

21st IUPAC International Symposium on Solubility Phenomena and Related Equilibrium Processes (ISSP21)

Book of Abstracts

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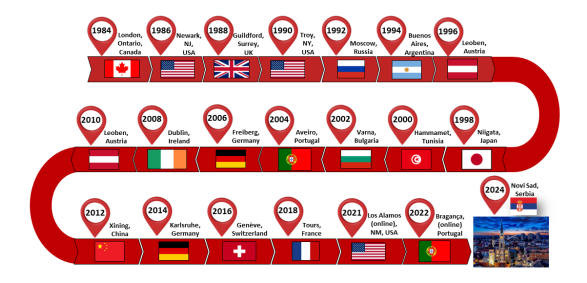
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Dear colleagues and friends,

It is a great pleasure for us to organize the 21st IUPAC International Symposium on Solubility Phenomena and Related Equilibrium Processes (ISSP21) in Novi Sad, Serbia, from 9th to 13th September 2024. ISSP became a traditional conference initiated in 1984, and the first event took place in London, Ontario Canada. It is a biennial symposium organized by the IUPAC Subcommittee on Solubility and Equilibrium Data committed to gather international experts on solubility and associated properties, to exchange scientific research and technological applications within the academic, scientific, and technical communities. It addresses the general importance of these phenomena in a variety of settings ranging from green chemistry to nuclear waste disposal, always envisaging applications for sustainable development.



After two successful online conferences organized by Los Alamos National Laboratory, USA in 2021 and Instituto Politécnico de Bragança, Portugal in 2022, ISSP21 including a Workshop on Data Analysis is organized in-person and will continue to celebrate creativity, diversity, and friendship among its participants, promoting the exchange of ideas and fruitful interactions. Traditionally, Franzosini Prize will be given in recognition of outstanding and sustained contributions to the field of critical evaluation of data in solubility and related chemical equilibria and memorializes the contributions to this field by the late Paolo Franzosini, Professor of Physical Chemistry at the University of Pavia. Also, the newly established Balarew Award will be awarded for the first time in the history of ISSP to an outstanding young scientist working in the field of critical evaluation of solubility and related chemical equilibria.

We are proud that ISSP21 is organized by the Faculty of Sciences University of Novi Sad in Serbia in the year when the Solubility Data Project celebrates 50 years, the Solubility Data Series, and the creation of the IUPAC Commission V.8 on Solubility Data 45 years, and 40 years since the first ISSP has been held in London, Ontario. City of Novi Sad, the second largest city in Serbia and the European capital of culture in 2021 has been selected for the conference venue. With rich cultural heritage, traditions, multinational diversity, six official languages, youthful vibrancy, and creativity, we are sure that Novi Sad will impress you and make this event memorable.

Let us come together to share our thoughts and shape the future in this field. Welcome to Serbia, welcome to Novi Sad.

Slobodan Gadžurić, Chair of ISSP21

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Conference partners







Sponsors

We would like to thank our sponsors for their support.





General Information

Venue

University of Novi Sad, Rectorate, dr Zorana Đinđića 1, Novi Sad Serbia

University of Novi Sad, Rectorate, address: dr Zorana Đinđića 1, Novi Sad Serbia. Our lectures are organized in the Amphitheatre on the ground floor of the building. The venue is within walking distance (about 15 minutes, 1 km) from the city center.





Registration

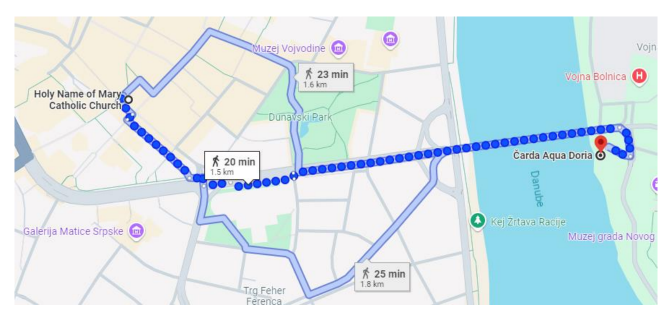
The registration desk will be open on Monday, September 9, 2024, from 8:00 am to 5:00 pm and every day during the conference from Tuesday to Friday from 9:00 am in front of the Amphitheater. Due to security reasons, participants must wear conference badges as their identification to enter the venue.

Conference tour Novi Sad city tour with wine tasting

The city tour is scheduled for Wednesday, September 11, 2024, in the afternoon for all participants. We will start our tour from the plateau in front of the Rectorate at 13:30 h directly after lunch. During the tour, we will have a tasting of the regional wines organized in the Museum of Vojvodina province. One of the wines will be a dessert wine that is a specialty of Fruška Gora wine region in northern Serbia. It was originally intended for medicinal purposes, but later it was produced for regular consumption. It is usually served as a dessert wine, but can also be served as an aperitif, much like Vermouth. However, *Bermet* is produced in a different manner than Vermouth, through maceration of 20 different herbs and spices. It can be made of red or white grapes, but the exact recipe is secret and held by only a handful of families in the town of Sremski Karlovci. According to some documents, *Bermet* was served on the Titanic, as well as in the Vienna royal palace as one of the most favorite wines of Austrian sovereign Maria Theresa. The end of the city tour is expected around 17h.

Conference dinner Aqua Doria restaurant, Kamenički put bb, 21000 Petrovaradin, Serbia

The conference dinner will be held on Thursday, September 12, 2024, at 20:00 in the traditional Aqua Doria Restaurant where you can enjoy traditional Serbian cuisine and music. The restaurant is located in one of the most attractive places in Novi Sad, at the foot of Petrovaradin Fortress on the bank of the Danube River (next to the bridge). You can take a taxi or just walk to the restaurant from the city center (20 min).



Topics

Aqueous Solutions

Biofuels

Computer Assisted Equilibrium Calculation

Deep Eutectic Solvents

Environmental Equilibrium Processes and Applications

Fluid Phase Equilibria

Molten Salts

Ionic Liquids

Nuclear Wastes

Solution Chemistry Complex Equilibria

Solubility Phenomena in Pharmaceutical Applications

Book of Abstracts

21st IUPAC International Symposium on Solubility Phenomena and Related Equilibrium Processes

PL1

Calorimetric and phase equilibrium data: from design of measurements methods to assessment of uncertainty and critical evaluation

Zdeněk Wagner

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Measurement is just one part of experimental scientific work in many branches including but not limited to physics and chemistry. A measured value is not much useful if its quality and reliability are not assessed. Data analysis is thus an important complement to experimental work. Mathematical analysis should often be employed even before the real measurement starts. The quality of the results of measurement techniques can be affected by settings of various parameters. For instance, the quality of the continuous measurement of heat capacity depends on the heating rate used. The optimum value should be selected by mathematical analysis of preliminary measurements.

Measurements can be affected by various types of errors. Each type of error requires its own methodology of analysis. We will limit our attention to measurement precision which is defined as closeness of agreement between indications of measured quantity values obtained by repeated measurements on the same or similar objects under specified conditions. It is also known as a repeatability as well as a random error because it behaves unpredictably but it is not necessarily a random variable in the statistical meaning of the word.

The final task is the comparison with literature values and critical evaluation. The aim is to find whether a subset of available values is possibly subject to a systematic error and after discarding them to provide a recommended value of the measured quantity. It is well known that experimenters sometimes assign to their measurements an uncertainty that is lower than reality. On the contrary, compilers of databases and other collections sometimes assign an uncertainty that is too high. Published values of uncertainty thus cannot be relied upon. The mathematical tools that are nowadays in use cannot reliably distinguish between poor precision (i.e. high scatter) and systematic deviation because there is no statistical model of systematic errors. A different approach is therefore needed.

The lecture will demonstrate the application of modern robust non-statistical methods of data analysis based on mathematical gnostics to all three interconnected parts of experimental work on real measurements of calorimetric and phase equilibrium data. Advantages, disadvantages, and limitations will be discussed.

PL2

Using the available pKa data in non-aqueous solvents

<u>Ivo Leito</u>, Ivari Kaljurand, Sofja Tshepelevitsh, Agnes Kütt, Märt Lõkov, Jaan Saame, Agnes Heering, Elisabeth Parman, Sigrid Selberg, Andre Leesment, Merili Tammiste, Helerin Roomet and Carmen Kesküla

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The acid and base strengths, typically expressed as pKa values, depend on the solvation of the proton, as well as of the neutral and ionized forms of the acid/base [1]. Every solvent has different solvation properties. Thus, the pKa values for the same acid/base in different solvents are also different (often dramatically different). In principle, whenever using pKa values for predicting or rationalizing chemical processes, the pKa values determined in the same solvent should be used. Two main problems arise:

- 1. In some solvents, e.g. water, DMSO or acetonitrile large bodies of pKa data exist, while in most solvents either very few pKa values are available or none at all.
- 2. Reliably measuring pKa values in non-aqueous solvents is difficult.

Thus, the pKa data in the literature is often unreliable. This presentation addresses both of these problems. The problem 1 can often be solved by estimating pKa values in one solvent from the data in other solvent(s) [2,3]. An overview to what extent such estimates can be usefully done, highlighting both successes and failures. Concerning the reliability of the non-aqueous pKa data a brief overview will be given of the IUPAC project 2015-020-2-500 "Critical compilation of acid pKa values in polar aprotic solvents" [4].

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PL3

Exploring the potential: design strategies and applications of deep eutectic solvents

Mirjana Minceva

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Deep eutectic solvents (DES) represent a promising class of designer solvents with diverse applications across various fields. DESs are eutectic mixtures of two or more compounds with a melting temperature significantly lower than that of the pure constituents. The DES physicochemical properties can be tuned by selecting its constituents. However, due to the large number of potential compounds that can be used to form DES, the design of these solvents is challenging. Therefore, understanding the DES formation is essential for selecting the constituents and defining their molar ratio to tune the solvent properties accordingly.

This talk will discuss approaches for designing DES and tailoring their compositions for specific applications. The importance of the solid-liquid equilibria (SLE) as a starting point in the DES design will be elaborated. Applications of DES in separation processes with a focus on extraction and chromatography, as well as current limitations and challenges, will be presented. The presentation aims to inspire further research and innovation in exploring DESs´ full potential as powerful solvents.

PL4 Thirty years of ionic liquids

Andrea Mele

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The history of ionic liquids (ILs) began with a chance discovery, the scientific implications of which were not understood at the time. Ethylammonium nitrate, prepared by Walden in 1914, was viewed as an academic oddity, a salt with a melting point so low that it remained liquid at room temperature. The world wasn't yet ready to grasp the exceptional potential of room-temperature liquid salts.

It took about 80 years to arrive at the first class of compounds -the alkylimidazolium salts- which we rightfully consider the forerunners of modern ionic liquids, organic salts with melting points below the ordinary boiling point of water.

