

Curriculum Vitae Dr. Dr. Danny E. P. Vanpoucke

Current Affiliation:

Universiteit Hasselt - Hasselt University
Wide Bandgap Materials (WBGm)
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Work Experience

2017– : **postdoc**
Hasselt University

In the Wide Bandgap Materials group (WBGm) of Prof. Dr. K. Haenen, at the UHasselt. (Belgium)

Teaching experience:
Teaching assistant : Biophysics
Co-lecturer : Functional Molecular Modeling

2014–2017: **FWO postdoctoral Fellow**
Ghent University and Hasselt University

In the Wide Bandgap Materials group (WBGm) of Prof. Dr. K. Haenen, at the UHasselt. (Belgium)

Teaching experience:
Teaching assistant : Biophysics
Co-lecturer : Functional Molecular Modeling
Promoter Bachelor projects:
- “*Applied Phonons – Vibrational properties of Metal-Organic Frameworks*”
- “*Germanium-Vacancy Complexes in Diamond*”

In the Center for Molecular Modeling (CMM) of Prof. Dr. Ir. V. Van Speybroeck, at the Ghent University. (Belgium)

Teaching experience:
Counselor of Master-thesis: “*Combined theoretical-experimental study of chromium doped zinc gallate phosphor.*”

2013–2014: **postdoc**
Ghent University

In the Center for Molecular Modeling (CMM) of Prof. Dr. Ir. V. Van Speybroeck, at the Ghent University. (Belgium)

- nov-dec 2012: **projectworker PMGE**
Ghent University
- At the Department of Administrative Affairs, supervising of the roll-out of the PMGE project:
- The introduction of new research groups and departments to the project
 - Monitoring, supervision, and providing support for previously introduced groups.
- 2009–2012: **Ph.D./postdoc**
Ghent University
- In the SCRiPTS group of Prof. Dr. I. Van Driessche, part of the Department of Inorganic and Physical Chemistry, at the Ghent University. (Belgium)
- Teaching experience:
Provided working classes for the courses:
- Quantum chemistry
 - Chemical bond
- Supervision of Bachelor project: “*Undecided electrons in ab initio calculations : What happens if the f electron in CeO₂ is uncertain whether to join in or not.*”
- 2005–2009: **Research assistant (OIO)**
Foundation for Fundamental Research on Matter, and University of Twente
- In the Computational Materials Science group of Prof. Dr. P. J. Kelly under supervision of Dr. G. Brocks, at the university of Twente. (The Netherlands)
- Teaching experience:
Provided working classes for the course “Introduction to Quantum Mechanics”
- 2004–2005: **Interim Teacher Physics/Chemistry**
KTA de Merodelei, Turnhout, Belgium
- 2003: **Tutor**
Memento-Mementec-Memodi, Ghent, Belgium

Education/Degrees

- 2009–2012: **Ph.D. in Chemistry**
Ghent University
- Ph.D. Thesis: Investigation of tunable buffer layers for coated superconductors: from solid state physics to quantum chemistry.
Defence: October 5th, 2012
Promoter: Prof. Dr. I. Van Driessche
Co-Promoter: Prof. Dr. P. Bultinck
- 2005–2009: **Ph.D. in Physics**
University of Twente, The Netherlands
- Ph.D. Thesis: *Ab Initio Study of Pt Induced Nanowires on Ge(001)*
Defence: September 11th, 2009
Promoters: Prof. Dr. P. J. Kelly and Dr. G. Brocks
- 1998–2004: **B. Sc. and M.Sc. Physics**
Ghent University, Belgium

Master thesis: *Bose-Einstein condensation in exactly solvable systems (in Dutch)*
Promoter: Dr. S. Rombouts
Supervisor: Ir. L. Pollet

Honors and Awards

- Cover Image of Crystal Engineering Communication Vol 17, Issue 45, page 8565, 2015
doi:10.1039/C5CE90198G
- Cover Image of Journal of Computational Chemistry Vol 34, Issue 5, pages i-ii, 2013
doi:10.1002/jcc.23239
- Young speakers award at the Belgian Physical Society annual general meeting 2011
- Finalist in the "Science as art competition" of the 2009 Materials Research Society (MRS) Spring meeting, San Fransisco, USA, april 13th-17th 2009

