KEMI

Forskerne er opdelt efter fagområde. I kemi skelnes der mellem Teoretisk, Biologisk, Analytisk, Organisk, Nucleinsyre og Medicinalkemi. Der udbydes også projekter fra fysik og farmaci.

Teoretisk Kemi	
Erik Donovan Hedegård	3
Hans Jørgen Aagaard Jensen	4
Himanshu Khandelia	5
Jacob Kongsted	6
Biologisk Kemi	
Changzhu Wu	7
Christine McKenzie	8
MiljøKemi	
Frants Roager Lauritsen	9
Organisk Kemi	
Jan O. Jeppesen	10
Steffen Bähring	11
Nucleinsyre Kemi	
Poul Nielsen	12
Stefan Vogel	13
Medicinal Kemi	

Jasmin Mcinovic

14

Vejledere fra andre fagområder

Du kan finde beskrivelse af personens forskning i det pågældende katalog.	
Carsten Svaneborg	Fysik
Francesca Serra	Fysik
René Holm	Farmac



Associate Professor Erik D. Hedegård

Topics: Theoretical chemistry, inorganic chemistry, bio-molecules, and enzymes



Focus of the group

See also https://erikh.gitlab.io/group-page/

- Development of theoretical methods for transition complexes
- Development of theoretical methods for (metallo)enzymes and chemistry in solution
- Theoretical / computational investigations of enzyme mechanisms and metal-containing drugs

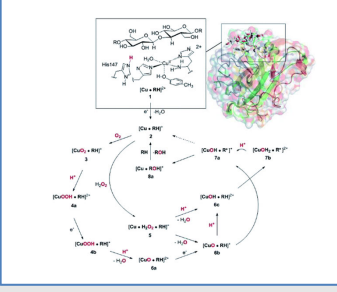
Examples of systems we currently investigate (see boxes 1–3):

- 1. Metalloenzymes mechanism and spectroscopy
- 2. Light-activated chemotherapy
- 3. Bio-mimetic complexes

1. Metalloenzymes

We employ hybrids of QM and classical methods to investigate LPMOs – a newly discovered class of enzymes used in bio-fuel production

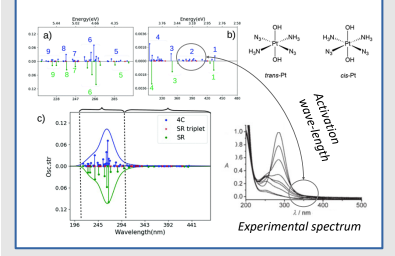
Calculated mechanism (Hedegård & Ryde, Chem. Sci. 2018)



2. Light-activated chemotherapy

We employ relativistic quantum chemical methods to investigate transition metal complexes that can be light-activated into anti-cancer agents

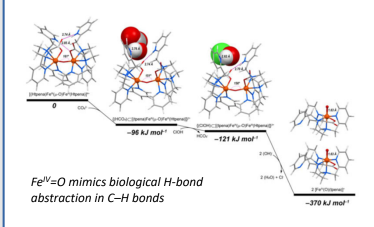
Blue: calculated (relativistic, spin-orbit) UV-vis **Green:** calculated (relativistic, no-spin orbit) UV-vis (Creutzberg & Hedegård, PCCP, 2020)



3. Bio-mimetic complexes

In collaboration with experimental groups, we investigate smaller inorganic complexes, mimicking actual enzymes

Calculated mechanism for binding of CO_3^{2-} to an iron-complex, replacement of CO_3^{2-} with HOCl, and formation of Fe^{IV} =O species (McPherson et al. submitted, 2021)



web: http://erikh.gitlab.io/group-page/

email: erdh@sdu.dk



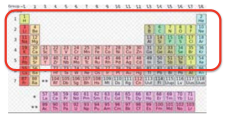
Professor Hans Jørgen Aagaard Jensen

Topics: non-relativistic (DALTON) and relativistic (DIRAC) quantum chemistry; MC-srDFT; 4c EPR and pNMR



http://daltonprogram.org

Computational quantum chemistry for primarily the *upper* half of the periodic table.

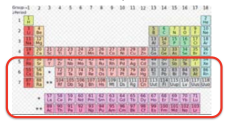


This code is based on the non-relativistic Schrödinger equation.

We develop new, efficient and flexible methods in Dalton, some of them together with Jacob Kongsted. My primary focus is currently on developing the MC-srDFT model described below.

http://diracprogram.org

Computational quantum chemistry for primarily the *lower* half of the periodic table.



This code is based on the relativistic Dirac equation. Relativistic effects are by definition effects which disappear if we artificially increase the speed of light. One example of a relativistic effect is the color of gold: if the speed of light had been the double of what it is, gold would have looked like silver!

We develop new, efficient and flexible metods in Dirac. In 2015 we were chosen by Oak Ridge Naitional Laboratory as one of 13 programs they would hellp optimizie for their supercompter Summit, which was opened for use Jan. 2019 and it is now number 1 in the world

MC-srDFT - beyond standard DFT

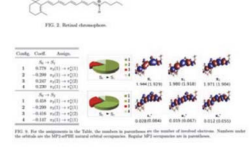
DFT – density functional theory – is today the most used method in computational quantum chemistry. Why? Because DFT often provides sufficiently accurate results without the need of a super computer, in fact many experimental papers are today supplemented with DFT calculations performed on a local work station.

However, there are many cases for which DFT is *not* sufficiently accurate. Examples are typically molecules or processes with a more complicated electronic structure as open *d*-shells of transition metal systems, chemical reactions involving surface hopping, or electronically excited states. These cases are all in principle well described by classical wave function theory (WFT), based on the Schrödinger equation as you learn it in introductory quantum chemistry/quantum mechanics courses. These WFT methods are unfortunately often very expensive, or even unfeasible ③.

My research idea with MC-srDFT is to combine the best of the two methods, and only use the more expensive WFT when needed. It looks promising \odot

If you want to participate, there are possibilities for projects with method development and implementation; or validation; or applications; or a combination.