The number of publications indexed with the keyword "ionic liquids" grew from 323 in 1992 to over 7,500 in 2022, reflecting the incredible success of this class of compounds.

In this presentation, we will trace the milestones in the study of ionic liquids through five pillars:

- Local structuring
- Gas absorption
- Interaction of ILs with biomacromolecules
- Use of IL with biomass
- Ionic liquids and energy storage devices

The central theme of the presentation is the close interconnectedness between the fundamental understanding of intermolecular interactions within ionic liquids and the innovative and diverse applications that characterize this class of compounds today.

PL5 Chemistry in art and art in chemistry

Clara Magalhães

School of Biological, Earth & Environmental Sciences, UNSW Sydney, NSW 2052, Australia LEAF, Linking Landscape Environment Agriculture and Food Research Centre, Associate Laboratory TERRA, Instituto Superior de Agronomia, University of Lisbon, Lisbon, Portugal

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Chemistry is the science of materials, their properties and transformations. Art is a human activity that uses a variety of languages combining color, space, image, movement, words, sound and volume. Combining Chemistry and Art the interested basic elements are color, image, space and volume. The interest lies in drawing, painting, photography, sculpture and architecture.

Chemistry appears in art, fundamentally in painting and drawing, long before its creation as a science, as many of the objects and methods are inherited from alchemy. Art has been used to disseminate the scientific knowledge. Nowadays, new substances, materials and structures can be used to built new artworks.

In today's world, with changes in the atmosphere, all of this heritage is subject to much more energetic solar radiation and intense climatic conditions. It is not only the outer atmosphere that has been changing but also the inner atmospheres. There are more and more visitors to exhibition sites (museums, caves, art galleries) with a strong impact on the indoor environment. How do these changes influence artworks?

Chemistry can help to restore damaged art pieces as well as in the rehabilitation of old buildings facing specific problems. Analytical techniques are very important to identify artworks, by studying the composition of the materials, their origins and also the artist techniques. Analytical techniques can help to determine the authenticity of an artwork and to identify forgeries.

Artists must know the materials they work with. Accidents can happen and should be minimized. Artists, like chemists, must know the safety procedures when using different materials.

IL1

Modeling the dependence of stability constants on medium and ionic strength. The pure water model

Clemente Bretti¹, Concetta De Stefano¹, Sofia Gama², Claudia Granata¹, Gabriele Lando¹, Salvatore Raccuia¹ and Demetrio Milea¹

Presenting author email: dmilea@unime.it

The dependence of stability constants (and activity coefficients) on medium, ionic strength and temperature can be modelled by several approaches. The most common (e.g., Davies, Bromley, Pitzer, Specific ion Interaction Theory – SIT) can be considered as evolutions of the (Extended) Debye-Hückel theory/equation. All models and theories have pros and cons, so that none can be considered better than others [1]. Between the '80s and '90s, the group of Prof. Silvio Sammartano from the University of Messina proposed, with the precious support of colleagues from the Italian Universities of Torino, Catania and Palermo, a model [2] (with the relative equation), for the ionic strength (and temperature) dependence of formation constants, based on three simple assumptions:

- [H1]: It is possible to express the dependence on ionic strength of formation constants by a simple equation, independently of the type of reactants and products, and only dependent on the type of reaction.
- [H2]: All the deviations from the predicted behavior are ascribed to weak complex formation between components and/or species under study and the background ions (e.g., the ionic medium). This implies that "pure water" is considered as reference state, and some ions as non-interacting with the reactants and/or products involved in the studied equilibrium.
- [H3]: Perchlorate does not interact with cationic species, tetraethylammonium cations (and higher tetraalkylammonium analogues) do not with O-donor ligands, and Na⁺ and K⁺ do not with N-donor ligands.

Evidences collected during more than half a century in those universities demonstrated the validity of this "pure water model", and showed the potential of this approach to model the speciation of several multicomponent complex systems in a very simple way. This contribution describes the main features of the pure water model through some examples, highlighting the theoretical and practical aspects of this approach in the speciation modeling of systems of different complexity, including real systems.

Acknowledgements: We acknowledge financial support under the National Recovery and Resilience Plan (NRRP), Mission 4, Component 2, Investment 1.1, Call for tender No. 1409 published on 14.9.2022 by the Italian Ministry of University and Research (MUR), funded by the European Union – NextGenerationEU–Project Title Efficient Sequestration of Metal Ions from Aqueous Systems for Green and Sustainable Applications - AquaGreen – CUP J53D23014430001- Grant Assignment Decree No. 1409 adopted on 14/09/2022 by the Italian Ministry of Ministry of University and Research (MUR). This contribution is dedicated to the memory of Prof. Silvio Sammartano, for the unquestionable impact he had in developing research in this and related fields.

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IL2

Thermodynamics of aqueous solutions of menthol-basedchiral ionic liquids from the macroscopic and molecularviewpoints

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Even though squeous solutions of organic and inorganic compounds, including of ionicliquids, are some of the most studied systems in physical chemistry, we still lack some understanding of the solute-solvent interactions of ionic liquid ions with water at the molecular level. In addition, such systems have an increasing practical relevance, as the properties of ionic liquids may be modified not only in terms of their structure, but also by adding suitable molecular solvents. Water seems to be one of the obvious choices, due to its availability and low toxicity. To better understand its interaction with ionic liquids, we propose an experimental and molecular dynamic simulation study of physicochemical properties and phase behaviour of these systems [1]. Experimental volumetric, osmometric, thermal and transport properties provide us with a fair amount of data on solution chemistry of aqueous solutions of ionic liquids in their own right. MD simulations then allow for an even deeper insight into the molecular structure of ILs in water, while estimates of bulk volumetric and transport properties by simulations can be validated by the corresponding experiments.

In this contribution, this approach will be illustrated by examples of studies of volumetric, transport and osmometric properties in aqueous solutions of conventional and chiral ionic liquids (e.g. in 1-alkyl-3-[(1R,2S,5R)-(-)-menthoxymethyl]imidazolium chlorides) by means of experiments and MD simulations. In addition to binary aqueous solutions, octanol-water partitioning of the studied ILs will be discussed.

Acknowledgements: Specific university research (A1_FCHI_2022_002), the Czech Science Foundation (grant 20- 24155S), MEYSe-INFRA CZ (ID:90140) and National Science Centre, Poland (Grant No. UMO-2020/37/B/NZ9/04201) are greatly acknowledged.

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Solute aggregates in solution and the crystallization of organic molecules

Ricardo Simões

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Crystallization from solution remains one of the most widely used, and yet poorly understood, methods for the preparation and purification of solid materials. Control over the crystallization process is often challenging, as even small variations in experimental conditions can lead to the production of different solid forms of a compound (polymorphism). As each polymorph displays unique physical properties (such as fusion temperature, colour, conductivity, or solubility), crystallization control is an important issue in many industries, with an impact on the reproducible manufacture, and safe use, of a product (e.g., variations in the solubility of pharmaceutical compounds changing their bioavailability).

It is generally assumed that crystal formation occurs through a mechanism of nucleation and growth. As the solution reaches a metastable state (supersaturation), small nuclei of the new phase begin to form, growing into macroscopic crystals by incorporating further solute molecules. Recent studies, however, have highlighted that stable aggregates of solute molecules are often already present in solution, even before reaching conditions of supersaturation. The role of these prenucleation clusters in the crystallization process is now being investigated [1].

The impact of solute aggregation on the precipitation of different solid forms is here explored for the crystallization of 4'-hydroxyacetophenone (HAP, Figure 1), from water. HAP is known to adopt two anhydrous polymorphs as well as three hydrated forms. When crystallized from water, the obtained solid form is highly sensitive to the experimental conditions. Indeed, two of the hydrates (H2, and H3) and the anhydrous form I can be selectively produced, through cooling crystallization of aqueous HAP solutions, simply by varying the initial concentration of the solute [2]. Variations in the solution structure, as a function of the temperature and composition, were assessed by combining volumetric and acoustic measurements with molecular dynamics simulations.

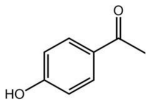


Figure 1. Molecular structure of 4'-hydroxyacetophenone.

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II.4

The intricate role of salts and sugars in biomolecular complexation

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The balance between attractive and repulsive interactions between protein and a synthetic polyelectrolyte (PE) in aqueous solutions determines the formation of protein-PE complexes. The strength of the interactions for a given protein-PE system and their modulation strongly depend on the PE charge density, the heterogeneous charge distribution of the protein surface, the pH and ionic strength of the solution, the presence of co-solutes, etc. and can lead to phase separation. Complexation is often initiated at pH values where both macromolecules have the same charge sign (so-called complexation on the "wrong side" of the isoionic point of the protein) [1]. Understanding how other components (such as salts, buffers, sugars, etc.) influence the onset of complexation is critical given its importance in numerous applications, including protein purification, drug delivery systems, biomaterials and biosensors.

We have investigated the modulating role of salts (NaCl, NaBr, NaI) and sugars (sucrose, sucralose) on complexation between a globular protein, bovine serum albumin (BSA), and a synthetic polyelectrolyte, sodium polystyrene sulfonate (NaPSS), at pH values above the isoionic point of BSA using a combination of experimental techniques and molecular modelling. The electrostatic attraction between positively charged patches on the BSA surface and the overall negatively charged NaPSS is responsible for the onset of BSA-NaPSS complex formation. Salt ions screen attractive electrostatic interactions and hinder complex formation in a concentration-dependent manner. Ion-specific effects have been observed (NaI > NaBr > NaCl) [1]. The effect of sugars on complex formation is less pronounced: sucrose showed no significant effect, while its chlorinated analogue, sucralose, prevented phase separation. The differences in the effect of these two sugars are due to their water-structuring capabilities [2,3]. Consistent with the so-called *preferential exclusion mechanism*, sucrose does not interact directly with the protein surface, whereas sucralose adheres to the BSA surface (*preferential interaction mechanism*) and can hinder contact between BSA and NaPSS.