Grants

- 2014 FWO grant postdoctoral fellowship: *"Elucidating the role of 1-dimensional metal-oxide chains in the physical and chemical properties of porous metal-organic frameworks by means of density functional theory calculations"*
- 2013-2018 6 HPC-projects: 12.2 M CPU hours (352k), **Flemish Supercomputer Center (VSC)**, Belgium
- 2018: 3069 node-days for the project: *"New luminescent centers in diamond: Eu and Ge single centers"*
 - 2017: 3100 node-days for the project: *"Breathing behavior of flexible mixed metal Metal-Organic Frameworks."*
 - 2016: 4970 node-days for the project: *"Breathing behavior of flexible Metal-Organic Frameworks with MIL-47/53 topology"*
 - 2015: 2905 node-days for the project: *"The electronic structure of functionalized luminescent Metal-Organic Frameworks"*
 - 2014: 4725 node-days for the project: *"The electronic and magnetic structure of Breathing Metal-Organic Frameworks"*
 - 2013: 8272 node-days for the project: *"Structural and atomic charge evolution in the breathing behavior of Metal Organic Frameworks"*
- 2009 Granted 20 000 hours CPU time by the **Stichting Nationale Computer Faciliteiten (NCF)** on the Dutch Huygens supercomputer, for the project: *"Ab initio modeling of metallic monatomic nanowire arrays on Ge(001)"*

Software

VASP

Fortran95, Fortran2003, Delphi, Pascal, Java, C++(basics)

bash script, HTML, CSS, Java script

\LaTeX 2_ε, MS Office, xmgrace, Maple, gnuplot, Jasc Paint Shop

Professional Membership

Belgian Physical Society (BPS)
European Physical Society (EPS)
Materials Research Society (MRS)

Science Communication

2018 Wetenschap Uitgedokterd
2018 de Wetenschapsbattle
2018-... EOS wetenschap blog [Dutch]
2015-... Blog on personal website [English]

Language Skills

Dutch (native speaker), English (fluent), French (good), German (basics), Latin (reading)

Interests

Travel, salsa, books, games, programming

Scientific Appendix

List of Publications

A1 Publications

- [1] *Revisiting the Neutral C-Vacancy in Diamond: Localization of Electrons through DFT+U*
Danny E. P. Vanpoucke and Ken Haenen
Diam. Relat. Mater. **79**, 60-69 (2017).
- [2] *A combined experimental and theoretical investigation of the Al-Melamine reactive milling system: a mechanistic study towards AlN-based ceramics*
Seyyed Amin Rounaghi, Danny E. P. Vanpoucke, Hossein Eshghi, Sergio Scudino, Elaheh Esmaeili, Steffen Oswald, and Jürgen Eckert
J. Alloys Compd. **729**, 240-248 (2017).
- [3] *Mechanochemical synthesis of nanostructured metal nitrides, carbonitrides and carbon nitride: A combined theoretical and experimental study*
Seyyed Amin Rounaghi, Danny E. P. Vanpoucke, Hossein Eshghi, Sergio Scudino, Elaheh Esmaeili, Steffen Oswald, and Jürgen Eckert
Phys. Chem. Chem. Phys. **19**, 12414-12424 (2017).
- [4] *Linker Functionalization in MIL-47(V)-R Metal-Organic Frameworks: Understanding the Electronic Structure*
Danny E. P. Vanpoucke
J. Phys. Chem. C **121(14)**, 8014-8022 (2017).
- [5] *Mechanochemical route to the synthesis of nanostructured Aluminium nitride*
Seyyed Amin Rounaghi, Hossein Eshghi, Sergio Scudino, Anastasia Vyalikh, Danny E. P. Vanpoucke, Wolfgang Gruner, Steffen Oswald, Ali-Reza Kiani-Rashid, Mohsen Samadi-Khoshkhoo, Ulrich Scheler, and Jürgen Eckert
Scientific Reports **6**, 33375 (2016).
- [6] *First-Principles Study of Antisite Defect Configurations in ZnGa₂O₄:Cr Persistent Phosphors*
Arthur De Vos, Kurt Lejaeghere, Danny E.P. Vanpoucke, Jonas J. Joos, Philippe F. Smet, and Karen Hemelsoet
Inorg. Chem. **55(5)**, 24022412 (2016).
- [7] *Understanding intrinsic light absorption properties of UiO-66 frameworks: A combined theoretical and experimental study*
Kevin Hendrickx, Danny E.P. Vanpoucke, Karen Leus, Kurt Lejaeghere, Andy Van Yperen-De Deyne, Veronique Van Speybroeck, Pascal Van Der Voort, and Karen Hemelsoet
Inorg. Chem. **54(22)**, 10701-10710 (2015).
- [8] *Mechanical Properties from Periodic Plane Wave Quantum Mechanical Codes: The Challenge of the Flexible Nanoporous MIL-47(V) Framework*
Danny E. P. Vanpoucke, Kurt Lejaeghere, Veronique Van Speybroeck, Michel Waroquier, and An Ghysels
J. Phys. Chem. C **119(41)**, 23752-23766 (2015).
- [9] *Fine-tuning the theoretically predicted structure of MIL-47(V) with the aid of powder X-ray diffraction*
Thomas Bogaerts, Louis Vanduyfhuys, Danny E.P. Vanpoucke, Jelle Wieme, Michel Waroquier, Pascal Van Der Voort, and Veronique Van Speybroeck,
Cryst. Eng. Comm. **17(45)**, 8612-8622 (2015).

