1.46. Revealing the multiple structures of Glutamine**

[Iker León](#); Elena Rita Alonso; Carlos Cabezas; Santiago Mata; José Luis Alonso

#### **P1.47. Linear Rayleigh and Raman scattering to second order**

Robert Cameron; [Neel Mackinnon](#)

#### **P1.48. The CRDS spectrum of natural and $^{13}\text{C}$ enriched carbon dioxide in the 1.73 $\mu\text{m}$ window**

[Ekaterina Karlovets](#); Anna Sidorenko; Peter Čermák; Didier Mondelain; Samir Kassi; Valery Perevalov; Alain Campargue

#### **P1.49. Rotational spectroscopy of organophosphorus chemical agents: Cresyl and Phenyl Saligenin Phosphate**

[Marcos Juanes San Jose](#); Rizalina T. Saragi; José L. Abad; Alberto Lesarri; Ruth Pinacho; José E. Rubio

#### **P1.50. FTIR laboratory measurement of O I spectra in the 0.77–12.5 $\mu\text{m}$ spectral range: Rydberg states and oscillator strengths**

[Vladislav E. Chernov](#); Svatopluk Civiš; Petr Kubelík; Adam Pastorek; Ekaterina Zanozina; Vladislav Chernov; Alexander Naskidashvili

#### **P1.51. Structural determination of aroma molecules in the gas-phase**

[Rihab Hakiri](#)

#### **P1.52. High resolution study of the strongly interacting $\nu_3(\text{F}_2)/\nu_1(\text{A}_1)$ bands of $^{28}\text{SiD}_4$**

[Natalia I. Raspopova](#); Olga V. Gromova; Elena S. Bekhtereva; Sigurd Bauerecker; Christian Sydow; Oleg N. Ulenikov

#### **P1.53. Molecular spectroscopy using radio-acoustic detection and high power radiation in THz region**

Mikhail Tretyakov; German Golubiatnikov; [Maxim Koshelev](#); Alexander Tsvetkov; Andrey Fokin; Mikhail Glyavin

#### **P1.54. Temperature dependence of collisional parameters of water 183-GHz line**

Mikhail Tretyakov; [Maxim Koshelev](#); Ilya Vilkov

#### **P1.55. Production of theoretical line lists for polyatomic molecules: $\text{SiH}_4$ , $\text{C}_2\text{H}_4$ , $\text{CH}_3\text{Cl}$ , $\text{CH}_3\text{F}$ , $\text{C}_2\text{H}_2$ , $\text{NH}_3$ , $\text{OH}_3^+$ , $\text{P}_2\text{H}_2$**

[Sergei N. Yurchenko](#); Jonathan Tennyson; Barry P. Mant; Alec Owens; Phillip Coles; Andrey Yachmenev; Alexander Fateev; Vladlen Melnikov

#### **P1.56. Indole rotational spectra**

[Karel Vávra](#); Kateřina Luková; Jan Koucký; Patrik Kania; Štěpán Urban

#### **P1.57. The visible spectrum of $\text{AlD}^+$ : the experimental and theoretical treatment**

[Rafal Hakalla](#); Wojciech Szajna; Keith Moore; Ian Lane; Malgorzata Ostrowska-Kopec; Izabela Piotrowska; Przemyslaw Kolek; Mirosław Zachwieja; Ryszard Kepa

#### **P1.58. High-resolution Fourier-transform spectroscopy of the comet-tail system in the $^{12}\text{C}^{17}\text{O}^+$ ion**

[Rafal Hakalla](#); Izabela Piotrowska; Malgorzata Ostrowska-Kopec; Wojciech Szajna; Przemyslaw Kolek; Marcin Rusznica; Mirosław Zachwieja; Ryszard Kepa

#### **P1.59. Self-broadening coefficients of rotational transitions of $\text{H}_2\text{S}$ measured by submillimeter-wave spectroscopy**

[Meriem Mouelhi](#); François Rohart; Francis Hindle; Cédric Bray; Gaël Mouret; Robin Bocquet; Hassen Aroui; Arnaud Cuisset

#### **P1.60. Photodissociation dynamics of $\text{CF}_2\text{I}_2$ in solution**

[Manho Lim](#)

#### **P1.61. Influence of the epigenetics marks in the non-covalent interactions between amino acids and DNA**

[Rodrigo Martínez](#); José A. Fernández; Judith Millán

#### **P1.62. Investigation on the degree of dissociation of Hydrogen plasma in PIG ion source via optical emission spectroscopy**

[Seyed Shahab Odin Salimi](#); Masoud Mahjour Shafiei; Masoomeh Yarmohammadi Satri

#### **P1.63. The optimal origin for computation of wave functions of polar molecules**

[Anastasiia Chervinskaja](#); Dmitry Dorofeev; Sergey Elfimov

#### **P1.64. 300 GHz Laboratory Heterodyne Emission Spectrometer**

[Jakob Maßen](#); Nadine Wehres; Bettina Heyne; Kirill Borisov; Katharina von Schoeler; Frank Lewen; Matthias Töpfer; Bernhard Schmidt; Henning Adams; Urs Graf; Netty Honingh; Stephan Schlemmer