Acknowledgements: financial support from the ARIS is acknowledged (research core funding no. P1-0201)

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IL5

Insight into the impact of ionic liquids on forming aqueous biphasic systems

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Aqueous biphasic systems (ABS) are formed when two water-soluble components are mixed in water at specific concentrations. Due to the negligible volatilities of the phase-forming constituents and the high water content, ABS are environmentally friendly and biocompatible alternatives to traditional liquid-liquid extraction methods. Rogers et al. [1] introduced ionic liquids (ILs) into the formation of ABS, spurring extensive research into IL-based ABS. The inherent advantages of ABS can be combined with the biocompatibility and favorable characteristics of ILs, including their tunability and wide solubility for both polar and non-polar compounds. Various combinations of ILs with inorganic and organic salts, alcohols, carbohydrates, and polymers have been reported to form ABS. The properties of ILs, such as density, viscosity, and surface tension, significantly influence the phase behavior in ABS. The potential of IL-ABS for the extraction, separation, and enrichment of diverse classes of compounds, including biomolecules, metal ions, pharmaceuticals, and dyes, has been extensively explored. Three major types of IL-based ABS have been extensively studied: IL-inorganic salt ABS, IL- carbohydrate ABS, and IL-polymer ABS. The formation and stability of IL-based ABS are influenced by the structure of the ILs, temperature, pH, and the nature and functionality of the other phaseforming components such as the charge density of the salts, the structure of the carbohydrates and the molar mass and functionality of the polymers. Firstly, the impact of the IL cation structure (such as alkyl chain length and substituents at the N-1 and N-3 positions of the imidazolium cation) and anion on forming ABS with inorganic salts was investigated. It was found that the ability to form ABS increases with the alkyl chain length on the imidazolium cation. Additionally, ILs with an ethyl group at the N-3 position form ABS more effectively compared to those with a methyl substituent, attributed to the increased hydrophobicity of the ILs and their lower affinity for water. Secondly, the formation of ABS based on symmetric imidazolium ILs, such as 1,3dibutylimidazolium and 1,3-diethylimidazolium combined with phosphate and citrate salts, was examined. It was found that the ABS based on 1,3-dibutylimidazolium dicyanamide exhibited an exceptionally large biphasic area, requiring avery low amount of salt (~2%) to induce phase separation. Thirdly, the effect of naturally derived anions (such as acetate, lactate, pyruvate, taurate, and nicotinate) in choline-based ILs on forming ABS with polymers was studied. The salting-out aptitude of these ILs is directly related to the hydration capacity of their ions. Since all the ILs share a common cation, choline, more hydrophilic anions with higher charge density, like acetate, exhibit greater hydration ability and stronger salting-out effects. To better understand the impact of ILs on ABS formation efficiency, computer simulations were conducted.

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Ionic liquids in organic synthesis: mechanistic aspects in Knoevenagel reactions

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Knoevenagel reaction, also known as Knoevenagel condensation, is a classical transformation between a carbonyl compound (aldehydes or ketones) and an activated methylene compound, such as a 1,3-dicarbonyl compound, with ammonia or other amines as catalysts in organic solvents, to obtain the Knoevenagel condensation product, a dicarbonyl α,β -unsaturated compound [1]. In spite of your 128 years old, this reaction remains a powerful tool to form new C-C bonds, employed as key-step in many synthetic strategies to afford complex structural compounds in linear or multicomponent processes [2-5].

In recent years, ionic liquids (ILs) have emerged as a greener alternative to commonly used organic solvents and catalysts. Their green characteristics, combined with their low volatility, non-inflammable nature, thermal stability, and capacity for reuse as catalysts, have made ILs an environmentally friendly option for organic synthesis [6]. However, a few studies have reported the use of ILs for both their solvent and catalytic functions; ILs have been used most commonly as solvents in the presence of another catalyst/promoter system or as catalysts in the presence of a solvent.

In this work, the use of Pyrrolidinium ionic liquids **3-5** (PyrrILS) as solvent and catalyst in the Knoenevagel reactions is described, using several classic 1,3-dicarbonyl compounds **2** (β -ketoesters, 1,3-diketones, Meldrum's acid, and malonates) with aliphatic aldehydes **1**, in a solvent-free media (Figure 1). Besides, the mechanistic aspects are discussed, supported by NMR experiments.

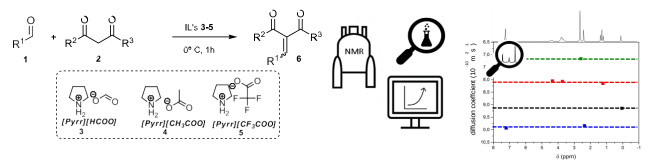


Figure 1. Mechanistic studies in Knoevenagel reaction and Ionic Liquids.

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IL7 Study of oxides solubility in fluoride molten salts

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Research on molten salts, especially fluoride systems, has experienced a significant renaissance in last decade. This focus is mainly on special applications that have the potential to replace old ones, while obtaining various benefits such as environmental, economic and, of course, functional. From a relatively wide range we can highlight in particular:

- use in metallurgy (e.g. solvent of electroactive component in electrolytic production of metals)
- use in energy applications: i) in Generation IV reactors; (ii) in the field of solar energy storage
- use in other areas: e.g. i) preparation of single crystals with desired physical properties (piezoelectrics, optical transparency, ion conductors ...), ii) preparation of substances serving as standards for various analytical methods, iii) preparation of substances with defined spectral properties (e.g. photoluminescence).

The recycling seems to be the efficient way how to reduce the dependence on imports of raw materials, reduce the volume of waste and how to increase the sustainable development. Molten salt electrolysis is an industrial method for metals or alloys production from corresponding mixtures, including oxides. It is known that the solubility of oxides is not high but the presence of fluorides increase the solubility, especially in the case of rare earth elements.

We have mainly used thermal analysis method and XRD analysis for investigation of oxides solubility. Oxides solubility differ from used fluoride molten salts.

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11.8

Drug solubility enhancement: from buffer complexes formation to acid-base supersolubilization

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A majority of the new chemical entities (NCE) that emerged as potential drug candidates in pharmaceutical development during the past 2-3 decades are practically insoluble solids consisting of ionizable molecules [1]. Class II BCS drugs (Biopharmaceutics Classification System) exhibit poor bioavailability due to insufficient absorption in the gastrointestinal tract with slow drug release from the dosage forms and low solubility as the limiting steps for their absorption [2]. Thus, detailed and precise study of compound solubility and the possibilities to increase solubility and dissolution rate, are at the core of the development of bioavailable drug dosage forms and clinically effective pharmaceutical products that would dissolve in gastric and intestinal fluids after oral administration or not precipitate in the blood after intravenous administration. A white paper on consensus recommendations for improving data quality in equilibrium solubility measurement of ionizable drugs [3] emphasizes the importance of precise solubility measurements. As a part of solubility studies of a group of tricyclic antidepressants (TCAs) we have shown the influence of competing counterions, such as buffering agents, complexing agents, salt coformers, tonicity adjusters, and solid-phase transformations on the aqueous solubility of studied drugs [4-5].

A variety of methods to increase solubility and/or dissolution rate, and thereby increase their bioavailability, have been developed. Still, most of them, like particle size reduction, salt formation, conversion to amorphous form, solid dispersion, and solubilization in lipids or lipid-surfactant mixtures have their own limitations. To mitigate some of the above limitations, a novel method of drug solubilization in aqueous media by acid—base interactions has been developed [6]. This novel approach of greatly increasing the solubility is based on interactions of a model low-soluble basic drug in an aqueous medium with acidic species that would not normally form salts with it. Although quite successful, the proposed model still needs additional work and some fine-tuning with additional low-soluble drugs to establish it as a widely accepted method for increasing solubility, dissolution rate and bioavailability of poorly water-soluble drugs. Our research team is working on it.

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OL1 My adventures with the solubility data

On the occasion of the Franzosini Prize

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The present speaker is co-author of 10 volumes [1-10] published within the IUPAC-NIST Solubility Data Series. He has been a four dacades member of the Solubility Data Commission and later the Subcommittee for Solubility and Equilibrium Data.

Preparation of the volumes for this series is really a time consuming process that needs a very systematic activity. Primary materials must be completed as much as possible. Some subjects need inspection of such rare sources as U.S. Atomic Energy Commission Reports, conference materials, rare journals and reports from Russia and China (not in English). After preparation of data sheets for every experimental study, critical evaluations of selected systems are prepared. Finally, authors of the critical evaluations select the best or recommended solubility data set to be further used by scientific as well as technical community. Therefore, the solubility volumes should not be treated as a kind of reviews where authors are not obliged for detailed assessment and a selection of recommended values.

This author was involved in elaboration of two kinds of systems: where liquid mercury or alkali metals (at elevated temperatures) were the solvents and where rare earth metal salts were dissolved in various aqueous media. Critical evaluations for a part of them were complicated by significant scatter of the solubility results. The most frequent reasons of the divergences were: purity of components of a systems being investigated, slow kinetics of an real equilibrium approach and side chemical reactions connected with solubility process.

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OL₂

Enhancing acetaminophen solubility in aqueous solutions with hydrotropes

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Hydrotropes are compounds capable of enhancing the solubility of sparingly soluble molecules in aqueous media, thereby expanding their effective use in many applications [1]. Typically, these compounds exhibit amphiphilic properties, characterized by both hydrophilic and hydrophobic domains within their molecular structure. And while an increase in solubility indicates hydrotrope efficiency, it also holds significance for water activity. In fact, the addition of hydrotropes is known to alter the free water within the system, potentially impacting its activity in pharmaceutical formulations [2,3]. Within the pharmaceutical industry, several traditional hydrotropes are commonly used to enhance the solubility of poorly water-soluble drugs, namely sodium-based salts like benzoate or salicylate, short alkylbenzene sulfonates, and selected small organic compounds such as urea or nicotinamide [2].

Acetaminophen (ACP), a widely used analgesic recognized as an essential medicine by the World Health Organization, remains surprisingly understudied in the hydrotropy field. Previous works have explored the use of organic solvents, biomolecules, salts, ionic liquids, and eutectic mixtures. Due to its significance as a model compound, ACP has been selected as the subject of study to explore different types of hydrotropes under similar conditions, aiming to establish connections between hydrotropes molecular structure and the increase in solubility.

The main goal of this study is to investigate the effect of several hydrotrope families (traditional organic compounds, ionic liquids, and organic or inorganic salts) at different concentrations on the solubility of ACP. New solubility data in aqueous solutions of sixteen hydrotropes were experimentally measured at 298.2 K using the shake-flask method coupled to UV-Vis spectroscopy or gravimetry as analytical techniques. The COSMO-RS model was used to analyze the hydrotrope effect on the solubility increase, through the determination of the hydrotrope charge density (the donor *DF*, apolar *AF*, and acceptor *AcF* factors). Experimental water activity data at 298.15 K for some specific organic salts are also included to better understand if the hydrotrope-induced depression of water activity is related to the solubility increase. Overall, the findings showed that among the tested hydrotropes, organic compounds (acetone or ethanol) had the highest potential for increasing solubility, followed by salts and ionic liquids. In general, according to COSMO-RS, compounds with higher *AF* values tend to increase solubility more effectively. Besides that, in the aqueous solutions of salts that promoted the highest ACP solubility increase, the highest water activity was observed. These findings provide a deeper understanding of the critical role of hydrotropes in improving drug solubility, with significant implications in the development of pharmaceutical formulations.