Example of validation calculations:



EPR/ESR and pNMR computations

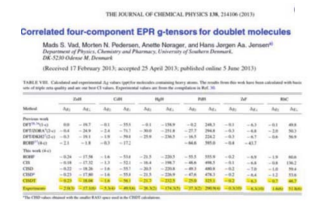
Electron paramagnetic resonance, also called electron spin resonance, is a powerful spectroscopic method for non-singlet molecules as organic radicals and inorganic chemistry involving transition metals with open \emph{d} -shells.

In an EPR/ESR experiment is measured how the electronic structure of the molecule make the g-factor deviate from the value of 2.0023 for the free electron. This deviation is in fact a relativistic effect, so with the pure non-relativistic Schrödinger equation one will always get 2.0023 – not very useful!

My research idea with an EPR/ESR module in Dirac is that using the correct theory for the effect will be the most efficient and most reliable in the long run. This project also looks promising ©

I hope soon to go on and extend this module to paramagnetic NMR – pNMR.

If you want to participate, there are possibilities for projects with method development and implementation; or validation; or applications; or a combination.



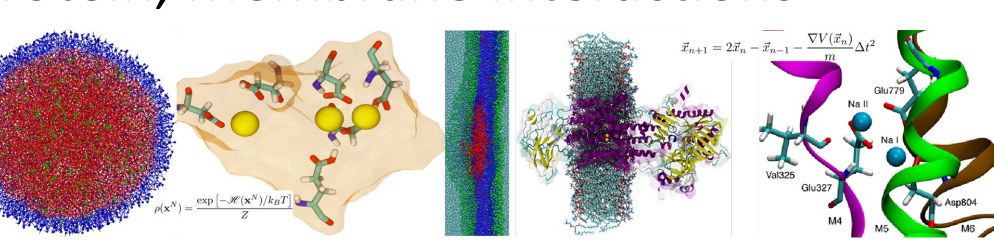
web: http://www.sdu.dk/ansat/hjj

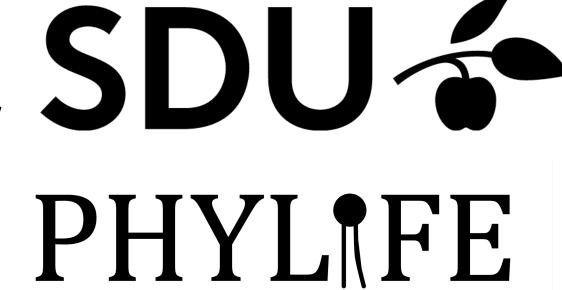
email: hjj@sdu.dk



Assoc. Professor Himanshu Khandelia SDU Topics: Computational Biophysics/Biochemistry

Protein, Membrane Interactions





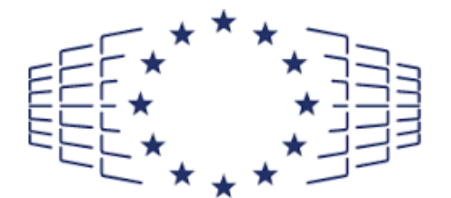
Physical LifeScience SDU &

Overview: Biomolecular Simulations

We (PhD1, 3 postdocs + students) use Molecular Simulations to explore a wide range of problem in biological systems, particularly near biomembranes, in collaboration with experiments. Examples of topics:

- The physics of biological membranes
- Nanoplastics
- Drug-membrane interactions
- Mechanisms of transport across membranes

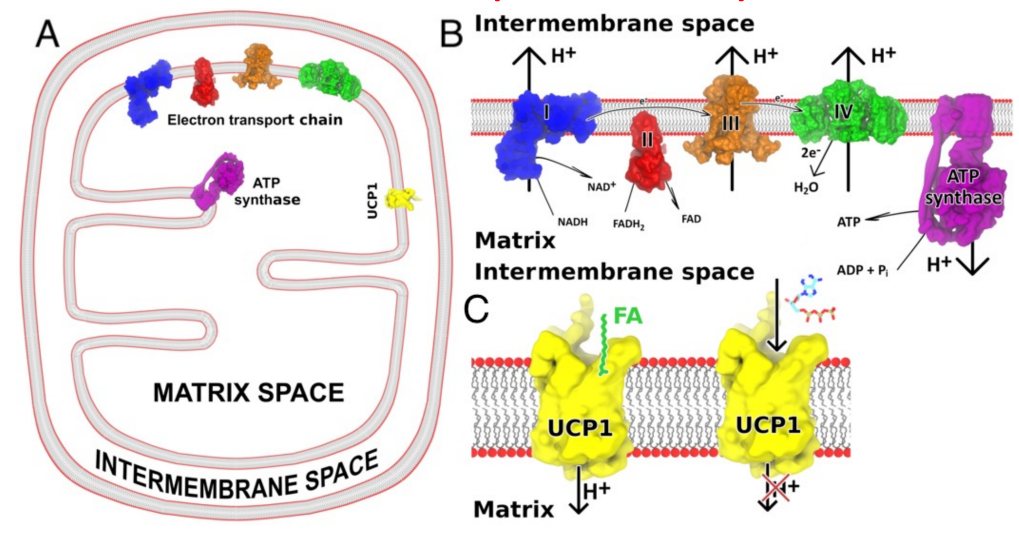
We are housed at PHYLIFE: Physical Life Sciences, and use **Supercomputers** for research