- [10] *A Flexible Photoactive Titanium Metal-Organic Framework Based on a $[Ti_3^{IV}(\mu_3-O)(O)_2(COO)_6]$ Cluster*
Bart Bueken, Frederik Vermoortele, Danny E. P. Vanpoucke, Helge Reinsch, Chih-Chin Tsou, Pieterjan Valvekens, Trees De Baerdemaeker, Rob Ameloot, Christine E. A. Kirschhock, Veronique Van Speybroeck, James M. Mayer and Dirk De Vos,
Angew. Chem. Int. Ed. **54(47)**, 13912-13917 (2015).
- [11] *Convergence of Atomic Charges with the Size of the Enzymatic Environment*
Danny E. P. Vanpoucke, Julianna Oláh, Frank De Proft, Veronique Van Speybroeck, and Goedele Roos
J. Chem. Inf. Model. **55(3)**, 564-571 (2015).
- [12] *Comment on "Europium doping induced symmetry deviation and its impact on the second harmonic generation of doped ZnO nanowires."*
Danny E. P. Vanpoucke
Nanotechnology **25(45)**, 458001 (2014).
- [13] *Quasi-1D physics in metal-organic frameworks: MIL-47(V) from first principles*
Danny E. P. Vanpoucke, Jan W. Jaeken, Stijn De Baerdemacker, Kurt Lejaeghere, and Veronique Van Speybroeck
Beilstein J. Nanotechnol. **5**, 1738–1748 (2014).
- [14] *Aliovalent Doping of CeO₂: DFT study of oxidation state and Vacancy effects*
Danny E. P. Vanpoucke, Patrick Bultinck, Stefaan Cottenier, Veronique Van Speybroeck, and Isabel Van Driessche
J. Mater. Chem. A **2**, 13723–13737 (2014).
- [15] *Rationality: A Social-Epistemology Perspective*
Sylvia Wenmackers, Danny E. P. Vanpoucke, and Igor Douven
Front. Psychol. **5**, 581 (2014).
- [16] *Modeling 1D structures on semiconductor surfaces: Synergy of theory and experiment*
Danny E. P. Vanpoucke
J. Phys.: Condensed Matter, **26(13)**, 133001 (2014). (Invited Topical Review)
- [17] *Tetravalent Doping of CeO₂: The impact of valence electron character on group IV dopant influence*
Danny E. P. Vanpoucke, Stefaan Cottenier, Veronique Van Speybroeck, Isabel Van Driessche, and Patrick Bultinck
J. Am. Ceram. Soc. **97(1)**, 258-266 (2014).
- [18] *New Functionalized Metal-Organic Frameworks MIL-47-X (X = -Cl, -Br, -CH₃, -CF₃, -OH, -OCH₃): Synthesis, Characterization and CO₂ Adsorption Properties*
Shyam Biswas, Danny E. P. Vanpoucke, Toon Verstraelen, Matthias Vandichel, Sarah Couck, Karen Leus, Ying-Ya Liu, Michel Waroquier, Veronique Van Speybroeck, Joeri Denayer, and Pascal Van Der Voort
J. Phys. Chem. C **117(44)**, 22784-22796 (2013).
- [19] *Reply to 'Comment on "Extending Hirshfeld-I to bulk and periodic materials"'*
D. E. P. Vanpoucke, I. Van Driessche, and P. Bultinck
J. Comput. Chem. **34**, 422-427 (2013).
- [20] *Extending Hirshfeld-I to bulk and periodic materials*
D. E. P. Vanpoucke, P. Bultinck, and I. Van Driessche
J. Comput. Chem. **34**, 405-417 (2013).
- [21] *Aqueous CSD approach for the growth of novel, lattice-tuned La_xCe_{1-x}O_δ epitaxial layers*
Vyshnavi Narayanan, Petra Lommens, Klaartje De Buysser, Danny E.P. Vanpoucke, Ruben Hue-