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Revealing the solubility enhancement of active pharmaceutical ingredients through eutectic mixtures formation: A parameter study

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The poor water solubility of active pharmaceutical ingredients (APIs) poses a significant challenge in pharmaceutical development, impacting their bioavailability and therapeutic efficacy. Enhancing API solubility in solvents through the formation of eutectic mixtures with excipients is a feasible approach. However, despite the experimental simplicity of this method, its exploration from a thermodynamic perspective remains relatively limited.

This study explores the influence of parameters on enhancing the solubility of an API with poor solubility in water through eutectic mixture formation. A hypothetical ternary system comprising an API, excipient, and water is considered. The investigation focuses on the impact of excipient melting properties and intermolecular interactions within the liquid solution on API solubility. The liquid phase nonideality is modeled using the two-suffix Margules equation.

The study finds that the eutectic point in a ternary phase diagram consistently represents the maximum solubility enhancement factor for the API. The position of the eutectic point is influenced by the melting properties of the compounds and their molecular interactions. Additionally, the investigation reveals that excipients with low melting temperatures and small melting enthalpies can increase the solubility enhancement factor of the API in water and affect the composition of the constituents at the eutectic point.

Stronger API-excipient interactions, compared to API-water interactions, significantly improve API solubility in the liquid solution. Conversely, if excipient-water interactions are favored over API-excipient interactions, the solubility enhancement of the API in the solution decreases.

The optimal scenario for enhancing API solubility involves selecting an excipient that strongly interacts with the API while maintaining moderate interactions with the solvent.

The current study could serve as a valuable tool in advancing pharmaceutical formulation science by aiding the design of effective drug delivery systems and the development of novel therapeutics with improved solubility and bioavailability.

OL₄

Determination and modelling of solubilities in aqueous vanadate solutions

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Aqueous solutions of vanadium (V) salts are in the center of many trending research topics such as hydrometallurgical vanadium recovery processes [1] and vanadium redox-flow batteries [2]. Despite this, physically sound models even for the simplest of binary vanadate salt – water systems that could describe their thermodynamic properties with reasonable accuracy have not appeared in the literature. It can be argued that currently the most important factor hindering the development of such models is the lack of experimental data which is needed in model fitting and validation.

Currently, the thermodynamic information available for aqueous V(V) systems is mainly limited to equilibrium constant, density, and solubility data [3-4]. However, in the case of NaVO₃ there seems to be some ambiguities in the solid phase for which solubility data is reported. Furthermore, many publications do not report sufficient details to allow reliable estimation of uncertainties in the reported solubilities. This is a common issue and there is currently an IUPAC project [5] ongoing with the aim of providing investigators working with solubility determinations with guidelines for good practices in uncertainty assessment.

A detailed investigation of solubility in the binary systems $NaVO_3$ – H_2O and NH_4VO_3 – H_2O was performed by conducting new experimental measurements and a review of previous literature [6]. The guidelines discussed by Hibbert [7] were followed in an attempt to provide reasonable uncertainty estimates. The capability to predict the solubility of the sparingly soluble NH_4VO_3 at 298.15 K based on presently available thermodynamic data was also briefly investigated. Although the magnitude of the solubility could be correctly predicted, considerable deficiencies in the models and the lack of data needed to improve them were recognized. The present work is part of the PhD studies of the first author focusing on the determination and modelling of thermodynamic properties of aqueous V(V) systems.

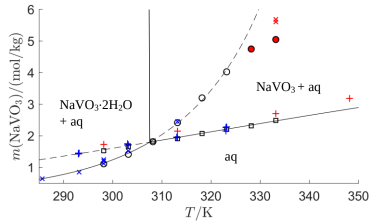


Figure 1. Solubility of NaVO₃ in water. Circles and squares: [5], Crosses: literature data, Red: rejected data, Curves: fitted correlation equations.

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OL5 Having fun with sulfate complexes

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A critical evaluation of the experimental data in the literature for the stability constants and related thermodynamic parameters of metal-ion/sulfate complexes is being conducted under IUPAC auspices. Such quantities are important for modelling natural and industrial water systems as sulfate is a significant constituent of ocean, estuarine, fresh, and waste waters. Metal-ion/sulfate complexes are also found in biological fluids such as blood plasma, in geochemical liquids and atmospheric aerosols, and are present in vast quantities in extractive metallurgy.

Because of its importance, this review began with the protonation of sulfate:

$$H^{+}(aq) + SO_4^{2-}(aq)$$
 $HSO_4^{-}(aq)$

Approximately 400 references containing quantitative information about this equilibrium were obtained from the literature, including databases and technical reports. About 60% were rejected outright for various reasons, such as inadequate specification of the measurement conditions, inconsistency with other studies, or because they didn't contain original experimental data. The values so obtained for the stability constants, enthalpies, entropies, and heat capacities for the protonation equilibrium will be discussed. Areas where the database needs further experimental investigation will be identified.

Current work in this project is focused on the *s*-block cations of Groups I and II of the Periodic Table. The database for the sulfate complexes of these ions varies enormously, with some systems (eg, Rb⁺, Ba²⁺) hardly studied while others (eg, Na⁺, Mg²⁺) have been comprehensively investigated. Progress on the evaluation of these apparently 'simple', yet diverse, systems will be presented.

T-dependent Pitzer model for carbonates and CO₂ in the THEREDA-database for aqueous geochemistry up to high ionic strengths

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The THEREDA-Database [1] represents a thermodynamic database providing ready-to-use downloads for widely used geochemical codes as PHREEQC and GWB. A particular feature of this database represents the Pitzer model of the oceanic salt system H^+ , Na^+ , K^+ , Mg^{++} , Ca^{++} // Cl^- , SO_4^{--} , OH^- - H_2O , which enables one to calculate the solubility equilibria in multicomponent solutions for a temperature range of $0-100\,^{\circ}C$. Recently, solubility constants of carbonates of sodium, potassium, magnesium and calcium have been implemented as well as the Pitzer interaction coefficients of CO_3^{2-} and HCO_3^- with all the other ions [2]. With the extended database the solubility of CO_2 and its interaction with the salt components can be calculated up to moderate pressures (see Figure 1).

In the contribution the data situation of the carbonates in respect to solubility at varying temperature, pH and partial pressure of CO₂ is discussed. At selected examples the description of experimental data by the model is demonstrated. Thereby, the uncertainty of experimentally determined solubilities of important minerals as magnesite, hydromagnesite or dolomite is discussed. The literature study revealed also that solubility data of these minerals in natural brines are nearly absent.

The incorporation of the solid-solution model of Königsberger and Gamsjäger [3] for magnesian calcites in a modified form allows the calculation of the equilibrium uptake of magnesium by calcium carbonate in magnesium salt containing solutions.

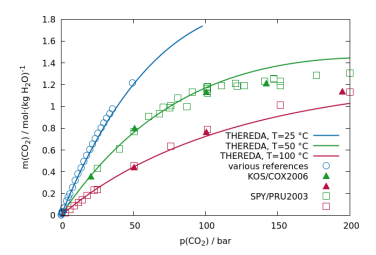


Figure 1. Solubility of CO₂ in water; comparison model (lines) and experimental data (symbols).

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The conversion of lithium carbonate to lithium hydroxide

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Lithium carbonate, Li₂CO₃, is an important precursor for the preparation of mixed-oxide cathodes in the production of Li-ion batteries [1]. Li₂CO₃ is then converted to the more precious lithium hydroxide, LiOH, also used for batteries, as a CO₂ sequestrant, and as starting material for various lithium salts. Conventionally, LiOH is produced by reacting the carbonate salt with Ca(OH)₂, which generates the sparingly soluble CaCO₃ [1,2]. Despite slaked lime being readily available, it needs to be of very high purity and more importantly, the overall process yields low conversions when using concentrated Li₂CO₃/Ca(OH)₂ mixtures, with a maximum attainable alkali concentration of 3.5 wt% [1–3]. However, the reason for low conversions has remained elusive; some suggested that is due to decreased solubility of either Li₂CO₃ or Ca(OH)₂ [1,3], while others observed the formation of an intermediate solid phase [2]. To shed light on the underlying mechanism of this reaction, we studied its time-dependence and equilibrium properties, analysing both the liquid and solid phases.

We performed all experiments at (30 ± 1) °C. First, conductometric measurements showed that most of the initial amount of Li₂CO₃ transforms to LiOH after 2 hours of contact time (using 10 g L⁻¹ Li⁺ and an equivalent concentration of Ca(OH)₂). Second, we performed conversion experiments varying both the amount of Li₂CO₃ (5–30 g L⁻¹ Li⁺) and the Ca(OH)₂:Li₂CO₃ molar ratio (25–200%), applying 2 hours of reaction time. We found the conversion ratios to be \leq 96% for 5 and 10 g L⁻¹ Li⁺, indicating that the process is essentially quantitative. However, for more concentrated suspensions, we indeed observed a dramatic decrease in yields, corresponding to an upper limiting concentration of LiOH of ~1.5 mol dm⁻³ or 3.5 wt%. For suspensions of 30 g L⁻¹ Li⁺, both sequential addition of the reactants and the order of their addition result in the same final concentration, suggesting that this is an equilibrium process and that heterogeneous reactions obey Le Chatelier's principle, similarly to liquid-phase reactions. Furthermore, no solid phase other than the starting materials and CaCO₃ is discernible on the powder diffractograms, supporting the equilibrium nature of the reaction and hence there is no need to assume intermediate phases.

In addition, concentration of CO_3^{2-} in the supernatants were unexpectedly high which on the one hand can be explained by the solubility of Li_2CO_3 . However, given the very low solubility of $CaCO_3$, most of the soluble CO_3^{2-} exist probably as $LiCO_3^0$ ion-pair.

In the second set of experiments, we investigated the backward reaction, starting from calcite and concentrated LiOH of 4.2 mol dm⁻³ (which otherwise could not be prepared from Li₂CO₃ and Ca(OH)₂). As expected from a true thermodynamic equilibrium, we now obtained Li₂CO₃ and Ca(OH)₂ as products. Strikingly, the final LiOH concentration was ~2.2 mol dm⁻³, contradicting the one obtained from the forward process (1.5 mol dm⁻³). However, when starting from freshly precipitated CaCO₃, we again obtained a LiOH solution of 1.5 mol dm⁻³. Since this solid contained vaterite to a significant degree, this apparent contradiction may be resolved by assuming that in situ forming CaCO₃ always contains some vaterite as opposed to phase-pure calcite. The corresponding equilibrium constant, consisting of the solubility products of Li₂CO₃, Ca(OH)₂ and CaCO₃, will then be different, depending on which polymorph of CaCO₃ is considered to govern the equilibrium. In turn, assuming different polymorphs will give rise to different equilibrium concentrations of LiOH.

