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🌐 <https://poranne-group.github.io/>

List of Publications and Software

February 2026

Peer-Reviewed Publications.....

53. S. J. Rodríguez, J. Santoyo-Flores, K. Młodzikowska-Pieńko, R. Gershoni-Poranne, and Sebastian Kozuch
Aromaticity Switching by Quantum Tunnelling
Chemical Science **2025**, 16, 21386.
52. K. Młodzikowska-Pieńko, J. Panda, S. Garhwal, A. Kaushansky, T. Krämer, R. Gershoni-Poranne*, and G. de Ruiter*
Toward Iron-Catalyzed Alkene Metathesis: Mapping the Reactivity and Deactivation Pathways of an Iron Metallacyclobutane
Angewandte Chemie International Edition **2025**, e202515731.
51. A. Wahab and R. Gershoni-Poranne*
From Rings to Properties: Understanding the Effect of Annulation on Pyrene
Journal of Organic Chemistry **2025**, 90, 12667.
50. S. Chakraborty, Itay Almog, and R. Gershoni-Poranne*
COMPAS-4: a Dataset of (BN)₁-Substituted Cata-Condensed Polybenzenoid Hydrocarbons – Data Analysis and Feature Engineering
Journal of Chemical Information and Modeling **2025**, 65, 5508.
⇒ **Featured as a Supplementary Front Cover**
49. B. K. Hillier, D. M. de Clercq, S. D. S. Bortolussi, S. S. Capomolla, M. P. Nielsen, K. Młodzikowska-Pieńko, Renana Gershoni-Poranne, Timothy W. Schmidt*, and Martin D. Peeks*
Photoexcited and Ground-State Diradical(oid) Character in a Triquinoc[3]radialene
Chemical Science **2025**, 16, 11331.
48. Y. Davidson, A. Philipp, S. Chakraborty, A. M. Bronstein*, and R. Gershoni-Poranne*
How Local is 'Local'? Deep Learning Reveals Locality of the Induced Magnetic Field of Polycyclic Aromatic Hydrocarbons
Journal of Chemical Physics **2025**, 162, 144101.
47. F. Khaleel, S. Chakraborty, and R. Gershoni-Poranne*
Polybenzenoid Hydrocarbons in the S₁ State: Simple Structural Motifs Predict Electronic Properties and (Anti)Aromaticity
Journal of Physical Organic Chemistry **2025**, 38, e70012.
⇒ **Featured on the Front Cover**
46. K. Dey, A. Gorai, K. Młodzikowska-Pienko, N. Fridman, I. Avigdori, R. Gershoni-Poranne*, and G. de Ruiter*
Manganese-Ketenimine Intermediates as Active Catalysts in the Michael Addition of Unactivated Nitriles to α,β-Unsaturated Ketones
Angewandte Chemie International Edition **2025**, e202423275.

45. Y. Zhu, Z. Zhou, Z. Wei, A. Tsybizova, R. Gershoni-Poranne*, and M. A. Petrukhina*
Stabilizing Contorted Doubly-Reduced Tetraphenylenes with Heavy Alkali Metal Complexation: Crystallographic and Theoretical Evidence
Chemistry – An Asian Journal **2025**, e202401498.
44. A. Wahab and R. Gershoni-Poranne*
Accelerated Diradical Character Assessment in Large Datasets of Polybenzenoid Hydrocarbons Using xTB Fractional Occupation
Physical Chemistry Chemical Physics **2025**, 27, 5973.
⇒ **Featured on the Inside Front Cover**
43. G. I. Warren, K. Mlodzikowska-Pienko, S. Jalife, I. S. Demachkie, J. I. Wu, M. M. Haley, and R. Gershoni-Poranne*
Effects of Benzoheterocyclic Annulation on the s-Indacene Core: a Computational Analysis
Chemical Science **2025**, 16, 575.
⇒ **Featured on the Inside Front Cover**
42. S. Garhwal, S. Raje, K. Mlodzikowska-Pienko, T. S. Mohammed, R. Rafaeli, N. Fridman, L. J. W. Shimon, R. Gershoni-Poranne*, and G. de Ruiter*
N₂ Dissociation vs. Reversible 1,2-Methyl Migration in PCNHCP Cobalt(I) Complexes in the Stereoselective Isomerization (E/Z) of Allyl Ethers
JACS Au **2024**, 4, 4234.
⇒ **Featured as a Supplementary Front Cover**
41. Y. Zhu, Z. Zhou, Z. Wei, A. Tsybizova, R. Gershoni-Poranne*, and M. A. Petrukhina*
What a Difference an Electron Makes: Structural Response of Saddle-Shaped Tetraphenylenes to One and Two Electron Uptake
ChemistryEurope **2024**, 2, e202400055.
⇒ **Featured on the Front Cover**
⇒ **Highlighted in ChemistryViews**
40. S. Chakraborty, E. Mayo Yanes, and R. Gershoni-Poranne*
Hetero-Polycyclic Aromatic Systems: A Data-Driven Investigation of Structure-Property Relationships
Beilstein Journal of Organic Chemistry **2024**, 20, 1817.
39. A. Wahab and R. Gershoni-Poranne*
COMPAS-3: a dataset of peri-condensed polybenzenoid hydrocarbons
Physical Chemistry Chemical Physics **2024**, 26, 15344.
38. S. Jalife, A. Tsybizova, R. Gershoni-Poranne*, and J. I. Wu*
Modulating Paratropicity in Heteroarene-Fused Expanded Pentalenes
Organic Letters **2024**, 26, 1293.
37. E. Mayo Yanes, S. Chakraborty, and R. Gershoni-Poranne*
COMPAS-2: a Dataset of Cata-Condensed Hetero-Polycyclic Aromatic Systems
Scientific Data **2024**, 11, 97.
36. Z. Yang, R. Nandi, A. Orieshyna, R. Gershoni-Poranne, S. Zhang*, and N. Amdursky*
Light-Triggered Enhancement of Fluorescence Efficiency in Organic Cages
Journal of Physical Chemistry Letters **2023**, 15, 136.
35. T. Weiss, E. Mayo Yanes, S. Chakraborty, A. M. Bronstein* and R. Gershoni-Poranne*
Guided Diffusion for Inverse Molecular Design