# Wednesday, September 5, 2018

## Morning Sessions

Plenary Session			
Chair: <b>Agnes Perrin</b> , <i>Laboratoire de Meteorologie Dynamique</i>			
9:00–9:45 Plenary Lecture 5			
Mixelena Auditorium	PT5. Spectroscopy for satellite remote sensing of greenhouse gases: recent advances and outstanding challenges		
Chair: <b>Vivienne Payne</b> , <i>Jet Propulsion Laboratory – NASA</i>			
9:45–10:30 Plenary Lecture 6			
Mixelena Auditorium	PT6. Odorants and broadband rotational spectroscopy: what can we learn?		
Chair: <b>Maria E. Sainz</b> , <i>King's College London</i>			
10:30–11:00 Coffee Break			
Main Hall	Sponsors Stands will be placed at Main Hall		
Oral Sessions 5			
11:00–12:40	Oral Session A5:	Oral Session B5:	Oral Session C5:
	Mixelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Vladimir Tyuterev</b> , <i>University of Reims</i>	Chair: <b>Mikhail Tretyakov</b> , <i>Institute of Applied Physics –RAS</i>	Chair: <b>Sang Kuk Lee</b> , <i>Pusan National University</i>
	SPECTRAL SIMULATION AND DATABASES	LARGE MOLECULES: MW	NITROGEN AND WATER CLUSTERS
11:00–11:20	A5.1. Spectroscopic databases for the VAMDC and dat@osu portals	B5.1. High-resolution rotational spectroscopy of 3-Methylbutyronitrile – a molecule of astrophysical interest	C5.1. Microwave spectrum and nuclear quadrupole structure analysis of the NH <sub>3</sub> -N <sub>2</sub> van der Waals complex
	<b>Vincent Boudon</b> , <i>Univ. Bourgogne Franche-Comté</i>	<b>Nadine Wehres</b> , <i>University of Cologne</i>	<b>Leonid Surin</b> , <i>Institute of Spectroscopy RAS</i>
11:20–11:40	A5.2. Computing spectra of open-shell diatomic molecules with duo	B5.2. Solving discrepancies between theory and experiment: Methyl Jasmonate and Zingerone	C5.2. New millimeter-wave study of the CO-N <sub>2</sub> van der Waals complex
	<b>Sergey Yurchenko</b> , <i>University College London</i>	<b>Isabelle Kleiner</b> , <i>LISA / CNRS</i>	<b>Ivan Tarabukin</b> , <i>Institute of Spectroscopy RAS</i>
11:40–12:00	A5.3. Accurate CO <sub>2</sub> Raman spectral simulation: an algebraic alternative based on anharmonic ladder operators	B5.3. Polyalcohols as artificial sweeteners: signs of a sweetness-structure connection	C5.3. Microsolvation of ethyl carbamate conformers: a rotational spectroscopy study
	<b>Miguel Carvajal</b> , <i>Universidad de Huelva</i>	<b>Elena R. Alonso</b> , <i>Universidad de Valladolid</i>	<b>Pablo Pinacho</b> , <i>Universidad de Valladolid</i>
12:00–12:20	A5.4. Chiral rotational spectroscopy	B5.4. Microwave spectroscopic and quantum chemical investigations on 2-Acetylthiophene	C5.4. Study of the molecular structure and microsolvation (H <sub>2</sub> O N=1-3) of 2-Phenylpyridine by rotational spectroscopy
	<b>Jörg B. Götte</b> , <i>University of Glasgow</i>	<b>Christina Dindic</b> , <i>IPC RWTH Aachen University</i>	<b>Alberto Macario</b> , <i>Universidad de Valladolid</i>

12:20–12:40	A5.5. Proton transfer tunneling splittings and the imaginary mode Hamiltonian: the beginning of a beautiful friendship	B5.5. Structure and rotational spectral features of N-Bu-tyraldehyde oxime: a study using MB-FTMW	C5.5. Structures of the complexes of Cyclooctanone with water
	<b>Antonio Fernández-Ramos</b> , <i>Universidade de Santiago de Compostela</i>	<b>Nobuhiko Kuze</b> , <i>Sophia University</i>	<b>Ecaterina Burevschi</b> , <i>King's College London</i>
13:00–14:30	<b>Lunch Break</b>		
	<b>Iberdola tower (black tickets) or Deusto Library (white tickets)</b>		

## Afternoon Sessions

Oral Session 6			
14:30–15:50	Oral Session A6:	Oral Session B6:	Oral Session C6:
	Mitzelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Nobuhiko Kuze</b> , <i>Sophia University</i>	Chair: <b>Masaru Fukushima</b> , <i>Hiroshima City University</i>	Chair: <b>Bruno Martínez-Haya</b> , <i>Universidad Pablo de Olavide</i>
	MOLECULAR STRUCTURE	CHIRPED–PULSED, LAMB–DIP AND IMAGING	CLUSTERS: IR
14:30–14:50	A6.1. Microwave spectrum and equilibrium structure of Diphenyl Disulfide	B6.1. Precise lamb-dip studies of monodeuterated Methanol	C6.1. Infrared spectrum and intermolecular potential energy surface of the CO–O <sub>2</sub> dimer
	<b>Jean Demaison</b> , <i>University of Ulm</i>	<b>Alexander Lapinov</b> , <i>Institute of Applied Physics of the RAS</i>	<b>Andrew R. McKellar</b> , <i>National Research Council of Canada</i>
14:50–15:10	A6.2. Semi-experimental structure of vinyl acetate determined by microwave spectroscopy and ab initio methods	B6.2. State-to-state rotational rate coefficients for NH <sub>3</sub> –NH <sub>3</sub> collisions obtained with pump-probe chirped-pulse experiments	C6.2. Production and study of ionic clusters by photodissociation spectroscopy
	<b>Stefanie Genuit</b> , <i>Leibniz Universität Hannover</i>	<b>Christian Endres</b> , <i>MPI for Extraterrestrial Physics</i>	<b>Raghd Bejjani</b> , <i>Université catholique de Louvain</i>
15:10–15:30	A6.3. Pure rotational spectrum of <sup>15</sup> ND and isotopic-independent Dunham-type analysis of imidogen radical	B6.3. Chirped-pulse millimetre-wave spectrometer for the 140–180 GHz region	C6.3. Unravelling non-covalent interactions in DNA-base ... Xanthine clusters
	<b>Luca Bizzocchi</b> , <i>Max-Planck-Institut für extraterrestrische Physik</i>	<b>Clément Lauzin</b> , <i>Université catholique de Louvain</i>	<b>Ander Camiruaga</b> , <i>Universidad del País Vasco (UPV/EHU)</i>
15:30–15:50	A6.4. Rotational spectroscopy of the two higher energy conformers of 2-cyanobutane	B6.4. Wave-packet imaging spectroscopy of the nitrogen dimer	C6.4. High-resolution spectroscopic study of the water-carbon dioxide van der Waals complex in the near-infrared range
	<b>Marius Hermanns</b> , <i>University of Cologne</i>	<b>Yasuhiro Ohshima</b> , <i>Tokyo Institute of Technology</i>	<b>Clément Lauzin</b> , <i>Université catholique de Louvain</i>
17:00–19:00	<b>XXV Jubilee Session – Guggenheim Museum</b> <b>Joyful Pursuit of Molecular Dynamics and Spectra</b> <b>Dudley R. Herschbach</b> , <i>Harvard University</i>		
19:00–20:00	<b>Optional Guggenheim Museum Visit (reduced admittance fee)</b>		