In conclusion, all experimental results suggest that the low conversion for concentrated suspensions stem from the decreased solubility of the Li_2CO_3 and $\text{Ca}(\text{OH})_2$ in concentrated LiOH solution, and this can be quantified by an equilibrium constant.

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Thermodynamic modelling of CuSO₄-H₂SO₄-H₂O system from 273.15 K to 353.15 K

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Modelling the solubility of copper sulfate in aqueous sulfuric acid is important for the development and optimization in hydrometallurgical processes for copper electrowinning and recycling.

A thermodynamic model for CuSO₄-H₂SO₄-H₂O system was developed covering temperature range from 273.15 to 353.15 K and concentration range up to ten molal sulfuric acid solutions. Pitzer model was used to describe the non-ideal behaviour of aqueous solution. The parameters for binary systems were obtained from literature. During the assessment the available solubility data of copper sulfate in sulfuric acid as well as activity of water data were critically evaluated.

Several Pitzer parameter sets for the system was also critically evaluated. It was found that following Pitzer parameters for the system $CuSO_4$ - H_2SO_4 - H_2O is required for Cu^{2+}/HSO_4 - interaction: $\beta(0)$, $\beta(1)$ and C^{φ} as well as ternary interaction ψ for $Cu^{2+}/H^+/SO_4^{2-}$. Total number of temperature depend terms in parameters were 3, 2, 2 and 2, respectively, from the following set of terms: a, bT and f/T. Calculated solubility versus experimental data at 298.15 K is shown in Figure 1.

The model was verified against electrochemical cell data and solubility of trihydrate.

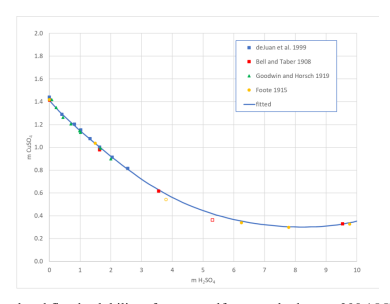


Figure 1. Experimental and fitted solubility of copper sulfate pentahydrate at 298.15 K. Open symbols was not included in the assessment.

Deliquescence of salt mixture: from water film to bulk liquid

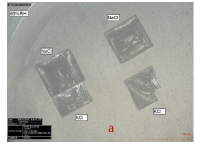
Shaoheng Wang¹, Sebastiaan Godts² and Michael Steiger¹

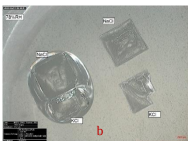
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Deliquescence is a first-order phase transition from solid to solution at a certain relative humidity (DRH), and it widely influences the process of aerosol formation, heritage conservation, the Martian water cycle, etc. [1-3]. In the mixture, the solid-solution transition humidity would be lower than each component, which impacts the physical stability of the salt mixture system. The dynamic mechanism that water sorption on salt surfaces has attracted a lot of attention. Many studies focused on predeliquescence of single salt have been reported [4-6], and raise the mechanism that there is nanoscale thickness water film forming before deliquescence, which leads to the salt's full dissolution at DRH. However, little understanding is gained of how the deliquescence starts in the salt mixture at a humidity that is lower than the DRH of each component.

In this work, we compare the water sorption curves of single salt NaCl, KCl, and the mixture NaCl + KCl. It is indicated that the deliquescence humidity of the mixture is lower than single salt. Subsequently, *insitu* observation of the water vapor sorption was performed in an optical microscope that cooperated with the environment chamber, as seen in Fig. 1a, 1b. We found the water condensation starts at the interface of salt particles, then the dissolution spreads to bulk salts at deliquescence humidity. We propose that the deliquescence initially takes place in the nanoscale hole generated by the rough surface of particles, where the water condensation pressure is much lower than the bulk system [7]. Then the solution film from each side, i.e. NaCl and KCl, converge in the interfacial gap, shown in Fig. 1c. This confluence is a crucial step for mixture deliquescence since it produces the saturated solution containing both salts, where the water activity or vapor pressure theoretically is lower than the single salt. When the environment humidity reaches the vapor pressure of the saturated solution, this interfacial reaction would extend to the whole system, until one of the components is depleted.





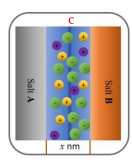


Figure 1. Water vapor sorption of single salts and their mixture.

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OL₁₀

Calcium phosphate precipitation in zwitterionic polymer solutions

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Zwitterionic polymers have found a wide range of applications due to their unique properties, which include high water solubility, biocompatibility, and resistance to fouling and protein adsorption. They can be incorporated into dental adhesives and restorative materials to improve their adhesion to tooth structure, preventing water infiltration that can lead to restoration failure over time.

The biomimetic precipitation of calcium phosphates in a physiological medium containing either polycarboxybetaine (PCB) or polysulfobetaine (PSB) was investigated in this study. PCB and PSB contain both positive and negative-charged functional groups within their structure, leading to interactions between the polymer and calcium and phosphate ions in the solution. Furthermore, the precipitation of calcium phosphate can be influenced by various factors, including pH, temperature, concentration of calcium and phosphate ions, etc. Several synthesis routes, differing in the mixing sequence of the starting solutions, amount of polymer, and pH were studied to elucidate their effect on the type of the pre-nucleation clusters and thus on the solid phases formed. The changes in pH and the concentration of Ca²⁺ ions during the syntheses were followed in polymer-free systems, in systems with PSB, and in systems with PCB to explore the effect of betaine zwitterionic functionality. Solid state NMR spectroscopy was used to identify the calcium phosphate phases formed as a function of the different synthesis conditions.

The results show that when the pH is not kept constant, and Ca-solution was added to the P-solution, HPO_4^{2-} ions dominate at the beginning, and the formation of $Ca(HPO_4)_3^{4-}$ pre-nucleation clusters is expected to prevail over other ones. The starting pH is 8.9, which results in the formation of amorphous calcium phosphate with common formula $Ca_x(HPO_4)_y(PO_4)_z$ and Ca/P = 1.1-1.6. In the second case (P-solution was added to Ca-solution), Ca^{2+} ions prevail in the initial solution. pH was 4.5 for the system with PSB, 5.7 for the system with PCB, and 6.3 for the polymer-free system. Thus, it was more probable that $CaHPO_4^{0}$ prenucleation clusters which are structural units of dicalcium phosphate dehydrate (DCPD) to be dominant. As a result, a mixture of octacalcium phosphate (OCP) and dicalcium phosphate dihydrate (DCPD) in different ratios was identified depending on the sequence of initial solution mixing and the type of negative functional group of the polymer used. The more acidic character of the sulfo group in PSB than the carboxy one in PCB determines the dominance of acidic solid phases, namely the acidic amorphous phase or DCPD. In the presence of PCB, the formation of ACP with acicular particles arranged in bundles with the same orientation was observed.

When the pH is kept constant, it is a dominant factor in the solid phase formation. Regardless of the amount of polymer and the method of reagent mixing the nano-crystalline hydroxyapatite with a Ca/P ratio of 1.63-1.71 was observed as a mineral phase in all systems.

The polymer type affects also the final microstructure and mineral particle size being thinner and smaller in the synthesis performed using PCB than using PSB. The final intermolecular interaction of the calcium phosphates was demonstrated to be stronger with PCB than with PSB.

The preliminary comparative study of the remineralization potential of the different prepared hybrid materials gives preference to that consisting of PCB.

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Terpene-based green solvents for the dissolution of plastics

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Plastics have become an integral part of our daily lives, offering cost-effective solutions that enhance food preservation, medical advancements, and other benefits. However, this convenience has led to an overreliance on these materials, contributing to a global crisis of plastic accumulation.

Despite the alarming production rate, which approaches 400 million tons per year, only 10% of plastics are currently recycled. Hence, there is an urgent need to develop more efficient and accessible recycling processes while ensuring the quality of recycled polymers.

Among the recycling processes, dissolution/precipitation stands out as a particularly promising option. Traditional organic solvents have been the most common choice for this method, but the quest for new solvents capable of efficiently dissolving and depolymerizing plastics under mild conditions and with high selectivity for valuable products has gained attention.

Eutectic solvents are promising alternatives in this endeavour. These are mixtures of pure compounds characterized by significant melting point depressions compared to those of the pure constituents. In the plastic dissolution field, natural terpene-based eutectic solvents have recently gained significant attention due to their dissolution potential.

Terpenes are a well-known class of natural compounds with important applications in various fields. Among them, limonene is a well-known solvent for different applications, and it has been reported as an attractive alternative for polystyrene solubilisation. Also, turpentine, a mixture of monoterpenes derived from the pine industry, has been used as a solvent for polyethylene and polypropylene.

This project aims to optimize dissolution/precipitation methods using environmentally friendly terpene-based biosolvents (and their mixtures) derived from industrial by-products like turpentine. Predictive tools such as the COnductor-like Screening MOdel for Realistic Solvents (COSMO-RS) and the Hansen solubility parameters are first used to identify the most promising systems, based on the chemical interactions between different polymers and solvents. Then, experimental solubility studies are carried out using an isothermal analytical methodology.

The findings gathered here are important contributions to revolutionizing plastics recycling, significantly reducing environmental pollution and fostering a more sustainable future.

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OL₁₂

Intermolecular interactions and sustainability: a pathway to exploring hydrophobic eutectics and eutectogels

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Hydrophobic non-ionic eutectic solvents (HES) have recently garnered attention amidst the array of (deep) eutectic solvents, (D)ES [1], thanks to their low viscosities, chloride-free nature, and water-immiscibility. A unique option to expand the design space of HES involves their immobilization within a jelly matrix to yield hydrophobic non-ionic eutectogels (HEG) [2].

While significant strides have been made in comprehending intermolecular interactions and mobility within conventional hydrophilic (D)ES, very little is known for HES, and even less so for HEG.

Here we demonstrate the pivotal role of a multifaceted investigation at the molecular level for unraveling the intricate interplay among H-bond, dispersive forces and steric factors, which drives the formation of HES and rules their peculiar properties [3].

A screening and ranking tool is proposed, which combines spectral descriptors of the intermolecular network that are easy to determine and of effective practical use, along with safety descriptors based on the hazards of the individual components.

We also show that non-ionic HES can be incorporated in stable and homogeneous supramolecular gel scaffolds using the low molecular weight gelator 1,3:2,4-dibenzylidene-D-sorbitol (DBS) at minimal loading, with a simple one-pot preparation process ranked as "excellent" in the EcoScale metrics [4]. A multidisciplinary approach is applied again to shed light on the structure and dynamics of both liquid and semi-solid systems, proving that the liquid-like nature of the hydrophobic mixture is retained upon gelation. Noteworthy, we observe an unexpected increase of the translational motion of the HES components under confinement compared to pure HES, with the effect of the gelator inversely proportional to the strength of pre-existing interactions.