Physics, Chemistry

Proton-Coupled Transport

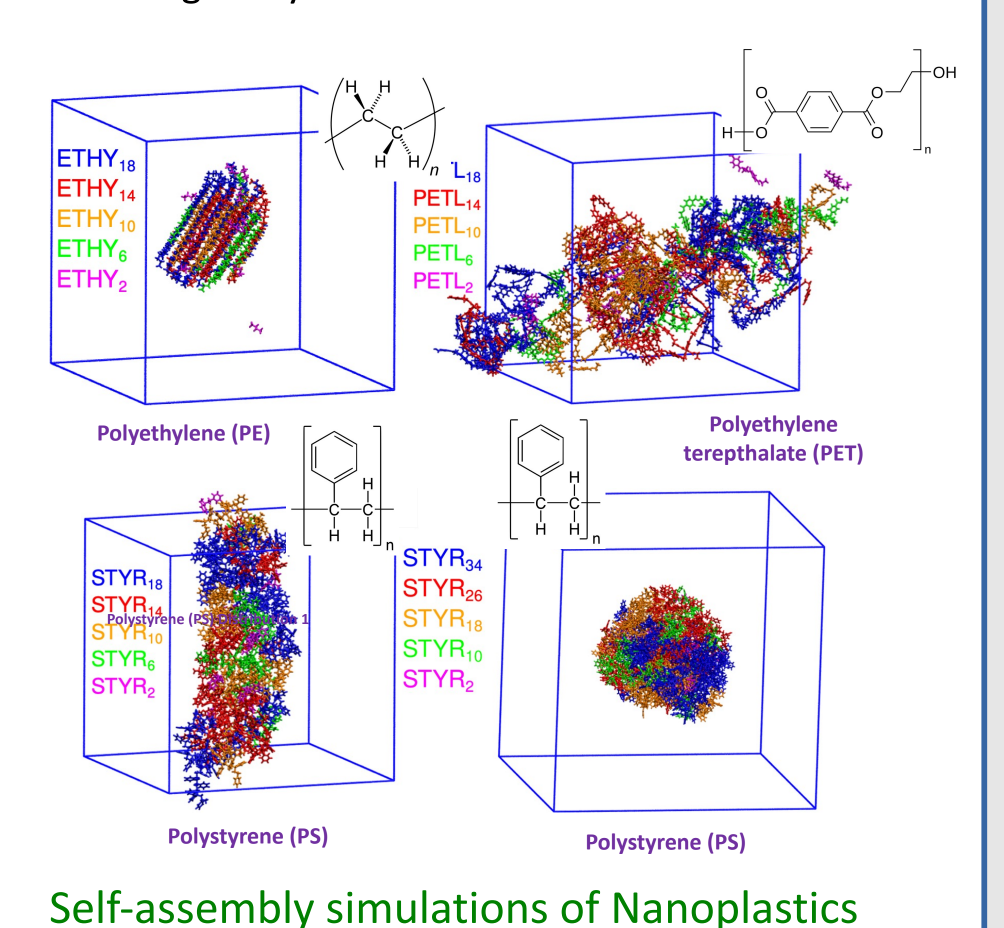


A: mitochondrion B. The electron transport chain establishes a proton gradient C. The uncoupling protein 1 dissipates the gradient to produce heat

Solute transport in mitochondrial solute carriers is often coupled to proton transport along a pH gradient. Disruption is linked to metabolic and obesity-related disease

Nanoplastics (Physics, Chemistry)

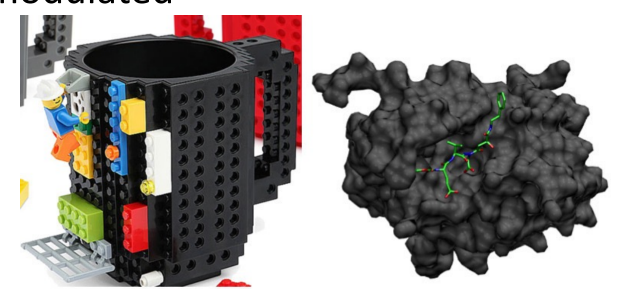
Most research on nanoplastics focusses on > 50 nm particles. What do < 50 nm particles look like? How can they be removed from water? How do they interact with biological systems?



Chemistry, Pharmacy

Protein-Drug Interactions

What are the molecular forces which determine the specificity and affinity of drugs to protein targets? How are these forces modulated

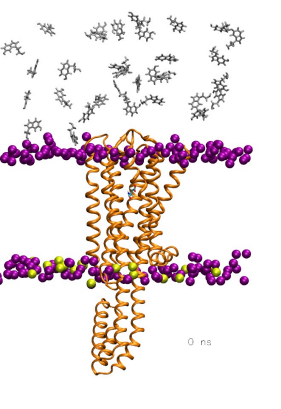


Physics, Chemistry, Pharmacy Magic Mushrooms

How does the active component of magic mushrooms cause hallucinations?

We also work with Other GPCRs

novo nordisk fonden







VILLUM FONDEN LUNDBECKFONDEN

web: hkgroup.sdu.dk

email: hkhandel@sdu.dk



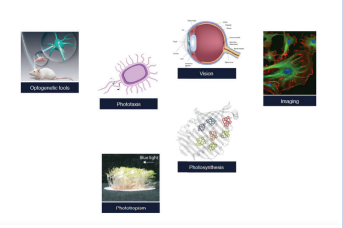
Professor Jacob Kongsted

Topics: Computational and Theoretical Physics, Chemistry and Biology



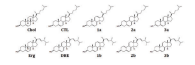
Light-driven biological processes

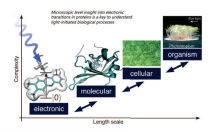
We develop and use theoretical and computational methods to study light-controlled biological processes such as photosynthesis, optogenetics etc. Our aim is to understand and develop functional biological materials at an atomistic level.



Development and applications of multiscale strategies

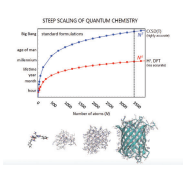
We develop and use multiscale strategies to (i) design small functional biological molecules and (ii) to rationalize biological functions. Our strategy is based on a hierarchy of methods based on either quantum or classical principles.





Method development within computational quantum chemistry

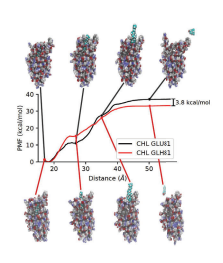
We develop novel computational methods and procedures within quantum chemistry aimed at being able to perform calculations on very big molecular systems of biological interest. Our developments are focused around quantum embedding methods and the applications are usually centered around biomolecular spectroscopy.