Thursday, September 6, 2018

Morning Sessions

Plenary Session			
Chair: <b>Leonid Sinitsa</b> , Zuev Institute of Atmospheric Optics – RAS			
9:00–9:45 Plenary Lecture 7			
Mitxelena Auditorium	PT7. Correlated rotational alignment spectroscopy: high-resolution, absolute frequency spectroscopy in the time domain		
Thomas Schultz, Ulsan National Institute Standards and Technology			
9:45–10:30 Plenary Lecture 8			
Mitxelena Auditorium	PT8. Space exploration of Venus, Mars and beyond using (relatively) high resolution spectroscopy		
Ann Carine Vandaele, Royal Belgian Institute for Space Aeronomy			
10:30–11:00 Coffee Break			
Main Hall	Sponsors Stands will be placed at Main Hall		
Oral Sessions 7			
11:00–12:40	Oral Session A7:	Oral Session B7:	Oral Session C7:
	Mitxelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Vincent Boudon</b> , ICB Bourgogne – CNRS	Chair: <b>Qian Gou</b> , Chongqing University	Chair: <b>Tom Varberg</b> , Macalester College
	SPECTRAL MODELLING	HOMOCLUSTERS	CALIBRATION, DATABASES AND SOFTWARE
11:00–11:20	A7.1. Universal behaviour of diatomic halo states and mass sensitivities of their properties <b>Vladimir Spirko</b> , Academy of Sciences of the Czech Republic	B7.1. Observation of two dimers of phenyl ethyl alcohol using rotational spectroscopy <b>Rizalina T. Saragi</b> , Universidad de Valladolid	C7.1. Application of molecular spectroscopic databases for certification of calibration gas mixtures <b>Vitaly Beloborodov</b> , D.I. Mendeleev Institute for Metrology
11:20–11:40	A7.2. High energy states of polyatomic molecules: application to hot spectra <b>Vladimir Tyuterev</b> , University of Reims	B7.2. Self-aggregation process in CH <sub>2</sub> F <sub>2</sub> : large homo-clusters studied by rotational spectroscopy <b>Camilla Calabrese</b> , Universidad del País Vasco (UPV/EHU)	C7.2. Comb-calibrated coherent Raman spectroscopy of molecular hydrogen <b>Davide Gatti</b> , IFN/CNR & Politecnico di Milano
11:40–12:00	A7.3. Absolute vibrational assignment from fragmentary spectroscopic data in two isotopologues <b>Asen Pashov</b> , Sofia University St. Kliment Ohridski	B7.3. The nitrogen–nitrogen noncovalent interaction in the gas phase <b>Lorenzo Spada</b> , Scuola Normale Superiore	C7.3. Design and fabrication of a high-resolution Fourier-transform spectrometer with a supercontinuum laser source <b>Clément Lauzin</b> , Université catholique de Louvain

12:00–12:20	A7.4. Atmospheric oxygen mm-absorption: models review and uncertainties evaluation	B7.4. Understanding structural similarities in the Homodimers of Fluorene, Dibenzofuran and Diphenylether via broadband rotational spectroscopy	C7.4. Line intensities for the $\nu_6$ and $2\nu_3$ bands of Methyl Iodide ( $^{12}\text{CH}_3\text{I}$ )
	<b>Dmitriy Makarov</b> , <i>Institute of Applied Physics</i>	<b>Mariyam Fatima</b> , <i>Deutsches Elektronen-Synchrotron</i>	<b>Agnes Perrin</b> , <i>Laboratoire de Meteorologie Dynamique</i>
12:20–12:40	A7.5. Anharmonic IR spectra of chemical warfare agents from ab initio molecular dynamics simulations	B7.5. Investigation of the products of thermal self-polymerization of Methyl methacrylate by means of microwave spectroscopy – the rotational spectrum of the dimer	C7.5. Automatic assignment and fitting of spectra with PGOPHER
	<b>Pawel Rodziewicz</b> , <i>University of Bialystok</i>	<b>Sven Herbers</b> , <i>Leibniz Universität Hannover</i>	<b>Colin Western</b> , <i>University of Bristol</i>
13:00–14:30	<b>Lunch Break</b>		
	<b>Iberdola tower (black tickets) or Deusto Library (white tickets)</b>		