- hne, Leopoldo Molina, Gustaaf Van Tendeloo, Pascal Van Der Voort, Isabel Van Driessche
J. Materials Chem. **22**, 8476 (2012).
- [22] *Models and simulations in material science: two cases without error bars*
Sylvia Wenmackers and Danny E. P. Vanpoucke
Statistica Neerlandica **66**, 339-355 (2012).
- [23] *Tuning of CeO₂ buffer layers for coated superconductors through doping*
Danny E. P. Vanpoucke, Stefaan Cottenier, Veronique Van Speybroeck, Patrick Bultinck, and Isabel Van Driessche
Appl. Surf. Sci. **260**, 32-35 (2012).
- [24] *Probability of inconsistencies in theory revision, A multi-agent model for updating logically interconnected beliefs under bounded confidence*
S. Wenmackers, D. E. P. Vanpoucke, and I. Douven
Eur. Phys. J. B **85**, 44 (2012).
- [25] *Density functional theory study of La₂Ce₂O₇: disordered fluorite vs pyrochlore structure*
Danny E. P. Vanpoucke, Patrick Bultinck, Stefaan Cottenier, Veronique Van Speybroeck, and Isabel Van Driessche
Phys. Rev. **B 84**, 054110 (2011).
- [26] *CO adsorption on Pt induced Ge nanowires*
D. E. P. Vanpoucke and G. Brocks
Phys. Rev. **B 81**, 235434 (2010).
- [27] *Pt-induced nanowires on Ge(001): A density functional theory study*
D. E. P. Vanpoucke and G. Brocks
Phys. Rev. **B 81**, 085410 (2010).
- [28] *Density functional theory study of Pt-induced Ge(001) reconstructions*
D. E. P. Vanpoucke and G. Brocks
Phys. Rev. **B 81**, 035333 (2010).
- [29] *Formation of Pt-induced Ge atomic nanowires on Pt/Ge(001): A density functional theory study*
D. E. P. Vanpoucke and G. Brocks
Phys. Rev. **B 77**, 241308(R) (2008).

non-A1 Publications

- [1] *Computational Materials Science: Where Theory Meets Experiments.*
Danny E. P. Vanpoucke
ICACC 2015 conference proceeding, Developments in Strategic Ceramic Materials: Ceramic Engineering and Science Proceedings, **Vol. 36(8)**, 323-334 (2016), ISBN: 978-1-119-21173-0.
- [2] *Doping of CeO₂ as a Tunable Buffer Layer for Coated Superconductors: A DFT Study of Mechanical and Electronic Properties.*
Danny E. P. Vanpoucke
ICACC 2015 conference proceeding, Developments in Strategic Ceramic Materials: Ceramic Engineering and Science Proceedings, **Vol. 36(8)**, 169-177 (2016), ISBN: 978-1-119-21173-0.
- [3] *Pt Nanowires on Ge(001): Sheep in Wolf's Clothing?*
Danny E. P. Vanpoucke
Belgian Physical Society Magazine **3**, 11-16 (2011) (Feature Article).
- [4] *The formation of self-assembled nanowire arrays on Ge(001): a DFT study of platinum induced nanowire arrays in Computational Nanoscience – How to Exploit Synergy between Predictive Simulations and*

Experiment.

D. E. P. Vanpoucke and G. Brocks

(Mater. Res. Soc. Symp. Proc. **Volume 1177E**, Warrendale, PA, 2009), 1177-Z03-09.

Ph.D. Theses and Book Chapter

- [1] *Book-Chapter: "Computational Chemistry Experiment Possibilities"*
Bartomiej M. Szyja and Danny E. P. Vanpoucke
In V. Blay, L. F. Bobadilla, & A. Cabrera (Eds.), *Zeolites and metal-organic frameworks. From lab to industry.* (pp. 235-264), Amsterdam University Press, (2018).
- [2] *Investigation of tunable buffer layers for coated superconductors*
D. E. P. Vanpoucke
Ph.D. Thesis Ghent University, Belgium (2012).
- [3] *Ab Initio study of Pt Induced Nanowires on Ge(001)*
D. E. P. Vanpoucke
Ph.D. Thesis University of Twente, The Netherlands, ISBN: 978-90-365-2873-3 (2009).