Computational medicinal chemistry

We develop and use computational procedures aimed at understanding the function of drug-like molecules at the atomistic level. In particular we study binding processes between macro-molecules and drugs.



email: kongsted@sdu.dk

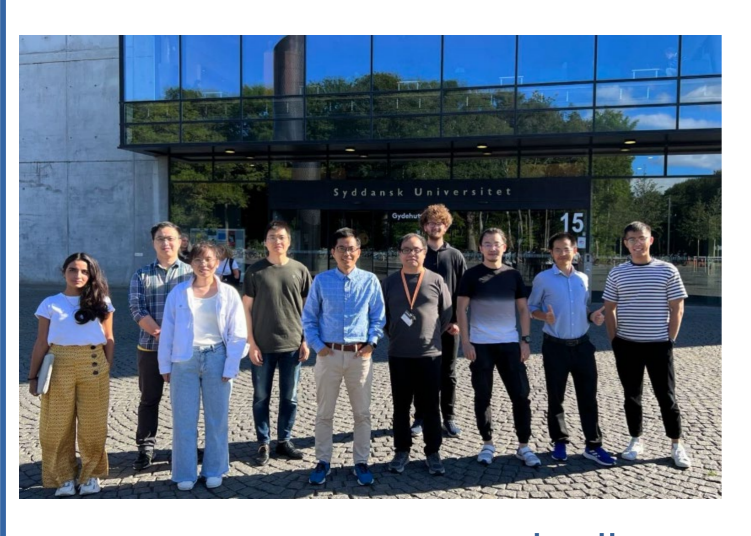


Associate Professor Changzhu Wu

Topics: Enzyme Catalysis and Protein Chemistry

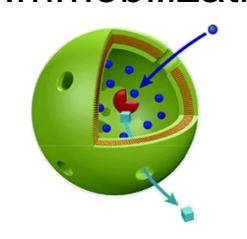


Who?

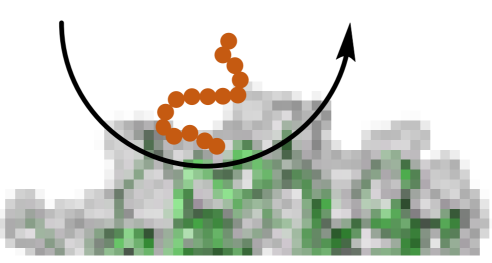


www.wugroup.sdu.dk
Enzyme Catalysis & Protein Chemistry

What? Immobilization



Proteins or enzymes



Chemical modification

Why?