## Afternoon Sessions

Oral Session 8			
14:30–15:50	Oral Session A8:	Oral Session B8:	Oral Session C8:
	Mixelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Pierre Carçabal</b> , <i>Institut des Sciences Moléculaires d'Orsay CNRS</i>	Chair: <b>Dennis W. Tokaryk</b> , <i>University New Brunswick</i>	Chair: <b>José Luis Doménech</b> , <i>Instituto de Estructura de la Materia CSIC</i>
	MATRIX AND CONDENSED PHASES	CAVITY RING-DOWN	CLUSTERS: IR
14:30–14:50	A8.1. Does one need a high resolution in matrix isolation studies? the symmetry changes of the $^{28}\text{SiH}_4$ molecule in low-temperature matrixes	B8.1. Wavelength-meter controlled cavity ring-down spectroscopy for measurement of trace water vapor at ppt levels	C8.1. High resolution infrared laser jet-cooled spectroscopy of small van der Waals clusters with rare gases
	<b>Ruslan Asfin</b> , <i>Saint Petersburg State University</i>	<b>Hisashi Abe</b> , <i>National Metrology Institute of Japan (NMIJ)</i>	<b>Pierre Asselin</b> , <i>CNRS</i>
14:50–15:10	A8.2. Gas to solid phase transition of theobromine: crystal vs amorphous structures	B8.2. CRDS of the collision induced absorption (CIA) band of $\text{O}_2$ at 1.27 $\mu\text{m}$	C8.2. Insights into the binding of protons, cations and anions by azamacrocycles
	<b>Imanol Usabiaga</b> , <i>Università di Bologna</i>	<b>Alain Campargue</b> , <i>CNRS/ Université de Grenoble</i>	<b>Bruno Martinez-Haya</b> , <i>Universidad Pablo de Olavide</i>
15:10–15:30	A8.3. Seeing more does not necessarily mean seeing clearly	B8.3. Cavity-enhanced dispersion spectroscopy for the highest accuracy applications	C8.3. Infrared spectroscopy and quantum-chemical calculations on diacetylene complexes: a joint experimental study and theoretical investigation on $\text{OCS-H}_2\text{C}_4$ and $\text{N}_2\text{O-H}_2\text{C}_4$ dimers
	<b>Ilya Shenderovich</b> , <i>University of Regensburg</i>	<b>Daniel Lisak</b> , <i>Nicolaus Copernicus University</i>	<b>Andrea Pietropolli</b> , <i>Università Ca' Foscari Venezia – Dipartimento Scienze Molecolari e Nanosistemi</i>

15:30–15:50	A8.4. Titania- and montmorillonite- photocatalyzed synthesis of methane from carbon dioxide and the subsequent formation of nucleobases on early Mars and Earth	B8.4. H <sub>2</sub> -He scattering states observation with cavity ring-down spectroscopy	C8.4. Infrared spectra of the carbon monoxide - water dimer and larger clusters
	<b>Svatopluk Civiš,</b> <i>J. Heyrovsky Institute of Physical Chemistry</i>	<b>Magdalena Konefał,</b> <i>Nicolaus Copernicus University in Toruń</i>	<b>N. Moazzen-Ahmadi,</b> <i>University of Calgary</i>
16:00–18:30 Axular Room and Chillida Room	<b>Poster Session 2 (P2.1. a P2.63.) and Coffee Break</b>		
19:00 –20:00 Meeting point: Maritime Museum, 7 pm	<b>Boat Tour</b>		

## Poster presentations – Session 2

### **P2.1. The Rotational study of vitamin B6 form pyridoxine**

Elena Rita Alonso; Iker León; Lucie Kolesniková; Santiago Mata; José L. Alonso

### **P2.2. Rotational spectrum of Methoxy Acetaldehyde: a combined CP-FTMW and millimeter wave study**

Lucie Kolesniková; Isabel Peña; Elena R. Alonso; Belén Tercero; José Cernicharo; Santiago Mata; José L. Alonso

### **P2.3. Conformational study of Cyclooctanone using broadband rotational spectroscopy**

Ecaterina Burevschi; Isabel Peña; Maria Eugenia Sanz

### **P2.4. The monohydrate and dimer of Phenyl Ethyl Mercaptan by rotational spectroscopy**

Rizalina Tama Saragi; Marcos Juanes; Alberto Lesarri; Ruth Pinacho; José E. Rubio

### **P2.5. Spectroscopy group at the University of the Basque Country**

Camilla Calabrese, Aran Insausti; Maider Parra, Cristobal Pérez, Iciar Uriarte, Imanol Usabiaga, Emilio J. Cocinero; Francisco J. Basterretxea

### **P2.6. Nuclear spin conversion of ammonia isolated in noble gas matrices**

Haruka Nagamoto; Takeru Sugimoto; Ichiro Arakawa; Koichiro Yamakawa

### **P2.7. Simulation of rotational energy levels of the ground and first excited vibrational states of H<sub>2</sub>S and SO<sub>2</sub> molecules using the effective Hamiltonian approach**

Irina Vasilenko; Olga Naumenko; Alexander Bykov

### **P2.8. Analytical extension of hard-collision model of velocity-changing collisions in the Hartmann-Tran profile**

Magdalena Konefał; Michał Słowiński; Mikołaj Zaborowski; Daniel Lisak; Piotr Wcisło

### **P2.9. Far- and mid-infrared spectroscopy of matrix-isolated clusters and matrix-sublimated ice of D<sub>2</sub>O**

Hirokazu Nasu; Kazuki Niwata; Yuichi Azuma; Toshiya Tanaka; Ichiro Arakawa; Koichiro Yamakawa