In preparation

- [1] *Synthesis, thermodynamic stability and magnetic properties of nanostructured epsilon iron carbonitride powder prepared by the solid-state mechanochemical route*
Seyyed Amin Rounaghi, Danny E. P. Vanpoucke, Elaheh Esmaeili, Sergio Scudino, and Jürgen Eckert
In preparation (2018).

Editorial/Reviewing work

Guest Editor Computational Nanoscience - How to Exploit Synergy Between Predictive Simulations and Experiment
Curran Associates, Inc. (Jun 2010)
series: Materials Research Society Symposium Proceedings Volume 1177
ISBN: 9781617383960

Review A complete overview can be found at <https://publons.com/a/1216610/>.

- Diamond and Related Materials
- Journal of Superconducting Science and Technology
- Journal of Physics: Condensed Matter
- International Journal of Applied Ceramics Technology
- Journal of Chemical Theory and Computation
- Journal of Computational Chemistry
- Journal of Physical Chemistry
- Crystal Growth & Design
- Applied Physics Letters
- Materials Science in Semiconductor Processing
- Surface Science
- Frontiers in Physics
- Nanotechnology
- ACS Catalysis

Review Editor Frontiers in Physics

List of Organized Workshops & Conferences

- [1] *Condensed Matter Science in Porous Frameworks: On Zeolites, Metal- and Covalent-Organic Frameworks*
Groningen, The Netherlands, September 4 – 5th 2016.
- [2] *Spring School on Computational Tools for Materials Science*
Ghent University, April 13th to April 17th 2015.

List of Oral Presentations (as presenting author)

- [1] *Ab initio investigation of Eu-doped diamond*
29th International Conference on Diamond and Carbon Materials, Valamar Lacroma Dubrovnik, Dubrovnik, Croatia, September 2-6th, 2018.
- [2] *Adding Solids and Molecules: Metal-Organic Frameworks as Hybrid Materials*
27th International Materials Research Congress, Cancun, Mexico, August 19-24th, 2018. **[Invited]**
- [3] *Luminescent Centres in Diamond: Ge & Eu Defect-Complexes*
27th International Materials Research Congress, Cancun, Mexico, August 19-24th, 2018.
- [4] *Doping Diamond with Luminescent Centres: The Electronic Structure of Ge and Eu defect complexes*
General scientific meeting of the Belgian Physical Society 2018, Antwerp University, Antwerp, Belgium, April 11th, 2018.
- [5] *Doping diamond with Ge and Eu: Understanding the electronic structure*
International Conference on Diamond and Carbon Materials 2017, Gothia Towers, Göteborg, Sweden, September 3-7th, 2017.
- [6] *Spin dependent breathing flexible hybrid materials*
E-MRS spring meeting 2017, Palais de la musique et des congrès, Strasbourg, France, May 22-26th, 2017.
- [7] *Understanding Breathing Metal-Organic Frameworks: Linking Theory to Experiment*
CMD26 Conference, Martiniplaza Conference Center, Groningen, The Netherlands, September 4-9th, 2016.
- [8] *Metal-Organic Frameworks: When the whole is more than the sum of its parts*
General scientific meeting of the Belgian Physical Society 2016, Ghent University, Ghent, Belgium, May 18th, 2016.
- [9] *Computational Solid State Physics and Chemistry.*
Virtual Winterschool on Computational Chemistry, February 3rd-9th, 2016. **[Invited]**
- [10] *Computational Materials Science: where theory meets experiment*
39th International Conference and exposition on Advanced Ceramics and Composites, Daytona Beach, Florida, United States, January 25-30th, 2015. **[Invited]**
- [11] *Doping of CeO₂ as a tunable buffer layer for coated superconductors: A DFT study of Mechanical and Electronic properties*
39th International Conference and exposition on Advanced Ceramics and Composites, Daytona Beach, Florida, United States, January 25-30th, 2015.