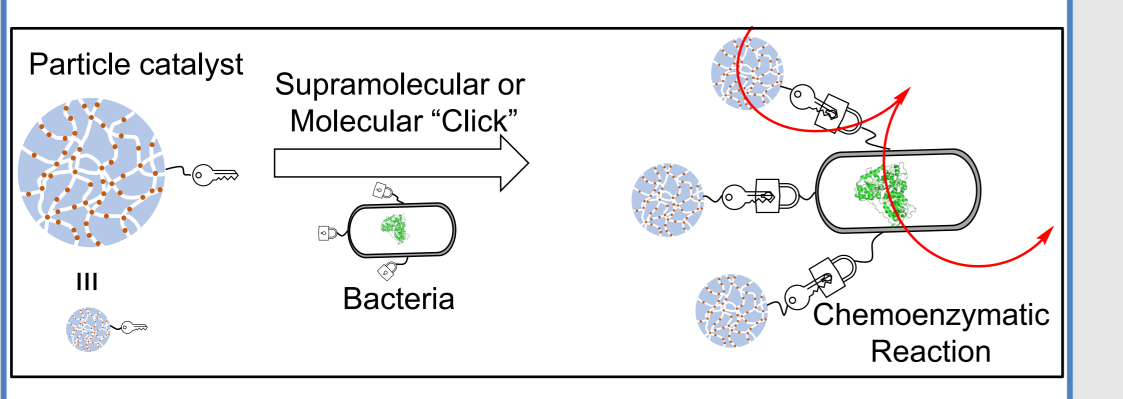
Industrial Biotechnology





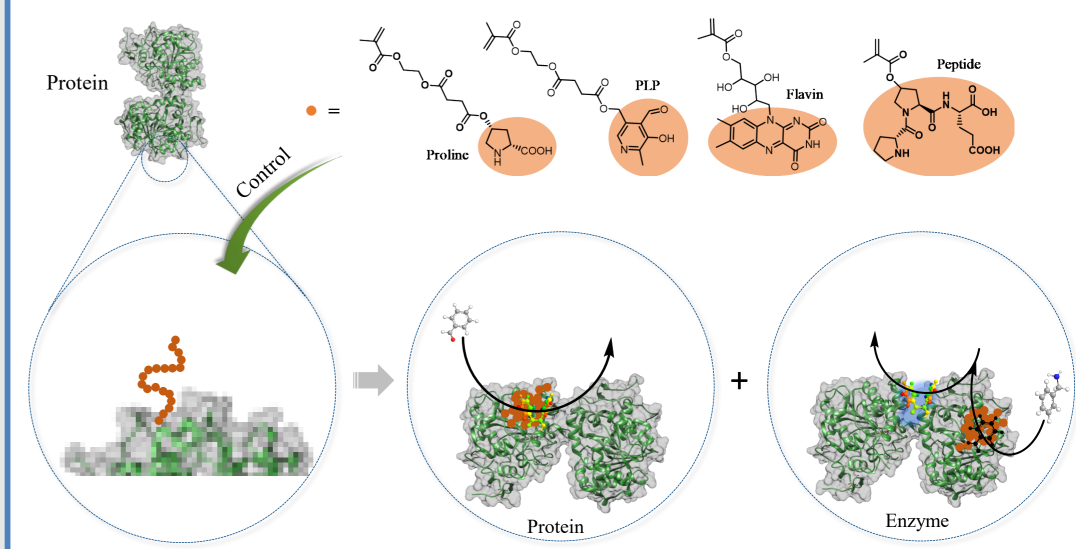
Medical Biotechnology

Project 1. Biohybrid catalysts



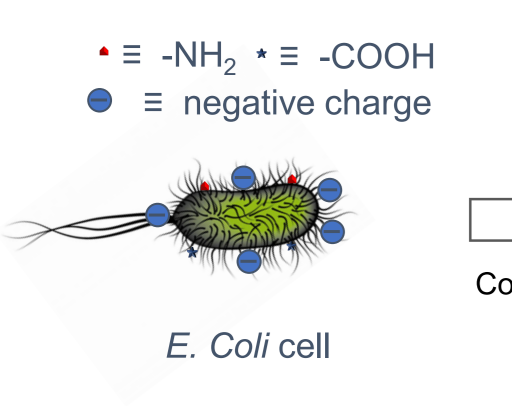
- Aim: Combine chemical catalysts with enzymes in bacteria for cascade reaction;
- **Method**: 1) Prepare particles that contain different catalysts, e.g., metal catalysts;
 - 2) modify the catalyst with active molecules;
 - 3) couple the particle with bacteria;
- Evaluation: Evaluate the catalytic efficiency.

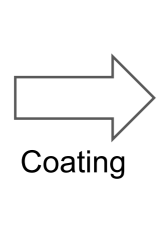
Project 2. Artificial enzymes

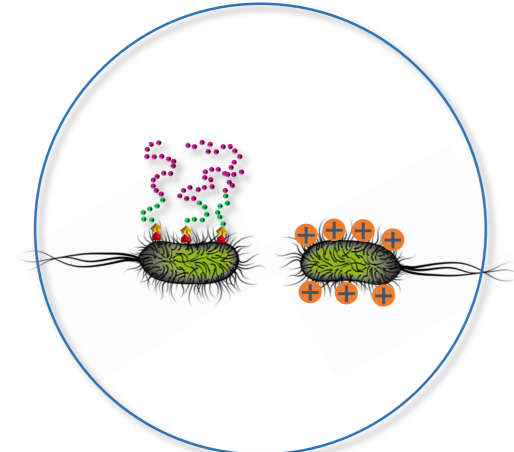


- Aim: Artificial or semisynthetic enzymes
- Method: 1) Prepare active organocatalysts;
 2) conjugate the catalysts to protein or enzymes as active polymer;
- Evaluation: 1) Characterize the conjugate; 2) evaluate the catalytic efficiency.

Project 3. Cell coating for biotechnology







- Aim: Protech cells for biocatalysis
- Method: 1) graft polymers from cells surfaces;
 2) coat cells using charge or chemical reactions;
- Evaluation: 1) characterize the system; 2)
 evaluate the catalytic efficiency

Ref.: Wu et al., Nature Communications, 13, 3142

Homepage: www.wugroup.sdu.dk

email: wu@sdu.dk



Professor Christine J. McKenzie

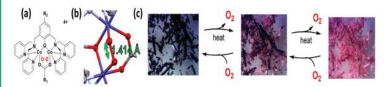
Metal-organic, Bioinorganic and Green Chemistry



Industrial processes are far from achieving enzyme-like selectivity, atom- and energy-efficiency. We synthesize new metal-organic compounds with *biomimetic* active sites with one aim being discovery of new technologies for alleviating the emission of Greenhouse (CO₂, CH₄) and toxic gases (NO_x) from anthropogenic activities.

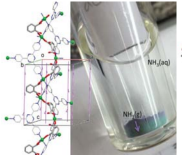
Solvent-free reactions

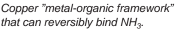
Just like the gas processing enzymes, small molecules are <u>chemi</u>sorbed by our new metal-organic materials. Reactions include reversible gas binding and toxic gas capture and conversion.

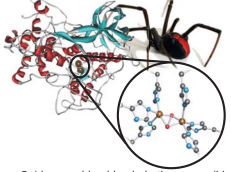


- (a) Active site structure of a material that grabs O_2 from air (b) Section of X-ray structure showing the chemisorbed O_2
- (c) Visible color changes on sorption and desorption.







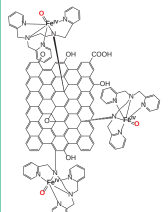


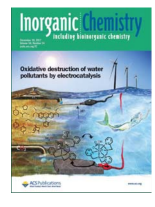
Spiders are blue-blooded - they reversibly bind O_2 at the dicopper site of hemocyanin.

Catalysis

Using our high-valent non-heme iron complexes we are developing a new electrocatalytic technology for the *total mineralization* of organic pollutants (e.g. pesticides) in contaminated water.

The next step is immobilization of the catalysts of on electrode materials e.g. graphene.





Collaboration with the Water Research Center, University of New South Wales, Sydney, AUS.

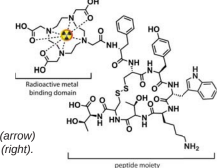
Single crystal X-ray crystallography

Co-supervisor Adjunkt prof. Vickie McKee

Radiotheranostics

Auger Electron Emitters (AEEs) hold great promise in targeted radionuclide therapy for cancer. Short decay paths means cell destruction can occur without extensive damage to surrounding tissues. We are synthesizing ligands for the *in vivo* transport of AEEs, 119-antimony (119Sb) and 58m-cobalt (58mCo). Sister isotopes (117Sb, 55Co) can be used for diagnostics.





PET/CT image of AR42J tumor (arrow) in mouse using a ⁵⁵Co complex (right).

Collaboration with Assoc. Prof. Helge Thisgaard at OUH-nuclear medicine

web: http://mckee.sdu.dk/cjmck/home.html email: mckenzie@sdu.dk



Professor Frants R. Lauritsen

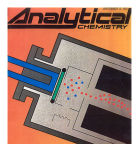


Topics: Analytical environmental chemistry, portable instrumentation, dynamics of chemical and biological systems

Membrane Inlet Mass Spectrometry

Membrane inlet mass spectrometry (MIMS) is a special type of mass spectrometry that makes it possible to monitor chemical and biological processes in real time and all kinds of samples being gaseous, liquid or solid can be analyzed directly for volatile organic compounds without sample preparation.