Figure 1. Sketched representation of the combined approach based on molecular properties and safety aspects for screening, ranking and investigating hydrophobic eutectic solvents (HES) and eutectogels (HEG).

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The role of atomistic calculations in modeling solvent effects

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The rapid advancement of computational technologies has fundamentally transformed the landscape of molecular and material discovery, making these processes more accessible and efficient than ever before. Just over a decade ago, researchers were limited by the computational capabilities of standard desktop computers, which restricted them to relatively simple calculations. More complex tasks, such as quantum-mechanical modeling, require specialized hardware, often associated with high costs and limited availability. Today, the power of modern desktop computers, combined with significant improvements in modeling software, allows researchers to perform a range of calculations on molecules of various sizes and complexities.

Incorporating solvent effects into these calculations is essential for accurately modeling molecular interactions, as solvents play a critical role in influencing the behavior, stability, and reactivity of molecules. By accounting for solvent environments, researchers can achieve a more realistic representation of molecular systems. This inclusion is particularly vital in applications such as pharmaceutical design (Figure 1), where solvent interactions can significantly impact the efficacy and binding of pharmaceutical compounds, as well as in materials science, where solvent effects can alter the properties and performance of new materials.

Investigation of solvent effects at the molecular level is now possible thanks to the emergence of state-of-the-art codes for atomistic calculations, such as ORCA [1] and xtb [2]. However, using a command prompt can be challenging for beginners or users with an experimental focus. In such cases, the graphical user interfaces provided by the Atomistica.online project (available at https://atomistica.online [3]) can be helpful in bridging the gap. These tools enhance our understanding of molecular behavior in realistic environments and provide accessible resources for researchers.

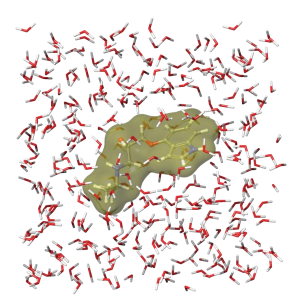


Figure 1. Molecular dynamics simulation of a pharmaceutical molecule solvated in water

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Recovery of pure lithium from salt lake brines at low temperatures: Parametrization of thermodynamic models and proposal of a novel extraction method

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One of the main challenges in the recovery of lithium from salt lake brines lies within the separation of lithium and magnesium salts. A thorough literature study revealed the potential of fractional crystallization at low temperatures as an effective method for the separation of magnesium salts.

Basic processes in the salar chemistry can be described within the quaternary reciprocal system $2Li^+$, $Mg^{2+} \parallel 2Cl^-$, SO_4^{2-} - H_2O . In order to estimate capabilities of the low temperature approach and to design respective processes, the knowledge of solubility equilibria in this system at temperatures below 25°C is essential.

The goal of the present work was to construct the phase diagram and describe equilibria down to about -30°C. For this purpose a PITZER model [1] has been parametrized in such a way that linear dependencies of the parameter allow for a reliable temperature extrapolation.

In the sulfate-free chloridic sub-system, LiCl-MgCl₂-H₂O, low water activities allow for the additional parametrization of a modified BET model according to Ally and Braunstein [2]. Though based on a completely different calculation approach, the BET model was able to validate the results of the Pitzer model. Also experimental solubility data and cooling experiments of salt brines confirm the model results.

Based on the model predictions, a process for direct crystallization of pure lithium chloride from highly concentrated lithium chloride - magnesium chloride mixed solutions between +9 °C and - 20 °C could be proposed. Chemical analyses and mass balances from crystallization experiments in pilot scale were in good agreement with model calculations.

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Exploring green solvent for lupin debittering via *in silico* COSMO-RS predictive screening

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Andean lupins have been widely cultivated in the Andean highlands for the enrichment of the soil, animal feeding and as a food crop. Their seeds are nutrient-rich in terms of their high content of protein (about 50%) and oil (about 20%). However, the presence of toxic quinolizidine alkaloids, including sparteine, lupanine, and 13-OH-lupanine, in the seeds limits their industrial applications.

Several ionic liquids (ILs) and eutectic systems (ESs) have been reported in the literature to extract alkaloids from plant materials with the advantages of sustainability, biodegradability as well as adjustable polarity to dissolve polar and non-polar compounds. However, identifying an optimal IL or ES from the large number of possible combinations is quite challenging. To address this issue, the COSMO-RS (COnductor-like Screening MOdel for Real Solvents) model has emerged as a reliable computational tool that can screen numerous compounds or mixtures based on different thermodynamic properties.

The current study demonstrates the dissolution behaviour of quinolizidine alkaloids in ILs and ES using the COSMO-RS model. Several combinations were attempted and evaluated by predicting the activity coefficients ate infinite dilution (γ_{13}^{∞}) of typical quinolizidine alkaloids (sparteine, lupanine, 13-OH-lupanine, and lupinine) in conventional solvents, ionic liquids and natural-based eutectic solvents.

Through this research, we aim to develop a novel separation method based on the use of green solvents, to debitter the Andean lupin beans, i.e., to remove the toxic alkaloids, and to unveil efficient and environmentally conscious strategies for maximizing their use in the food industry.

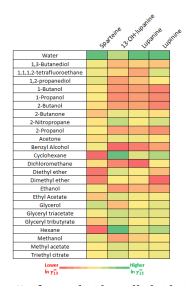


Figure 1. Preliminar results on the $\ln \gamma_{13}^{\infty}$ of quinolizidine alkaloids in different pure solvents at 298.15 K.

Acknowledgements: This work was supported by national funds through FCT/MCTES (PIDDAC): CIMO, UIDB/00690/2020 (DOI: 10.54499/UIDB/00690/2020) and UIDP/00690/2020 (DOI: 10.54499/UIDP/00690/2020); and SusTEC, LA/P/0007/2020 (DOI: 10.54499/LA/P/0007/2020).

Interaction and aggregation behavior of rhodamine B dye in water, ethanol and aqueous solutions of ionic liquids

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Xanthene dyes are the most widely investigated class of luminescent dyes, because of their special characteristics which make them useful for many applications. One of the most used xanthene dye is rhodamine B emitting in the red region of visible spectrum. Rhodamine dyes are used extensively in biotechnology applications such as fluorescence microscopy, flow cytometry and fluorescence correlation spectroscopy Dye aggregation in ionic liquid-based solutions is a crucial topic because dyes and these ecologically friendly designer solvents have a wide range of applications. The self-association or aggregation of rhodamine B (Rh B) in aqueous and aqueous.

The self-association or aggregation of rhodamine B in water, ethanol and aqueous solutions of Ionic liquids (ILs) has been studied by using spectrophotometry. The concentration ranges from $\sim 1 \times 10^{-5}$ M to $\sim 3 \times 10^{-4}$ M were selected to avoid the formation of higher aggregates such as trimer, tetramer etc. For instance, bathochromic J-aggregation can be easily distinguished from hypsochromic H-aggregation using UV/vis light spectroscopy. The impact of ILs namely tetramethylammonium Mercaptopropionate [TMA][MPA] and tetramethylammonium indole 3 acetate [TMA][IAA], a water-miscible room-temperature IL, on the behavior of an aggregation of dyes is evaluated. The addition of ~ 0.5 M ILs resulted in the precipitation of Rh B dye from the solution. The results have been explained in terms of bonding between dye molecule and solvents which affects the arrangement of molecules there by affecting extent of aggregation.

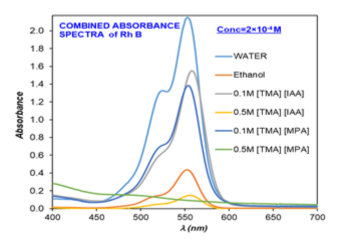


Figure 1. Absorbance spectra of RH B in water, ethanol and aqueous solutions of ILs $(2 \times 10^{-4} \text{ M})$ at room temperature

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Residual volume approach as a tool for prediction of biological activity of ionic liquids

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Enzymes are large protein biomolecules responsible for many chemical reactions that are necessary to sustain life. They are an integral part of any living organism, providing its proper functioning by regulation of processes such as growth, healing, digestion, reproduction, to name just a few [1].

This study examines how various hydrophilic ionic liquids (ILs) affect the diphenolase activity of mushroom tyrosinase, an enzyme responsible for the pigmentation of mammals' skin and hair, and involved in the molting process of insects and the browning of fruits and vegetables, as a model enzyme. It also aims to investigate the potential for predicting the biological activity of these ILs using The Residual Volume Approach (RVA), an empirical model originally developed for predicting IL properties such as density, viscosity, surface tension, etc [2-4].

The ILs studied were based on various cations and anions such as N,N,N',N',N'',N''-hexaalkylguanidinium $\{[N22N33N3ngua]^+\}$, 1-alkyl-3-methylimidazolium $\{[CnC1im]^+\}$, N,N,N,N-tetralkylammonium $\{[CnCnCnCnN]^+\}$ or N,N-dialkylpyrrolidinium $\{[CnCnpyrr]^+\}$ cations, coupled with anions such as chloride (Cl^-) , bromide (Br^-) , acetate $\{[AcO]^-\}$, trifluoroacetate $\{[TFA]^-\}$, thiocyanate $\{[SCN]^-\}$, dicyanamide $\{[dca]^-\}$, saccharinate $\{[Sac]^-\}$ and acesulfamate $\{[Ace]^-\}$.

The wide range of ILs used allowed several generalizations for their effect to be drawn: (i) ILs can be considered as potent tyrosinase inhibitors; (ii) guanidinium ion ensures the highest inhibitory activity; (iii) increase in the alkyl side chain length enhances the inhibitory effect; (iv) anion is of immense importance toward high inhibitory activity. Kinetic analysis was also performed to establish the inhibitory mechanism for the most active compounds.

These findings offer valuable insights into the interactions between hydrophilic ionic liquids and enzymes, providing a foundation for further research. Moreover, the RVA correlations demonstrated that this approach could be successfully applied to predict the enzyme-inhibitory, and possibly other biological activities of ILs.

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Separation of cobalt from nickel via polymer inclusion membrane with ionic liquid $[P_{66614}][Dec]$

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It is estimated that more than 1.2 million tons of batteries enter the European Union each year, with global demand predicted to grow considerably over the next five years [1]. As a result, the demand for Co is projected to increase 20 times by 2050 [2]. Proper disposal of used LIBs is essential to manage resources effectively and prevent environmental pollution from toxic materials [3]. LIBs contain Critical Raw Materials (CRMs) like cobalt, lithium, and graphite, which have high supply risks and price variability [4]. Recent research has focused on sustainable methods for recovering materials from spent LIBs, with hydrometallurgical recycling gaining attention for its low energy and resource use and high metal purity. Liquid-liquid separations using ionic liquids (ILs) have attracted interest due to their potential selectivity and superior characteristics compared to volatile organic solvents (VOCs), such as low volatility, safety, chemical and thermal stability, and recyclability [5]. Immobilizing ILs into polymer inclusion membranes (PIMs) reduces IL volumes which reduces the price, enables extraction and stripping in a single step which makes PIMs suitable for scaled-up processes [6].