- [12] *Tailoring metal-organic frameworks for adsorption applications.*
E-MRS Spring Meeting, Lille, France, May 26–30th, 2014.
- [13] *Calculating Hirshfeld-I charges in solids: implementation, pitfalls and applications*
Belgian Physical Society General Scientific Meeting 2013, Louvain-la-Neuve, Belgium, May 22nd, 2013.
- [14] *Investigation of tunable buffer layers for coated superconductors*
2013 meeting of the Belgian Ceramic Society, Diepenbeek, Belgium, April 19th, 2013.
- [15] *Atomic charges in solids: Calculating Hirshfeld-I charges for overlapping atoms from the electron density distribution*
ICAMM 2012 & VASP workshop, Nantes, France, June 11-16th, 2012.
- [16] *Tuning of CeO₂ buffer layers for coated superconductors through metal doping*
E-MRS 2011 Fall Meeting, Warsaw, Poland, September 19–23rd, 2011.
- [17] *Pt nanowires on Ge(001): Sheep in Wolf's Clothing?*
(First prize at the '2011 Young Speaker Contest)
Belgian Physical Society General Scientific Meeting 2011, Namur, Belgium, May 25th, 2011.
- [18] *DFT study of La₂Ce₂O₇: a Question of Order*
2011 meeting of the Belgian Ceramic Society, Mons, Belgium, February 7th, 2011.
- [19] *The formation of self-assembled nanowire arrays on Ge(001): a DFT study of platinum induced nanowires*
MRS Spring meeting, San Fransisco, USA, April 13–17th, 2009.
- [20] *DFT Study of Pt Induced Nanowires*
DPG Spring meeting, Dresden, Germany, March 22–27th, 2009.
- [21] *Nanowires on the Germanium-Platinum(001)-surface*
CW-study group Spectroscopy and Theory, Luntheren, The Netherlands, January 28–29th, 2008.

List of Poster Presentations (as presenting author)

- [1] *Doping Diamond with Luminescent Centres: The Electronic Structure of Ge and Eu defect complexes*
VSC userday 2018, Paleis der Academiën, Brussels, Belgium, may 22nd, 2018.
- [2] *Doping Diamond with Luminescent Centres: The Electronic Structure of Ge and Eu defect complexes*
SBDD XXIII, Cultureel Centrum Hasselt, Hasselt, Belgium, March 7-9th, 2018.
- [3] *Revisiting the neutral C-vacancy in diamond: Localization of electrons in DFT*
QCB13: Quantum Chemistry in Belgium, Universite Libre de Bruxelles, Brussels, Belgium, January 30th, 2018.
- [4] *Tuning the electronic structure in Metal-Organic Frameworks: When molecules and metal oxides come together*
Summer school: Upscaling techniques for mathematical models involving multiple scales, Hasselt University, Hasselt, Belgium, June 26-29th, 2017.
- [5] *HPC-based materials research: From Metal-Organic Frameworks to Diamond*
VSC userday 2017, Paleis der Academiën, Brussels, Belgium, June 2nd, 2017.