MIMS principle and typical applications



On-site measurement of off odours from pig manure



Ground water characterization at a nuclear power station



Online monitoring of trihalomethanes in swimming pools at Gladsaxe Sportscenter

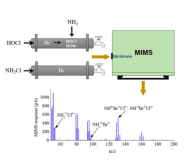


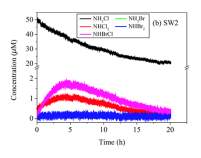
Disinfection of water and the derivative formation of disinfection byproducts

Experimental setup to investigate disinfection of surface water using chlorination (OCI or chloramination).

As part of the disinfection chemistry chloramines are transformed into more reactive bromamines

A typical kinetic experiment showing the disinfection of surface water and the associated transformation of monochloramine into dichloramine and bromchloramine





Science of the Total Environment 751 (2021) 142303

Hot Cell MIMS

In hot cell MIMS solids can be analysed directly for their content and liberation of chemicals to the surroundings. This makes it possible to evaluate the dangers associated with physical contact to the materials.

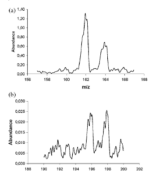
EXAMPLE: Direct analysis of soil from a factory site in Sweden that was heavily contaminated with herbicides (2,4-D and 2,4,5-T in particular) also known as agent orange. For analysis a spoonful of contaminated soil was simple dumped into the MIMS as is.

Picture of the contaminated site



Waste sampling. Half masks was needed because of the odour.

Mass spectrometric verification of the presence of the two precursors to 2,4-D and 2,3,5-T

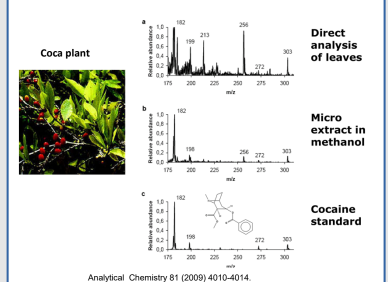


Rapid Communications of Mass Spectrometry 22 (2008) 2234-2240

Analysis of drugs

In addition to environmental samples the hot cell MIMS technique can also be used for direct analysis of active compounds in plant materials and for detection of counterfeit medicines.

EXAMPLE: Leaves from a plant analysed as is reveals its origin and content of cocaine



email: frl@sdu.dk



Professor Jan O. Jeppesen

Topics: Organic Chemistry, Supramolecular Chemistry, Analytical Chemistry and Tetrathiafulvalene



Tetrathiafulvalene (TTF)







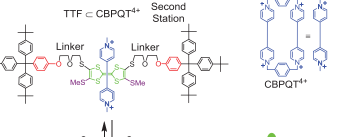
Tetrathiafulvalene's (TTF's) Properties

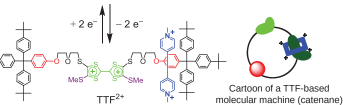
- Strong electron donor
- Readily oxidised in a stepwise and reversible manner to TTF*+ or TTF2+, either chemically or electrochemically
- 3 oxidation states (TTF, TTF*+ and TTF2+) easily distinguished using spectroscopy
- Stable under most synthetic conditions, allowing incorporation into larger systems - see below

TTF-Derivatives

- Various substituents can be added to TTF to build useful molecules
- Pyrrolo annelated TTFs, monopyrrolo-TTF (MPTTF) and bispyrrolo-TTF (BPTTF), are offen used to avoid isomerism
- MPTTF and BPTTF have comparable properties to TTF but different positions to connect to other groups - this can be helpful when designing new molecules

Molecular Machines $\mathsf{TTF} \subset \mathsf{CBPQT}^{4+}$





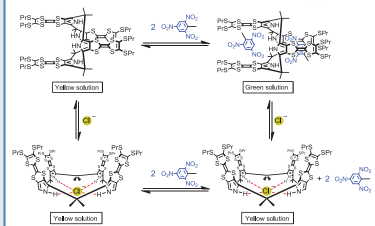
Advantages of Using TTF

- TTF is an electron donor which can interact (complexate) with cyclobis(paraquat-p-phenylene) (CBPQT4+) which is an electron acceptor
- Allows preparation of interlocked systems: rotaxanes (linear) and catenanes (cyclic)
- Oxidation of TTF destroys the interaction between TTF and CBPQT4+ and induces a movement

Challenge

Design of systems capable of using the induced movement to perform unidirectional motion (linear or rotary)

Molecular Sensors –TetrakisTTF-calix[4]pyrrole



Advantages of Using TTF

- tetrakisTTF-calix[4]pyrrole binds nitrated benzene derivatives (e.g. trinitro-toluene (TNT)) - can be used to detect explosives
- Distinct, easily visible colour change when TNT binds to tetrakisTTFcalix[4]pyrrole – helpful for use in devices

Challenges

- Chloride anions bind stronger than TNT to tetrakisTTF-calix[4]pyrrole and disrupts sensing behaviour (see above)
- TNT binds only to the 1,3 alternated configuration (top) can this be locked?

Molecular Wires

Advantages of Using TTF

- TTF's redox properties can be used to add switching behaviour to wires
- BPTTF allows for highly conjugated molecular wires this improves conductance

Challenges

- Improve solubility of wires so conductance studies can be done
- Find good anchoring groups for different electrodes (e.g. gold or graphene)

Jeppesen, J. O.; Becher, J. Eur. J. Org. Chem. 2003, 3245-3266

Andersen, S. S.; Share, A. I.; Poulsen, B. L. C.; Kørner, M.; Duedal, T.; Benson, C. R.; Hansen, S. W.; Jeppesen, J. O.; Flood, A. H. *J. Am. Chem. Soc.* 2014, 136, 6373–6384
Nielsen, K. A.; Cho, W. S.; Jeppesen, J. O.; Lynch, V. M.; Becher, J.; Sessler, J. L. *J. Am. Chem. Soc.* 2004, 126, 16296–16297
Solano, M. V.; Della Pia, E. A.; Jevric, M.; Schubert, C.; Wang, X.; van der Pol, C.; Kadziola, A.; Nørgaard, K.; Guldi, D. M.; Nielsen, M. B.; Jeppesen, J. O. *Chem. Eur. J.* 2014, 20, 9918–9929.