### **P2.10. Laser absorption spectroscopy of <sup>13</sup>CH<sub>4</sub> at 80 K and 296 K near 1.73 μm**

Magdalena Konefał; Mélanie Ghysels; Didier Modelain; Samir Kassi; Alain Campargue

### **P2.11. Nuclear spin conversion in matrix-isolated water monomer and dimer**

Koichiro Yamakawa; Kiwa Yamaguchi; Ichiro Arakawa



**P2.12. Understanding metal compound chemistry in Late-Type star environments using chirped-pulse FT millimeter Wave Spectroscopy of supersonic Jets**

[Pascal Stahl](#); Guido W. Fuchs; Thomas F. Giesen

**P2.13. Tropospheric reactions of triazoles with hydroxyl radicals**

[Najoua Derbel](#); Olfa Ferchichi; Alexander Alijah

**P2.14. Flexibility at the fringes: conformations of the steroid hormone  $\beta$ -estradiol**

[Sabrina Zinn](#); Melanie Schnell

**P2.15. Franck-Condon simulated electronic spectra of gold carbene complexes**

[Agnes H H Chang](#); Bing Jian Sun; Ian J. Lin

**P2.16. FTIR measurements of the third overtone band of  $^{12}\text{C}^{16}\text{O}$**

[Gang Li](#); Alexandra Domanskaya; Alekandra Kyuberis; Volker Ebert

**P2.17. High resolution Ro-Vibrational analysis of  $\text{C}_2\text{HD}_3$  in the region of  $600 - 1150 \text{ cm}^{-1}$**

Olga Gromova; Nadezda V. Kashirina; [Elena S. Bekhtereva](#); Sigurd Bauerecker; Christian Sydow; Oleg N. Ulenikov

**P2.18. Fourier transform spectrum of  $^{34}\text{SO}_2$  in the region of the  $\nu_2$  bending fundamental band**

[Anastasiia G. Ziatkova](#); Olga V. Gromova; Elena S. Bekhtereva; Sigurd Bauerecker; Christian Sydow; Georg Mellau; Martin Quack; Oleg N. Ulenikov

**P2.19. Realistic vibrational spectra of Ethylene Based on a 3-band model**

[Sergey Yurchenko](#); Barry Mant; Jonathan Tennyson; Andrey Yachmenev

**P2.20. Line positions and strengths in the  $\nu_{10}/\nu_7$  bands of the  $\text{C}_2\text{D}_4$  molecule**

[Anna Fomchenko](#); Elena S. Bekhtereva; Olga V. Gromova; Maria A. Merkulova; Sigurd Bauerecker; Christian Sydow; Oleg N. Ulenikov

**P2.21. Direct perturbation analysis of the L-mixed  $\text{B}^1\Pi$  and  $\text{C}^1\Sigma^+$  states of LiRb**

[Ekaterina Bormotova](#); Elena Pazyuk; Andrei Stolyarov; Asen Pashov

**P2.22. High resolution study of the decades of  $\text{H}_2\text{S}$**

Tolganay Yersin; Fangce Zhang; [Elena S. Bekhtereva](#); Olga V. Gromova; Sigurd Bauerecker; Christian Sydow; Oleg N. Ulenikov

**P2.23. Absolute line strengths in bands  $\nu_9$  and  $\nu_{11}$  of the  $^{12}\text{C}_2\text{H}_4$  molecule**

[Yulia S. Aslapovskaya](#); Aleksei V. Kuznetsov; Olga V. Gromova; Elena S. Bekhtereva; Sigurd Bauerecker; Christian Sydow; Oleg N. Ulenikov

**P2.24. Analysis of resonance interactions in the bands located in the region of  $1500-1750 \text{ cm}^{-1}$  in the  $\text{C}_2\text{H}_2\text{D}_2$ -cis molecule**

Yuliya V. Konova; Ivan A. Konov; [Olga V. Gromova](#); Elena S. Bekhtereva; Sigurd Bauerecker; Christian Sydow; Oleg N. Ulenikov

**P2.25. High resolution study of  $^{73}\text{GeH}_4$  in the dyad and pentade regions**

[Natalia I. Raspopova](#); Olga V. Gromova; Elena S. Bekhtereva; Petr G. Sennikov; Maxim A. Koshelev; Irina A. Velmuzhova; Aleksandr P. Velmuzhov; Oleg N. Ulenikov

**P2.26. High resolution study of Hexades of  $\text{H}_2\text{MS}$**

Fangce Zhang; Petr A. Glushkov; Anastasiia S. Belova; [Elena S. Bekhtereva](#); Olga V. Gromova; Sigurd Bauerecker; Christian Sydow; Christof Maul; Oleg N. Ulenikov

**P2.27. First high resolution analysis of SiD<sub>4</sub> in the dyad region**

Natalia Raspopova; [Olga V. Gromova](#); Elena S. Bekhtereva; Christian Sydow; Sigurd Bauerecker; Oleg N. Ulenikov

**P2.28. Ab initio calculations of the potential energy curves and intramolecular interactions in LiRb and LiCs including spin-dependent effects**

[Ekaterina Bormotova](#); Alexander Medvedev

**P2.29. Comprehensive update of structural data in the MOGADOC database**

Rainer Rudert; Natalja Vogt; [Jürgen Vogt](#)

**P2.30. A highly accurate ab initio dipole moment surface for water: transitions extending into the ultraviolet**

[Aleksandra Kiuberis](#); Eamon Conway; Oleg Polyansky; Irina Mizus; Jonathan Tennyson; Nikolay Zobov