- [6] *Revisiting the neutral C-vacancy in diamond: Localization of electrons in DFT*
E-MRS Spring Meeting 2017, Palais de la musique et des congrès, Strasbourg, France, May 22-26th, 2017.
- [7] *Tuning the electronic structure in Metal-Organic Frameworks: When molecules and metal oxides come together*
E-MRS Spring Meeting 2017, Palais de la musique et des congrès, Strasbourg, France, May 22-26th, 2017.
- [8] *Revisiting the neutral C-vacancy in diamond: Localization of electrons in DFT*
SBDD XXII, Cultureel Centrum Hasselt, Hasselt, Belgium, March 8-10th, 2017.
- [9] *Breathing Metal-Organic Frameworks: From spin-dependent breathing to electronic structure tuning*
tUL Life Science Research Day 2016, Alden Biesen, Bilzen, Belgium, October 5th, 2016.
- [10] *Understanding Experiments: the Power of First Principle Simulations of Metal-Organic Frameworks*
Quantum Chemistry In Belgium 12, KULeuven, Kasteelpark Arenberg 41, 3001 Heverlee, Belgium, February 16th, 2016.
- [11] *Understanding Experiments: the Power of First Principle Simulations of Metal-Organic Frameworks*
Annual Scientific Meeting IAP P7/05, Hasselt, Belgium, September 11th, 2015.
- [12] *Low dimensional physics in Metal-Organic Frameworks: a DFT study of breathing MIL-47*
Belgian Physical Society Meeting, Liège, Belgium, May 13th, 2015.
- [13] *Quasi-1D Physics in Breathing Metal-Organic Frameworks*
Meeting of the Dutch Zeolite Association On hybrids and Zeolites, Ghent, Belgium, October 7th, 2014.
- [14] *New Quasi-1D Materials: DFT study of Breathing Metal Organic Frameworks.*
Annual Scientific Meeting IAP P7/05, Louvain-La-Neuve, Belgium, September 19th, 2014.
- [15] *Hirshfeld-I charges: from Molecules to Solids. Implementation, pitfalls and applications*
Quantum Chemistry in Belgium, eleventh edition, Namur, Belgium, January 23rd, 2014.
- [16] *Functionalized Metal-Organic Frameworks: MIL-47(V)+X a computational investigation of its properties*
IAP7 meeting, Ghent University, September 18th, 2013.
- [17] *Calculating Hirshfeld-I charges in solids: implementation, pitfalls and applications*
DFT2013, Durham, UK, September 9–13th, 2013.
- [18] *Chemistry in Solids; Extending the Hirshfeld-I method.*
Quantum Chemistry in Belgium, tenth edition, Etterbeek, Belgium, February 10th, 2012.
- [19] *Tuning of CeO₂ buffer layers through doping*
Seventh International Conference on Inorganic Materials, Biarritz, France, September 12–14th, 2010.
- [20] *CO Adsorption on Pt induced Nanowires*
Mesa⁺-day, Enschede, The Netherlands, September 21st, 2009.
- [21] *Road To Nanotechnology*
Science as Art competition at the MRS Spring meeting, San Fransisco, USA, April 13–17th, 2009.

- [22] *Formation of Pt Induced Nanowires on Ge(001): a DFT study*
CW-study group Spectroscopy and Theory, Luntheren, The Netherlands, January 26–27th, 2009.
- [23] *The Germanium-Platinum(001) surface as a template for nanowires*
ICTP 14th international workshop on computational physics and material science: Total energy and force methods, Trieste, Italy, January 8–10th, 2009.
- [24] *The Germanium-Platinum(001) surface as a template for nanowires*
CECAM Workshop: Structural, electronic and transport properties of quantum wires, Lyon, France, June 9–12th, 2008.
- [25] *The Germanium-Platinum(001) surface as a template for nanowires*
FOM Physics@Veldhoven, Veldhoven, the Netherlands, January 22–23rd, 2008.
- [26] *Depositing Pt on Ge(001): Where do the Pt Atoms Go?*
Mesa⁺-day, Enschede, The Netherlands, September 11th, 2007.
- [27] *Depositing Pt on Ge(001): Where do the Pt Atoms Go?*
CW-study group Spectroscopy and Theory, Luntheren, The Netherlands, January 29–30th, 2007.
- [28] *Depositing Pt on Ge(001): Where do the Pt Atoms Go?*
FOM Physics@Veldhoven, Veldhoven, the Netherlands, January 23–24th, 2007.
- [29] *Surface and Interface calculations: GaAs/Ge, LaAlO₃/SrTiO₃ and Pt on Ge(001)*
NIC winterschool 2006: Computational Nanoscience: Do It Yourself!, Forschungszentrum Jülich, Germany, February 14–22nd, 2006.

International Schools and Conferences

- [1] *Summer school: “Upscaling techniques for mathematical models involving multiple scales”*
Hasselt University, Hasselt, Belgium, June 26–29th, 2017.
- [2] *SBDD XXI (Hasselt Diamond Workshop 2016)*
Cultuur Centrum Hasselt, Hasselt, Belgium, March 9–11th, 2016.
- [3] *CECAM Workshop: “Basic techniques and tools for development and maintenance of atomic scale software”*
CECAM-HQ-EPFL, Lausanne, Switzerland, October 13–17th, 2014.
- [4] *Challenges towards Exascale Computing*
Ghent University, Ghent, Belgium, May 16th, 2011.
- [5] *Frontiers of Density Functional Theory: A One Day Symposium in Honor of Weitao Yang*
Academy Palace of the Royal Flemish Academy of Belgium for Science and the Arts, Brussels, Belgium, November 15th, 2010.
- [6] *QCB9 (Quantum Chemistry in Belgium)*
Louvain-La-Neuve, Belgium, January 26th, 2010.
- [7] *Symposium on Carbon Nanostructures*
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