In this work we prepared and characterized a PIM membrane based on PVC and IL [P₆₆₆₁₄][Dec] for separation of Ni and Co. Salts of cobalt and nickel were dissolved in 100 ml of HCl acid and used as a Feed phase in reactor shown in Figure 1. The best conditions (temperature, acid molarity, strip phase composition) were determined for metal separation. Metal content in Feed and Strip phase for kinetics and separation study was determined by ICP-OES, and absorption spectra was collected by UV-VIS.

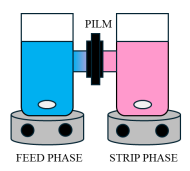


Figure 1. Setup for separation of cobalt from nickel. The volume of each compartment is 100 ml, contact surface with the membrane 28.27 cm², stirring speed 700 rpm.

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Solubilization of aqueous-insoluble phenothiazine drug in Triton X-100 micellar media

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The solubilization of poorly soluble drug phenothiazine (PTZ) in triton X-100 (TX-100) micellar aqueous media at physiological pH and the interaction of cationic surfactants, viz., tetradecyltrimethylammonium bromide (TTAB), hexadecyltrimethylammonium bromide (CTAB), cetylpyridinium chloride (CPC) and cetylpyridinium bromide (CPB) in presence of drug PTZ in TX-100 are discussed. The interaction of surfactants in PTZ–TX-100 media has been studied using conductometric technique at three temperatures. The conductometric results reveal the sequence of critical micelle concentration (CMC) values of surfactants as TTAB > CPC > CPB > CTAB. UV-vis spectroscopy studies were used to learn the interactions of surfactants with the drug PTZ in TX-100 aqueous media (at 303.15 K and pH 7.4) which suggest the interactions are highly feasible (change of Gibbs energy; $\Delta G_b < 0$) and greatly improve beyond the CMC values of corresponding surfactants. Fluorescence technique has been employed to study the aggregation number of TTAB/CTAB in PTZ–TX-100 systems at 303.15 K.

Figure 1. Molecular structure phenothiazine drug (PTZ).

Relative pK_a measurements in non-aqueous solvents

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Acids and bases are widely used in organic synthesis, chemical analysis, electrochemical processes, and the pharmaceutical industry. The pK_a value used to describe the strength of acids and bases is one of their most important properties. The strength of acids or bases affects their reactivity in chemical reactions, bioactivity, absorption and distribution in living organisms, etc. Accurate pK_a values in water have been determined for a myriad of compounds, but because a significant degree of the usage of acids and bases takes place in non-aqueous solvents, it is crucial to have reliable acidity and basicity data also in these solvents. There is significantly less acidity and basicity data available in non-aqueous solvents because of difficulties in pK_a measurements in such solvents. Moreover, although a significant amount of pK_a values have been determined, e.g. in acetonitrile (MeCN), dimethylsulfoxide (DMSO) and dimethylformamide (DMF), their quality is sometimes doubtful. For instance, in some cases, even simple and widely used carboxylic acids have published pK_a values in the literature that differ by more than an order of magnitude between different authors. For this reason, a reevaluation and revision of pK_a values in MeCN, DMSO and DMF is needed.

We have established measurement capability for pK_a measurements in non-aqueous solvents on the basis of a spectrophotometric method of measuring relative acidity/basicity (ΔpK_a) values. The main advantages of the relative pK_a measurement method are that there is no need to measure pH values and a number of uncertainty sources cancel out, either partially or fully. The most prominent result that we have obtained are extensive self-consistent pK_a scales for acids and bases in acetonitrile (MeCN), each containing over 230 compounds. Because of the nature of the measurement method, involving a number of check and continuous cross-validation, the quality of the results is considered high. Determining high-quality pK_a values also in other solvents (e.g. DMSO and DMF) is underway.

This presentation will describe the principle of our relative pK_a measurement method and show its advantages. A short overview of our recent results and ongoing research will also be provided.

OL21 Uncertainty analysis of solution chemistry data

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Uncertainty and sensitivity analysis is an invaluable tool for any modelling attempt, be it a simple chemical speciation calculation to far more complex analyses such as transport of radionuclides from a repository for used nuclear fuel. The founding principle is that we cannot determine anything precisely even of the level of precision may vary. Thus, all measured entities will be encumbered with uncertainties and as these values are subsequently used in e.g. modelling attempt also these results will be uncertain. It is therefore clear that in order to have any reliability in any results a through uncertainty analysis must be done and described together with the determined entity. In this presentation some techniques for uncertainty analysis of solution chemistry data will be given together with an analysis of these results in some modelling attempts.

Physicochemical investigation of ionic additives for improved lithium-ion battery performance and reliability

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The development of high-energy density, safe, affordable, and environmentally friendly energy storage devices is crucial for the energy transition. Lithium-ion batteries (LIBs) currently dominate the global energy storage market, but their safety, cost, and environmental impact are significant concerns. The highly flammable and toxic state-of-the-art electrolytes used in LIBs are a major concern, as they can pose a risk to both people and the environment.

The safety issues associated with LIBs are closely linked to their high energy density and the use of volatile and flammable organic-solvent-based electrolytes. As such, improving the thermal stability of electrolytes and enhancing the safety of LIBs is a key priority for our research.

To address this challenge, we are developing electrolytes that can perform multiple functions simultaneously, including stabilizing electrode materials and functionalizing electrodes. In this context, we conducted a comprehensive study involving physicochemical, thermal, and electrochemical testing to evaluate the stability of an electrolyte containing newly synthesized zwitterionic compounds as additives.

Our study utilized $1.0 \, M \, LiPF_6$ in a $50/50 \, mixture$ of ethylene carbonate and diethyl carbonate (EC/DEC) as our battery-grade electrolyte solution. We compared the electrochemical performance of the electrolyte with additives to commercially used electrolytes to assess their effectiveness.

Overall, our research aims to develop safer and more environmentally friendly energy storage devices by improving the thermal stability of electrolytes and exploring alternative additive compounds that can enhance the performance of lithium-ion batteries.

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Residual volume approach as a tool for prediction densities of ionic liquids at different pressures and temperatures

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A new method has been developed to predict the densities of ionic liquids (ILs) at higher temperatures and pressures. This method is based on an adjusted linear regression equation using the Residual Volume Approach (RVA). The method relies on the linear density dependency on the organic residue's molar volume. The correlation between density and molecular volume has been studied, and adjusted equations have been proposed to accommodate the effects of increased temperature and pressure. These newly defined equations depend on eight coefficients calculated from observed changes in the density of N-alkyl-N-methylimidazolium bis(trifluoromethylsulfonyl)imide type of ILs, as well as the combined effect of changes in molecular volume, temperature, and pressure. These equations can be used with acceptable accuracy. Additionally, a simplified equation requiring only five coefficients is presented. It is also possible to customize the equations to a specific type of cation-anion IL pair by following the method used herein to yield more accurate results. Using eight constants dependent only on residual volume, temperature, and pressure eliminates the need to know IL-specific coefficients, such as isobaric expansivity or isothermal compressibility, which are widely used in similar equations from literature.

The approach proposed can also be used to calculate the properties of other combinations of ions. The calculated results are more accurate when the nature of the IL pair is similar to that of the original. It is possible to fine-tune the equations by adjusting the parameters to accommodate the specific ionic pair, provided that enough density points are available while the equations retain their original form.

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Physicochemical characteristics and antimicrobial activity of nicotine-based ionic liquids with different length alkyl chains

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Nicotine is a pyridine alkaloid widely known for its recreational use as a stimulant and anxiolytic in the form of tobacco products and e-cigarettes. Nicotine causes addiction in humans and can lead to increased health risks and even cancer [1]. However, nicotine is a natural compound found in a large array of plant species. It is widely believed that the role of this secondary metabolite is that of a pesticide, due to its ability to bind to acetylcholine receptors in the nervous systems of animals, exhibiting harmful effects. For this reason it was used as an agricultural pesticide [2].

Due to nicotine's structure (Figure 1a) the tertiary nitrogen atom is susceptible to alkylation. The increase of an alkyl side-chain has shown to lead to, among other effects, the increase in antimicrobial activity and toxicity of a substance. Further, ionic liquids (ILs), ionic compounds with melting points below 100°C, are also known to have high biological activity [3]. Due to potential synergistic effects of the base substance, an added alkyl side-chain and the form of an IL, we synthesised four nicotine-based ionic liquids with varying alkyl chain lengths: N-octylnicotinium bromide (Figure 1b), N-decylnicotinium bromide (Figure 1c), N-dodecylnicotinium bromide (Figure 1d) and N-tetradecylnicotinium bromide (Figure 1e).

In order to understand the behaviour of these newly synthesised nicotine-based ILs, density, viscosity and sound velocity measurements of their water solutions were performed in a concentration range from 0.02 to 0.12. mol·kg⁻¹ and in a temperature range from 293.15 to 313.15K. The gathered data was further used to determine, among others, apparent molar volume, apparent molar volume at infinite dilution, *Jones-Dole* viscosity coefficient *B*, apparent molar compressibility and apparent molar compressibility at infinite dilution. The values gained are further discussed in terms of solute-solute and solute-solvent interactions. The antimicrobial activity of these ILs and nicotine was also tested against several strains of gram-positive and gram-negative bacteria, yeasts and molds and compared. It was found that all nicotine-based ILs have significantly higher antimicrobial activity than nicotine, for all examined species (Figure 1f).

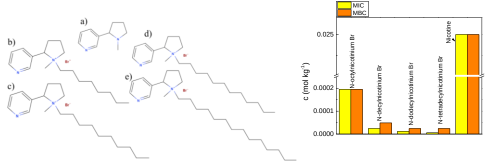


Figure 1. Structures of a) nicotine and nicotine-based ILs: b) N-octylnicotinium bromide c) N-decylnicotinium bromide d) N-dodecylnicotinium bromide e) N-tetradecylnicotinium bromide and f) a graph showing antimicrobial activity of these substances *Escherichia coli* bacteria

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Impact of the environmental conditions on the resilience of bioregulators in the aquatic environment

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Bioregulators encompass natural and synthetic compounds utilized directly on plants to enhance agricultural yields. Among the synthetic variants, Globaryll (G) and Ormoroc (O) stand out, aiding in regulating plant growth and development. However, their prolonged presence may lead to environmental accumulation at diminishing concentrations, potentially posing risks as organic pollutants [1].