**P2.31. Far IR continuum absorption of H<sub>2</sub><sup>16</sup>O and H<sub>2</sub><sup>18</sup>O**

Mikhail Tret'yakov; [Tatyana Odintsova](#); Aleksandra Zibarova; Olivier Pirali; Pascale Roy; Alain Campargue

**P2.32. A complete and coherent spectral line parameter set for the full 2ν<sub>3</sub>-band of <sup>14</sup>N<sub>2</sub><sup>16</sup>O including line strengths, air- and self-induced broadening and shift coefficients**

[Gang Li](#); Viktor Werwein; Olav Werhahn; Volker Ebert

**P2.33. Rovibrational laser jet-cooled spectroscopy of the NH<sub>3</sub>-Ar complex in the ν<sub>2</sub> umbrella region of NH<sub>3</sub>: comparison between new infrared data and an ab initio calculated spectrum**

[Yacine Belkhdja](#); Pierre Asselin; Atef Jabri; Alexey Potapov; Jérôme Loreau; Ad van der Avoird

**P2.34. Exploring interstellar chemical processes of polycyclic aromatic hydrocarbons using opposite sides of the electromagnetic spectrum**

[Pragya Chopra](#); Donatella Loru; Sébastien Gruet; Amanda Steber; Hansjochen Köckert; Jasper Pechel; Jan Lahl; Sylvain Maclot; Jason Lee; Rebecca Boll; Simon Dörner; Sadia Bari; Benjamin Erk; Lanhai He; Mehdi Mohammad Kazemi; Alexander Lemmens; Felix Allum; Robert Mason; David Heathcote; Michael Burt; Dimitrios Rompotis; Pavel Olshin; Christopher Passow; Daniel Ramm; Farzaneh Ziaee; Xuemei Cheng; Nora Schirmel; Joss Wiese; Melby Johnny; Daniel Rolles; Sebastian Trippel; Terence Mullins; Anouk Rijs; Jochen Küpper; Mark Brouard; Claire Vallance; Per Johnsson; Bastian Manschewitz; Melanie Schnell

**P2.35. Laser-based hydrogen chloride measurements for biogas and biomethane applications**

Javis Nwaboh; [Alexandra Domanskaya](#); Zhechao Qu; Olav Werhahn; Volker Ebert

**P2.36. Bimolecular absorption in dry atmospheric gases at millimeter waves: new experimental data and high accuracy modeling**

Mikhail Tret'yakov; [Evgeny Serov](#); Aleksandr Balashov; Tatyana Odintsova; Maksim Koshelev; Aleksandra Zibarova

**P2.37. Exoplanetary atlas of molecular opacities: ExoMol Gallery**

[Sergei N Yurchenko](#); Jonathan Tennyson; Oleg N. Polyansky; ExoMol team

**P2.38. Advanced fitting method for mid-infrared spectra**

[Risto Sarjonen](#); Teemu Kääriäinen; Albert Manninen

**P2.39. Nuclear spin conversion among three isomers in crystal II methane measured by infrared spectroscopy**

[Takeru Sugimoto](#); Hirokazu Nasu; Ichiro Arakawa; Koichiro Yamakawa

**P2.40. LED-based Fourier transform spectroscopy of HD<sup>16</sup>O and H<sub>2</sub><sup>16</sup>O in 14800–15500 cm<sup>-1</sup> spectral region**

[Irina Vasilenko](#); Victor Serdyukov; Olga Naumenko; Leonid Sinit'sa

**P2.41. Collisional broadenings and shifts in the S<sub>0</sub>(0), S<sub>0</sub>(1) and S<sub>0</sub>(2) rotational lines of HD perturbed by He.**

[Raúl Martínez Torres](#); Dionisio Bermejo; Franck Thibault; Piotr Wcisło

**P2.42. Terahertz spectra of Ambroxol**

[Jan Koucký](#); Tereza Uhlíková; Jitka Bartnická; Ivana Wurmová; Patrik Kania; Štěpán Urban

**P2.43. Rotational spectra of 1,2,3,4-Tetrahydroquinoline**

[Karel Vávra](#); Kateřina Luková; Patrik Kania; Štěpán Urban

**P2.44. Evaluation of the current spectroscopic databases, such as GEISA-2015 and HITRAN-2016, in the frame of the preparation of the new release of GEISA**

[Agnes Perrin](#); Raymond Armante; Nicole Jacquinet; Noelle Scott; Alain Chedin

**P2.45. High-resolution FTIR spectrum of SO<sub>2</sub> molecule between 2400 and 2650 cm<sup>-1</sup>**

[Irina Vasilenko](#); Olga Naumenko; Veli-Matti Horneman

**P2.46. Precision spectroscopy and global deperturbation analysis of the A<sup>1</sup>Π(v = 0) state in <sup>13</sup>C<sup>18</sup>O**

[Rafal Hakalla](#); Talluri Trivikram; Alan Heays; Edcel Salumbides; Nelson de Oliveira; Robert Field; Wim Ubachs

**P2.47. Precision FT spectroscopy and deperturbation analysis of the  $A^1\Pi(v=0)$  state in  $^{12}C^{17}O$**

[Rafal Hakalla](#); Wojciech Szajna; Alan Heays; Nelson de Oliveira; Edcel Salumbides; Malgorzata Ostrowska-Kopec; Izabela Piotrowska; Przemyslaw Kolek; Mirosław Zachwieja; Ryszard Kepa; Robert Field; Wim Ubachs