Assistant Professor Steffen Bähring

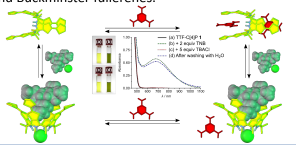
Topics: Supramolecular Chemistry and Synthetic Organic Chemistry

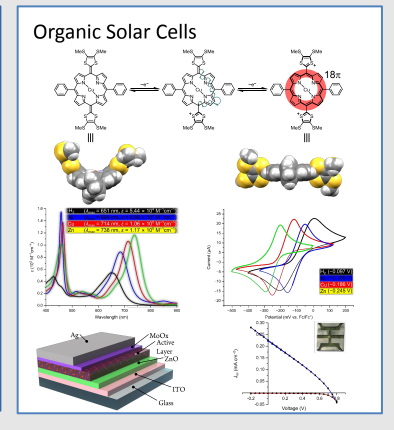


Supramolecular Chemistry

The area of Supramolecular Chemistry covers non-covalent interactions between molecules to assemble larger ordered structures. The non-covalent nature of these structures makes them highly sensitive towards external stimuli, which can be exploited in e.g. sensors and electron-transfer processes.

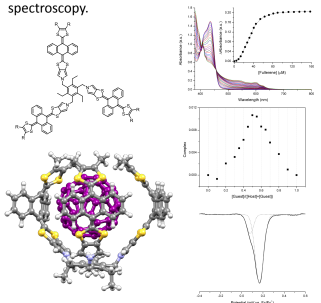
Tetrathiafulvalene is a highly electron-rich molecule that functions as a π -electron donor. This has made it ideal as a building block in Supramolecular Chemistry and the sensing of explosives (e.g. TNT) and Buckminster fullerenes.





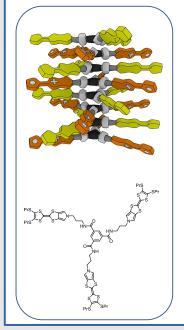
Fullerene Receptor

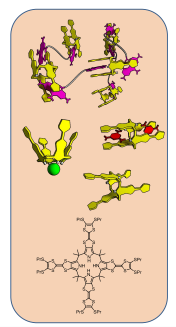
B. Sc. Project Jesper Tversted. Synthesis and investigation of extended tetrathiafulvalene receptor for fullerene by computational modelling, NMR, absorption, ITC, CV and transient spectroscopy.



Supramolecular Polymers

Concentration and temperature dependent polymerization of TTF-monomers investigated by absorption spectroscopy, NMR, AFM and DLS.







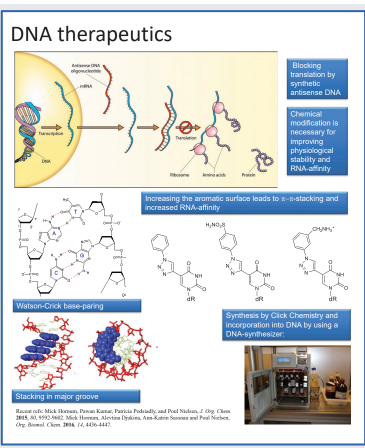
email: sbahring@sdu.dk

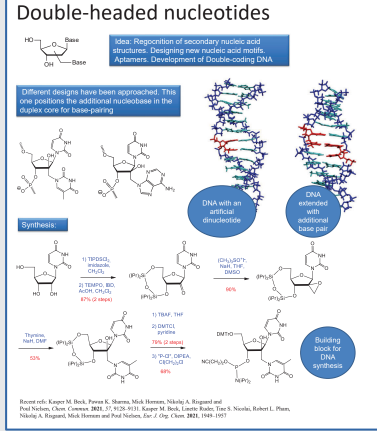


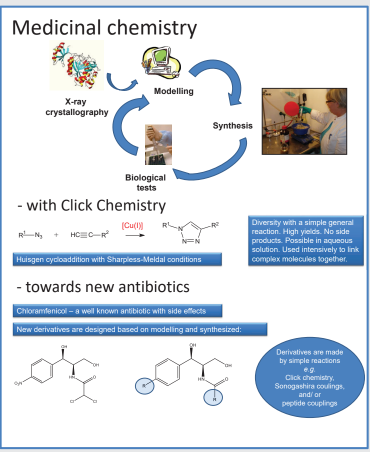
Professor Poul Nielsen

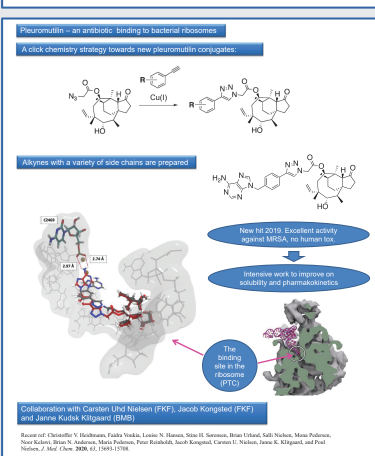
Organic Chemistry;
Synthetic DNA and Medicinal Chemistry
towards new antibiotics

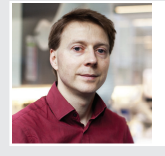












Assoc. Prof. Stefan Vogel

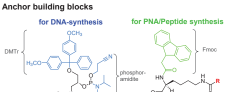


Topics:

Chemistry & Catalysis in Nanoreactors and Liposomal Drug Delivery of DNA & Peptides

Synthesis of Lipid building blocks for automated DNA & Peptide synthesis

We synthesize lipophilic building blocks for DNA, PNA and Peptides – and incorporate them into oligomers with therapeutic applications.