Nature Computational Science **2023**, *3*, 873.

⇒ **Featured on the *Front Cover***

⇒ **Highlighted in *News & Views***

34. M. Pennachio, Z. Wei, R. G. Clevenger, K. V. Kilway, A. Tsybizova, R. Gershoni-Poranne* and M. A. Petrukhina*
Repercussions of Multi-Electron Uptake by a Twistacene: A Reduction-Induced Double Dehydrogenative Annulation
Organic Chemistry Frontiers **2023**, *10*, 5823.
⇒ **Featured on the *Inside Front Cover***
33. R. Gershoni-Poranne* and A. Tsybizova
A Crowning Achievement: The First Solution-Phase Synthesis of Circumcoronenes
Angewandte Chemie Int. Ed. **2023**, *62*, e202305289.
32. T. Weiss, A. Wahab, A. M. Bronstein and R. Gershoni-Poranne*
Interpretable Deep-Learning Unveils Structure-Property Relationships in Polybenzenoid Hydrocarbons
Journal of Organic Chemistry **2023**, *88*, 9645.
⇒ **Featured on the *Front Cover***
31. M. Pennachio, Z. Zhou, Z. Wei, A. Tsybizova, R. Gershoni-Poranne*, and M. A. Petrukhina*
Interplay of Charge and Aromaticity Upon Chemical Reduction of p-Quinquephenyl with Alkali Metals
Organometallics **2023**, *42*, 2492.
30. S. Fite, A. Wahab, E. Paenurk, Z. Gross and R. Gershoni-Poranne*
Text-Based Representations with Interpretable Machine Learning Reveal Structure-Property Relationships of Polybenzenoid Hydrocarbons
Journal of Physical Organic Chemistry **2022**, *36*, e4458.
⇒ **Invited contribution for the special issue on Excited State Aromaticity and Antiaromaticity**
29. A. Wahab, L. Pfuderer, E. Paenurk, and R. Gershoni-Poranne*
The COMPAS Project: A Computational Database of Polycyclic Aromatic Systems. Phase 1: catalyzed Condensed Polybenzenoid Hydrocarbons
Journal of Chemical Information and Modeling **2022**, *62*, 3704.
⇒ **Featured on the *Front Cover***
28. Z. Zhou, D. T. Egger, C. Hu, M. Pennachio, Z. Wei, R. K. Kawade, Ö. Üngör, R. Gershoni-Poranne*, M. A. Petrukhina*, and I. V. Alabugin*
Localized Antiaromaticity Hot-spot Drives Reductive Dehydrogenative Cyclizations in Bis- and Mono-Helicenes
Journal of the American Chemical Society **2022**, *144*, 12321.
⇒ **Featured on the *Front Cover***
27. E. Paenurk* and R. Gershoni-Poranne*
Simple and Efficient Visualization of Aromaticity: Bond Currents Calculated from NICS Values
Physical Chemistry Chemical Physics **2022**, *24*, 8631.
⇒ **Featured on the *Front Cover***
26. R. Thenarukandiyil, E. Paenurk, A. Wong, N. Fridman, A. Karton, R. Carmieli, G. Ménard, R. Gershoni-Poranne*, and G. de Ruiter*
Extensive Redox Non-Innocence in Iron Bipyridine-Diimine Complexes: a Combined Spectroscopic and Computational Study
Inorganic Chemistry **2021**, *60*, 18296.