**P2.48. Cavity ring-down spectroscopy of CH in the UV**

[Chris Medcraft](#); Harold Linnartz; Wim Ubachs

**P2.49. THz Spectroscopy: a solution to monitor spoilage indicators. The case of Atlantic salmon**

[Meriem Mouelhi](#); Lotta Kuuliala; Cédric Bray; Robin Bocquet; Arnaud Cuisset; Gaël Mouret; Frank Devlieghere; Francis Hindle

**P2.50. Structure and dynamics of substituted imidazoles in the gas phase**

[Eva Gougoula](#); Nicholas R. Walker

**P2.51. New spectral characterization of dimethylether isotopologues in the THz region**

[José M. Fernández](#); Guzmán Tejada; Miguel Carvajal; M. Luisa Senent

**P2.52. Ab Initio potential energy surface and vibration-rotation energy levels of Germanium Dicar-bide,  $GeC_2$**

[Jacek Koput](#)

**P2.53. The influence of water vapor isotopologues on the reflection coefficient of multilayer mirrors**

[Leonid Sinitisa](#); Victor Serdyukov; Alexei Lugovskoi; Michail Arshinov

**P2.54.  $H_2^{16}O$  and  $H_2^{18}O$  absorption spectra between 16,460 and 17,200  $cm^{-1}$**

[Leonid Sinitisa](#); Semen Mikhailenko; Victor Serdyukov

**P2.55. Monosulfur derivatives of methyl formate: millimeter and submillimeter wave spectra of S- and O-methyl thioformates**

[Vadym Ilyushyn](#); Atef Jabri; Roman Motiyenko; Laurent Margulès; Jean-Claude Guillemin; Olga Dorovskaya; Eugene Alekseev; Isabelle Kleiner; Belén Tercero; José Cernicharo

**P2.56. Analyses of various  $^{17}O$  and  $^{18}O$  enriched isotopic species of ozone from FTS high resolution spectra**

[Vladimir Tyuterev](#); Evgeniya Starikova; Alain Barbe; Marie-Renée De Backer

**P2.57. New version of S&MPO database on the ozone spectroscopy**

[Vladimir Tyuterev](#); Yurii Babikov; Semen Mikhailenko; Alain Barbe; Evgeniya Starikova

**P2.58. Quantitative influence of the ozone potential energy surface upon dynamics of the  $^{18}O + ^{32}O_2$  reaction**

[Vladimir Tyuterev](#); Grégoire Guillon; Pascal Honvault; Roman Kochanov

**P2.59. Analysis and theoretical modelling of the  $^{18}O$  enriched carbon dioxide spectrum by CRDS near 1.74  $\mu m$**

[Ekaterina Karlovets](#); Peter Čermák; Didier Mondelain; Samir Kassi; Alain Campargue; Sergey Tashkun; Valery Perevalov

**P2.60. Electronic structure of the molecular system  $HPS^+ / HSP^+$**

[B. Mehnen](#), [M. Hochlaf](#), [Saida Ben Yaghlane](#)

**P2.61. A global approach for deperturbation of NiH lowest-lying electronic states**

[Ilvie Havalayova](#); Ivayla Bozhinova; Amanda Ross; Patrick Crozet; Asen Pashov

**P2.62. Dispersed fluorescence from NiD excited with a CW laser, taken in a novel way with a BOMEM DA3 interferometer**

Amanda J. Ross, Patrick Crozet, Bradley G. Guislain, Ryan A. R. Harvey, Allan G. Adam, [Dennis W. Tokaryk](#)

**P2.63. Observation of a periodic many-body system**

[Klaus Müller-Dethlefs](#), [Francois Michels](#)

**P2.64. Creating, imaging, and controlling chiral molecules with electric fields**

[Andrey Yachmenev](#), Alec Owens, Jochen Küpper

Friday, September 7, 2018

Morning Sessions

Plenary Session			
Chair: <b>Amanda Ross</b> , <i>Université Lyon 1 &amp; CNRS</i>			
Plenary Lecture 9			
Mitxelena Auditorium	PT9. Nature and strength of unconventional hydrogen bonds		
Sanjay Wategaonkar, <i>Tata Institute of Fundamental Research</i>			
Plenary Lecture 10			
Mitxelena Auditorium	PT10. High resolution spectra of floppy molecules: neutral and ionic species		
Koichi M. T. Yamada, <i>National Institute of Advanced Industrial Science and Technology</i>			
Coffee Break			
Main Hall	Sponsors Stands will be placed at Main Hall		
Oral Sessions 9			
11:00–12:40	Oral Session A9:	Oral Session B9:	Oral Session C9:
	Mitxelena Auditorium	Baroja Auditorium	Oteiza Auditorium
	Chair: <b>Walther Caminati</b> , <i>University of Bologna</i>	Chair: <b>Thomas Giesen</b> , <i>University of Kassel</i>	Chair: <b>Francisco Basterretxea</b> , <i>Universidad del País Vasco (UPV/EHU)</i>
	HALOGEN AND CHALCOGEN INTERACTIONS	ASTROPHYSICAL MOLECULES	BIOMOLECULES
11:00–11:20	A9.1. Effect of fluorine substitution on non-covalent interactions in molecular complexes of aromatic compounds	B9.1. Spectroscopy of new Imine astrophysics target: Methylimino-acetonitrile ( $\text{CH}_3\text{N}=\text{CHCN}$ )	C9.1. Rotational spectroscopy signature of smelling allyl alcohols
	<b>Sonia Melandri</b> , <i>University of Bologna</i>	<b>Laurent Margules</b> , <i>Laboratoire PhLAM UMR CNRS 8523</i>	<b>Assimo Maris</b> , <i>University of Bologna</i>
11:20–11:40	A9.2. Chalcogen bond formation: The non-covalent, intermolecular sulfur-sulfur interaction of dimethyl sulfide and sulfur dioxide	B9.2. Imidazole rotation spectrum investigation for astrophysical search	C9.2. DNA sugars: in the gas phase and in solution
	<b>Daniel Obenchain</b> , <i>Leibniz Universität Hannover</i>	<b>Barbara Michela Giuliano</b> , <i>Max-Planck-Institut für extraterrestrische Physik</i>	<b>Iciar Uriarte</b> , <i>Universidad del País Vasco (UPV/EHU)</i>
11:40–12:00	A9.3. Halogen bond and internal dynamics in Perfluorocyclobutane-water	B9.3. Laboratory rotational spectroscopy of isotopic species of Methyl Mercaptan, $\text{CH}_3^{34}\text{SH}$ and $\text{CH}_3\text{SD}$ , and search for them in ISM	C9.3. Multiresponsive chromic soft materials: formation of macrocycles from carbazole-based biradicaloids
	<b>Qian Gou</b> , <i>Chongqing University</i>	<b>Olena Zakharenko</b> , <i>I. Physikalisches Institut, Universität zu Köln</i>	<b>Maria Carmen Ruiz</b> , <i>Universidad de Málaga</i>