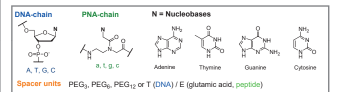




Lipid-DNA or Lipid-PNA/Peptide conjugates



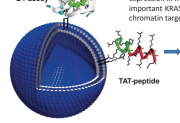




Ries, O., et al., Organic & Biomolecular Chemistry 2015, 13 (37), 9673-9680; Rabe, A. et al. Chem Commun 2017, submitted.

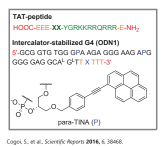
Liposomal platform for targeted drug delivery

Liposomes allow free variation of payload vs. targeting vector concentrations. Their size controls bio-distribution. Investigation for the treatment of pancreatic cancer:



Membrane-anchored DNA with therapeutic activity. A specific G-Quadruplex (G4) forming sequence can inhibit KRAS expression in mammalian cell. It is called "decoy" as it scavenges an important KRAS transcription factor (MAZ) by mimicking its natural

> Membrane anchored cell-penetrating peptide. Cell penetrating peptide aids uptake of the G4 into the cell.

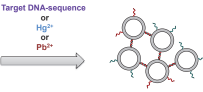


Environmental and Health Diagnostics using Nanoparticle assembly

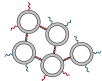
DNA-encoded liposomes assemble into large aggregates (visible by eye). Assembly can be triggered by binding of a biological target strand for DNA/RNA diagnostics or toxic metals such as mercury (stabilizes T:T mismatch), or lead (stabilizes G-quadruplexes).

Non-covalent attachment of lipid-DNA-conjugates to soft nanoparticles, like liposomes, is an attractive technology as it needs no surface chemistry.

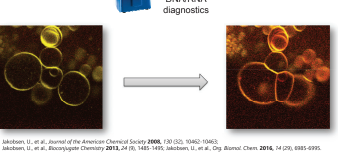






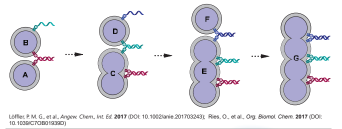


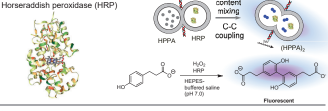
Liposome assembly

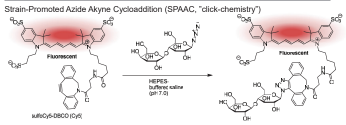


Programmed chemical and enzymatic reactions in liposomes

Liposomes as reaction flasks: DNA-programmed mixing of tiny volumes.







email: snv@sdu.dk

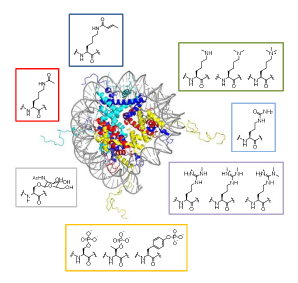


Professor Jasmin Mecinović

Topics: Medicinal Chemistry, Chemical Biology, Organic Chemistry, Bioorganic Chemistry

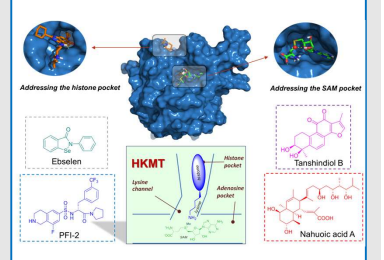


Chemical Basis of Epigenetics



- Aim: to unravel the chemical basis of epigenetics
- Objectives: to explore diverse posttranslational modifications of histone proteins
- Approach: chemical biology and physical-organic chemistry
- Techniques: synthesis, peptide chemistry, protein chemistry, enzymology
- References: Nat. Commun. 2015, 6, 8911, Commun. Chem. 2019, 2, 112, Commun. Chem. 2020, 3, 69, Commun. Chem. 2022, 5, 27.

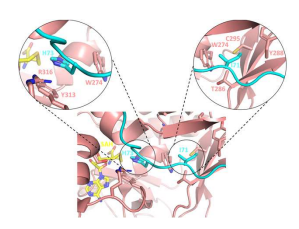
Discovery of Epigenetic Inhibitors



- Aim: to develop new epigenetic inhibitors of therapeutic potential
- Objectives: to design novel chemical probes for histone-modifying enzymes
- Approach: medicinal chemistry
- Techniques: synthesis, inhibition assays
- References: ChemMedChem 2018, 13, 1405-1413, Bioorg. Med. Chem. Lett. 2018, 28, 1234-1238, Org. Biomol. Chem. 2022, 20, 173-181.

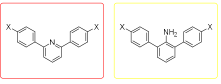
Actin Modifications

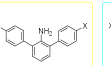


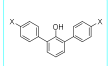


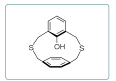
- Aim: to probe substrate scope and inhibition of actin-modifying enzymes
- Objectives: to investigate enzyme catalysis of histidine methyltransferase SETD3
- Approach: bioorganic chemistry and chemical biology
- Techniques: synthesis, peptide chemistry, enzymology
- References: ChemMedChem, 2021, 16, 2695-2702, Org. Biomol. Chem. 2022, 20 1723-1730. Protein Sci. 2022, 31, e4305.

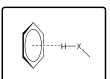
Molecular Recognition

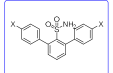




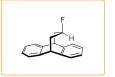


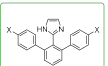


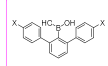




Polar-π interactions







- Aim: to probe noncovalent interactions in chemistry and biology
- Objectives: to understand interactions between aromatic rings and polar groups
- Approach: organic chemistry
- Techniques: synthesis, NMR, crystallography
- References: Org. Lett. 2020, 22, 7870-7873, Chem. Eur. J. 2021, 27, 5721-5729, Chem. Eur. J. 2022, 28, e202104044, J. Org. Chem. 2022, 87, 6087-6096.

email: mecinovic@sdu.dk