25. Z. Zhou, Ö. Üngör, Z. Wei, M. Shatruk*, A. Tsybizova, R. Gershoni-Poranne*, and M. A. Petrukhina*
Tuning Magnetic Interactions Between Triphenylene Radicals by Variation of Crystal Packing in Structures with Alkali Metal Counterions
Inorganic Chemistry **2021**, *60*, 14844.
24. G. Markert, E. Paenurk, and R. Gershoni-Poranne*
Prediction of Spin Density, Baird-Antiaromaticity, and Singlet-Triplet Energy Gap in Triplet-State Polybenzenoid Systems from Simple Structural Motifs
Chemistry - A European Journal **2021**, *27*, 6923.
⇒ Selected for a *Cover Feature*
⇒ Denoted as a *Hot Paper*
23. E. Paenurk, S. Feusi, and R. Gershoni-Poranne*
Predicting Bond-currents in Polybenzenoid Hydrocarbons with an Additivity Scheme
Journal of Chemical Physics **2021**, *154*, 024110.
⇒ Invited contribution for the Issue Honoring Women in Chemical Physics and Physical Chemistry
22. M. A. Hope, T. Nakamura, P. Ahlawat, A. Mishra, M. Cordova, F. Jahanbakhshi, M. Mladenović, R. Runjhun, L. Merten, A. Hinderhofer, B. I. Carlsen, D. J. Kubicki, R. Gershoni-Poranne, T. Schneeberger, L. C. Carbone, Y. Liu, S. M. Zakeeruddin, J. Lewinski, A. Hagfeldt, F. Schreiber, U. Rothlisberger, M. Grätzel*, J. V. Milić*, and L. Emsley*
Nanoscale Phase Segregation in Supramolecular pi-Templating for Hybrid Perovskite Photovoltaics from NMR Crystallography
Journal of the American Chemical Society **2021**, *143*, 1529.
21. T. Schnitzer, E. Paenurk, N. Trapp, R. Gershoni-Poranne, and H. Wennemers*
Peptide–Metal Frameworks with Metal Strings Guided by Dispersion Interactions
Journal of the American Chemical Society **2021**, *143*, 644.
20. A. Wahab, F. Fleckenstein, S. Feusi, and R. Gershoni-Poranne*
Predi-XY: A Python program for automated generation of NICS-XY-Scans based on an Additivity Scheme
Electronic Structure **2020**, *2*, 047002.
⇒ Invited contribution for the Emerging Leaders issue
⇒ Selected as Editor's Choice paper
19. E. Solel, D. Pappo, O. Reany, T. Mejuch, R. Gershoni-Poranne, M. Botoshansky, A. Stanger, and E. Keinan*
Flat corannulene: when a transition state becomes a stable molecule
Chemical Science **2020**, *11*, 13015.
18. S. Eichenberger, M. Hönig, M. J. R. Richter, R. Gershoni-Poranne*, and E. M. Carreira*
Ring-fused cyclobutanes via cycloisomerization of alkylidenecyclopropane acylsilanes
Chemical Science **2020**, *11*, 5294.
17. M. A. Ruiz-Preciado, D. J. Kubicki, A. Hofstetter, L. McGovern, M. H. Futscher, A. Ummadisingu, R. Gershoni-Poranne, S. M. Zakeeruddin, B. Ehrler, L. Emsley*, J. V. Milić*, and M. Grätzel*
Supramolecular Modulation of Hybrid Perovskite Solar Cells via Bifunctional Halogen Bonding Revealed by Two-Dimensional ¹⁹F Solid-State NMR Spectroscopy
Journal of the American Chemical Society **2020**, *142*, 1645.
16. Z. Zhou, R. K. Kawade, Z. Wei, F. Kuriakose, Ö. Üngör, M. Jo, M. Shatruk, R. Gershoni-Poranne*, M. A. Petrukhina*, and I. V. Alabugin*

Negative charge as a lens for concentrating antiaromaticity: using pentagonal "defect" and helicene strain for cyclizations

Angewandte Chemie Int. Ed. **2020**, 59, 1256.