12:00–12:20	A9.4. Rotational spectroscopy of the dimers and monohydrates of furfuryl alcohol and thenyl alcohol	B9.4. Accurate millimetre and submillimetre rest frequencies for cis- and trans-dithioformic acid, HCSSH	C9.4. Raman spectroscopy as a versatile tool to study organic biradicals
	<b>Marcos Juanes</b> , <i>Universidad de Valladolid</i>	<b>Domenico Prudeniano</b> , <i>Max-Planck-Institut für extraterrestrische Physik</i>	<b>Jose Luis Zafra</b> , <i>Universidad de Málaga</i>
12:20–12:40	A9.5. Microwave spectroscopic characterization of the S...O/N chalcogen bond	B9.5. The nanocosmos gas cell: A broadband Fourier transform millimeterwave spectrometer based on radio astronomy receivers.	C9.5. Molecular spectroscopic study on a natural uric acid type of Kidney Stone
	<b>Gang Feng</b> , <i>Chongqing University</i>	<b>Celina Bermúdez</b> , <i>Instituto de Física Fundamental CSIC</i>	<b>Mustafa Kumru</b> , <i>University of Freiburg</i>
13:00–14:30	<b>Lunch Break</b>		
	<b>Iberdola tower (black tickets) or Deusto Library (white tickets)</b>		
14:30–15:30	<b>Science Salon: Discussion Forum for students</b>		
Mixelena Auditorium	Informal meeting with Dudley R. Herschbach, Nobel Laureate in Chemistry, 1986		

## Free Afternoon

21:00–24:00	<b>Conference Dinner</b>
	<b>Euskalduna Concert Hall</b>
	Including:
	<ul style="list-style-type: none"> <li>– Award of the Pliva Prize, Prizes of the University of Valladolid and PCCP Prize to the best oral presentations from students</li> <li>– Award of the International Mez–Starck Prize 2018 of the Dr. Barbara Mez–Starck Foundation for the Advancement of Science and Research.</li> <li>– Announcement of the 20 student travel grant recipients.</li> </ul>

# Direct deperturbation analysis of the L-mixed B<sup>1</sup>Π and C<sup>1</sup>Σ<sup>+</sup> states of LiRb

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In recent years, ultracold alkali metal dimers have been found to have many promising implementations both in fundamental physics and quantum engineering. Amongst these are ultracold chemistry reactions, quantum information processing, and even the testing fundamental laws of physics. Li-containing heterodimers are particularly attractive candidates due to their high ground state dipole moment that makes it possible to manipulate them with external electric fields. However, to be able to utilize these molecules, one must first create a stable ensemble of ultracold molecules. One of the most widespread methods to accomplish this is the adiabatic transfer of ultracold Feshbach molecules from the lowest triplet electronic state to their absolute ground state ( $v=0, J=0$ ). This transfer is a multistep process which *a priori* requires precise and comprehensive spectroscopic information on the involved electronic states.

In this work, we performed a direct deperturbation analysis of all experimental term values of the B<sup>1</sup>Π and C<sup>1</sup>Σ<sup>+</sup> states that have recently become available in the literature for the <sup>6,7</sup>Li<sup>85,87</sup>Rb isotopologues [1–4]. The rigorous coupled-channel modeling Hamiltonian accounts for the local L-uncoupling effect between the crossing B<sup>1</sup>Π and C<sup>1</sup>Σ<sup>+</sup> states explicitly while the regular interactions with the remote states manifold are introduced into the Hamiltonian by the 2-nd order van Vleck's contact transformation. The initial set of the required interatomic potentials and L-uncoupling electronic matrix elements as a function of internuclear distance are obtained within the framework of high level *ab initio* electronic structure calculations [5]. The fitted non-adiabatic matrix elements and potential energy curves reproduce the overall experimental data of the interacting B<sup>1</sup>Π and C<sup>1</sup>Σ<sup>+</sup> states with experimental accuracy (0.01 cm<sup>-1</sup>). Particular attention is paid to the probe mass-invariant properties of the deperturbed structure parameters.

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[1] J. Chem. Phys. 2013, **138**, 94315

[2] Chem. Phys. Lett. 2011, **7**, 511

[3] Rev. A, 2016, **94**, 062503

[4] Phys. Rev. A, 2014, **90**, 062513

[5] PCCP, 2018, **20**, 1889