15. P. Finkelstein and R. Gershoni-Poranne*
An Additivity Scheme for Aromaticity: The Heteroatom Case
ChemPhysChem **2019**, 20, 1508.
14. J. V. Milić, C., N. Hellou, F. Isenrich, R. Gershoni-Poranne, D. Neshchadin, S. Egloff, N. Trapp, L. Ruhlmann, C. Boudon, G. Gescheidt, J. Crassous, and F. Diederich*
Light-Responsive Pyrazine-Based Systems: Probing Aromatic Diarylethene Photocyclization
Journal of Physical Chemistry C **2018**, 122, 19100.
13. R. Gershoni-Poranne,* A. P. Rahalkar, and A. Stanger*
The Predictive Power of Aromaticity: Quantitative Correlation between Aromaticity and Ionization Potentials and HOMO-LUMO Gaps in Oligomers of Benzene, Pyrrole, Furan, and Thiophene
Physical Chemistry Chemical Physics **2018**, 20, 14808.
12. R. Gershoni-Poranne*
Piecing it Together: An Additivity Scheme for Aromaticity using NICS-XY-Scans
Chemistry – A European Journal **2018**, 24, 4165.
11. S. Künzi, R. Gershoni-Poranne, and P. Chen*
Mechanistic Studies on the Nickel-Catalyzed Cyclopropanation with Lithiomethyltrimethylammonium Triflate
Organometallics **2019**, 38, 1928.
10. P. Chen* and R. Gershoni-Poranne
Response to "Covalent Bonding and Charge Shift Bonds: Comment on 'The Carbon–Nitrogen Bonds in Ammonium Compounds Are Charge Shift Bonds'"
Chemistry – A European Journal **2017**, 23, 18325.
9. E. Paenurk, R. Gershoni-Poranne, and P. Chen*
Trends in Metallophilic Bonding in Pd-Zn and Pd-Cu Complexes
Organometallics **2017**, 36, 4854.
8. R. Gershoni-Poranne and P. Chen*
The C-N Bonds in Ammoniums are Charge Shift Bonds
Chemistry – A European Journal **2017**, 23, 4659.
7. R. Gershoni-Poranne and A. Stanger*
Magnetic Criteria of Aromaticity
Invited Review Chemical Society Reviews **2015**, 44, 6597.
6. M. Schaffroth, R. Gershoni-Poranne, A. Stanger*, and U. H. F. Bunz*
Tetraazacenes Containing Four-membered Rings in Different Oxidation States. Are They Aromatic? A Computational Study
Journal of Organic Chemistry **2014**, 79, 11644.
5. R. Gershoni-Poranne and A. Stanger*
The NICS-XY-Scan: Identification of Local and Global Ring Currents in Multi-Ring Systems
Chemistry – A European Journal **2014**, 20, 5673.

4. R. Gershoni-Poranne, C. M. Gibson, P. W. Fowler, and A. Stanger*
Concurrence between Current Density, Nucleus-Independent Chemical Shifts, and Aromatic Stabilization Energy: The Case of Isomeric [4]- and [5]Phenylenes
Journal of Organic Chemistry **2013**, *78*, 7544.
3. R. Gershoni-Poranne and A. Stanger*
An MO-Based Identification of Charge-Shift Bonds
ChemPhysChem **2012**, *13*, 2377.
2. M. Standera, R. Haefliger, R. Gershoni-Poranne, A. Stanger, G. Jeschke, J. D. van Beek, and A. D. Schlüter*
Evidence for Fully Conjugated Double-Stranded Cycles
Chemistry – A European Journal **2011**, *17*, 12163.
1. R. Gershoni-Poranne, D. Pappo, E. Solel, and E. Keinan*
Corannulene Ethers Via Ullmann Condensation
Organic Letters **2009**, *11*, 5146.

Preprint / Under Review.....

3. M. U. G. Khan, K. Młodzikowska-Pieńko, R. Gershoni-Poranne,* and J. I. Wu*
Three Wrongs Can Make a Right: Concealing Antiaromaticity in π -Expanded [4n]–[4n]–[4n] Frameworks
September **2025**, *Under review at Chemical Science*
2. N. Barel, S. Mujaheed, K. U. Ansari, K. Młodzikowska-Pieńko, R. Gershoni-Poranne,* and Y. Tulchinsky*
Cationic Sulfonium-based Tripodal Ligand and its Rh(I) Complexes
August **2025**, *Under review at Chemical Science*
1. S. Raje, K. Młodzikowska-Pieńko, A. Stanger, R. Gershoni-Poranne,* and G. de Ruiter*
Metallaaromaticity Reimagined: Metallaaromatic Cobalt Macrocycles Through Metal-Ligand Coordination Chemistry
June **2025**, *Under review at the Journal of American Chemical Society*

Book Chapters.....

1. R. Gershoni-Poranne* and A. Stanger*
Chapter 4: NICS – Nucleus Independent Chemical Shifts
in Aromaticity: Modern Computational Methods and Applications, **2021**
Edited by I. Fernandez.

Software.....

Notes: All of our software is freely available to download from the *Poranne Group Repository*.

2. BC-Wizard
Python package implementing the NICS2BC method for calculating bond-currents from NICS values.
1. Predi-XY
Python package implementing an additivity scheme for rapid generation of NICS-XY-Scans for polycyclic aromatic systems.