We investigated the photolytic stability of G and O in a mixture of higher (100 and 20 mg/dm³, respectively) and lower (50 and 10 mg/dm³, respectively) concentrations in ultrapure (UPW) and stream water (SW) to understand their environmental stability and behavior. After applying bioregulators in the field, higher concentrations of both substances can be found in the natural environment. However, as the plant partially absorbs the bioregulators over time, their concentrations decrease. Therefore, it is essential to study the stability of bioregulators in both scenarios. The degradation efficiency of G in higher concentrations resulted in 38% degradation observed in UPW (Fig. 1a) and 55% in SW (Fig. 1b). In contrast, G in lower concentrations showed consistently high photolytic stability, whereby 8% in UPW (Fig. 1a) and 2% in SW was degraded (Fig. 1b). On the other hand, O showed high stability for both concentrations. In UPW, 4% degradation was observed for lower and 5% for higher concentrations. The degradation efficiency of O significantly increased in SW, with 13% degradation in lower concentration and 50% in higher concentration, likely due to additive or synergistic effects facilitated by the presence of inorganic ions.

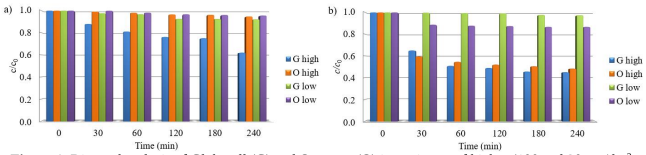


Figure 1. Direct photolysis of Globaryll (G) and Ormoroc (O) in a mixture of higher (100 and 20 mg/dm³, respectively) and lower (50 and 10 mg/dm³, respectively) concentrations in (a) UPW and (b) SW.

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The behavior of hydroxyapatite ceramics in the mimic de- and remineralization solutions

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The compositional and structural similarity of hydroxyapatite to solid tissues makes it suitable for bone reconstruction in the form of ceramic materials, coatings of metal devices, etc. When placed in the body, hydroxyapatite interacts with the body fluids, and depending on the environment, processes of dissolution (demineralization) or reverse deposition of mineral crystals (remineralization) can occur.

The behavior of hydroxyapatite ceramics in the mimic de- and remineralization solutions from the oral cavity was investigated in this study. Dense ceramic tablets from monophasic hydroxyapatite were prepared by pressing and sintering at 1000°C of nanocrystalline carbonate apatite. The precursor was synthesized via wet precipitation method from 1M solution of Ca(NO₃)₂ and 0.6M solution of (NH₄)₂HPO₄ at a Ca/P ratio of 1.67, pH 12 (keeping with NH₄OH), open-air atmosphere and continuous stirring. The resulting precipitate was matured for 24 h, centrifuged, washed, and dried at 90°C for 15 h. A solution containing 0.1 mM of lactic acid, 2.2 mM CaCl₂, and 2.2 mM NaH₂PO₄ was chosen as a demineralizing agent with pH 4.5. It mimics the conditions in the oral cavity at the bactericidal decomposition of sugars producing lactic acid. Remineralization was examined in the presence of artificial saliva after biostimulating the tablet surface with a solution containing polycarboxybetaine (PCB). PCB is a zwitterionic polymer with unique properties – it creates conditions for mineral nucleation and can serve as an antifouling coating. The experiments were carried out under such conditions as to avoid mineral deposition resulting from sedimentation.

The results of X-ray photoelectron spectroscopy (XPS) reveal that in the initial ceramic tablets, Ca, P, O, and C are the main elements present on the surface in the form of hydroxyapatite and calcium carbonate. The element concentrations are close to the one in the tooth email. Insignificant amounts of Na, Cl, and Si were also found. During the demineralization process, no significant changes were observed in the state of the main elements, except for an increase in the amount of carbonate-type carbon and absorbed water and the appearance of less than 3% nitrogen. The quite noticeable surface change was detected after the remineralization procedure of the sample staying for 12 hours in a demineralizing solution. All peaks become asymmetric indicating a new microenvironment on the surface originating from the components of artificial saliva, namely the appearance of C–N and Na-P-O bonds and increasing the amount of hydroxy and carboxylic groups, which may serve as centers of mineralization.

The images from scanning electron microscopy show a difference in the surface morphology after 6 and 12 h exposure to demineralization respectively. During the first hours, the dissolution proceeds in a spiral fashion, with a greater rate in the center and a lower rate in the periphery. As a result, pits on the grain surface and holes at the grain boundary are formed. As time increases, the concentration of Ca²⁺ ions in the solution increases, the driving force of dissolution decreases, and the two rates equalize. Thus, after 12 h of stay in a demineralizing environment, the surface is smoother, without sharp boundaries between individual hydroxyapatite crystals.

After the remineralization procedure, crystal grains are covered with a diaper due to the deposition of the newly formed mineral phase. The Ca/P ratio of 1.58 - 1.71 reveals apatite type mineral phase in which Na⁺ ions are included. The C–N bond, detected by XPS analysis is due to the absorbed organic compounds from the artificial saliva.

In conclusion, hydroxyapatite ceramic obtained by us reveals de and remineralization behavior similar to the tooth enamel and could be used both for tooth restoration and as a primary model for enamel research. **Acknowledgments**: The authors thank the Bulgarian Ministry of Education and Science for the financial support under the project KP-06-H49-6/2020. In this investigation, we used the research equipment of Distributed Research Infrastructure INFRAMAT, part of the Bulgarian National Roadmap for Research Infrastructures, supported by the Bulgarian Ministry of Education and Science.

Graphitic carbon nitride and tetrabutylammonium chloride modified carbon paste electrode for efficient electrochemical sensing of carbamazepine

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Carbamazepine (CBZ) is an anticonvulsant, antiepileptic and mood stabilizing drug used primarily in the treatment of epilepsy and bipolar disorder [1]. As a result of its widespread use, CBZ determination in biological and pharmaceutical samples is of great importance. Different analytical methods are reported for CBZ determination, whereby there is an increasing interest in the development of electrochemical methods for determining CBZ due to their advantages such as simplicity, portability, reliability, and cost-effectiveness. Although CBZ direct electrochemical measurement is possible at different electrodes, researchers mostly used carbon-based electrodes in their modified form [2]. Ionic liquids (ILs) draw attention in electroanalysis as modifiers of carbon paste electrodes (CPE) because of their specific features, such as possible catalytic activity, wide electrochemical window, electron-transfer acceleration and high conductivity [3]. Additionally, IL modified electrodes showed an excellent electroanalytical performance in detection of electroactive species by enhancing faradaic responses and consequently the signal-to-noise ratio [4].

This study utilizes the preparation of CPE bulk-modified with a type of two-dimensional conjugated polymer, i.e., graphitic carbon nitride (g- C_3N_4), and IL tetrabutylammonium chloride (TBACl) to obtain advanced electrochemical sensor for CBZ. Cyclic voltammetric experiments showed that TBACl-g- C_3N_4 -CPE has an improved electrochemical response compared to unmodified CPE and g- C_3N_4 -CPE due to the synergistic effect of electrode modifiers, as well as that the CBZ oxidation process is irreversible and diffusion-controlled. Using direct anodic square-wave voltammetry (SWV) under optimized conditions at pH 7.0, the CBZ sensor showed a linear dynamic range from 0.10 to 2.20 μ g mL⁻¹, a limit of detection (LOD) of 0.030 μ g mL⁻¹ and relative standard deviation lower than 3%. No significant influence of inorganic ions or organic compounds on sensor response was verified. The practical applicability of TBACl-g- C_3N_4 -CPE was tested for quality control of CBZ pharmaceutical formulation Karbapin® and for the determination of CBZ in spiked human urine sample. Based on the obtained results it can be concluded that TBACl-g- C_3N_4 -CPE could be successfully applied as a very simple and perspective analytical tool for the determination of CBZ in different matrices with excellent recovery and reproducibility.

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Exploring protic ionic liquids-water interactions: Experimental and computational analysis of 2-hydroxypropylammonium formate

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Ionic liquids (ILs) are a unique class of compounds that have garnered significant academic and industrial attention due to their distinct properties such as low volatility, high thermal stability, and a broad liquidus temperature range. These salts, which are liquid below 100°C, exhibit remarkable solvating abilities owing to their ionic nature. Among them, protic ionic liquids (PILs) are of particular interest for various industrial applications due to their economic feasibility and straightforward synthesis. The production of PILs involves the direct neutralization of a Brønsted acid with a Brønsted base, often in a solvent-free process, making it cost-effective and environmentally friendly [1,2].

The versatility of PILs is further enhanced by the wide variety of available Brønsted acids and bases, allowing for the creation of PILs with tailored properties. This customization is crucial for industrial processes where specific physicochemical properties, such as viscosity, conductivity, and thermal stability, can be fine-tuned. Adjusting the hydrophobic or hydrophilic nature of PILs by selecting appropriate acid-base combinations also improves their efficacy as solvents, catalysts, or electrolytes in various chemical reactions and processes.

PILs exhibit exceptional physicochemical properties, including low volatility, high thermal stability, high ionic conductivity, and wide electrochemical windows. These attributes make them suitable for applications in energy storage and conversion systems, such as batteries and fuel cells. Moreover, PILs are generally biodegradable and less toxic than traditional organic solvents and aprotic ionic liquids, making them more environmentally friendly [3,4].

In this study, the protic ionic liquid 2-hydroxypropylammonium formate [2-OHC₃NH₃][Fo] was synthesized through a solvent-free neutralization process using non-toxic, cost-effective formic acid and (\pm)-1-amino-2-propanol. The physicochemical properties of the synthesized ionic liquid and its mixtures with water were measured across the entire range of mole fractions and within the temperature interval of T = (278.15 - 333.15) K. These properties included density, viscosity, and electrical conductivity. Additionally, the thermal profile of the synthesized ionic liquid was presented.

The study provided an in-depth analysis of the interactions between the ionic liquid and water, focusing on how variations in temperature and composition affect the hydration properties and ionization degree of the ionic liquid. Computer simulations were also conducted to gain further insights into the structural organization of water molecules around the ions of the ionic liquid.

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Synthesis, characterization, and antimicrobial activity of an ionic liquid 1-butyl-3-methylimidazolium 2-hydroxy-2-phenylacetate

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Industrial processes often use organic solvents that are toxic and harmful to the environment and human health. To protect our environment, harmful solvents must be replaced with non-toxic and biodegradable solvents.

That is why green chemistry was developed, i.e., procedures for obtaining chemical products and processes that reduce or eliminate the use and creation of hazardous substances. It deals with the development of safe, non-flammable, and non-toxic compounds, and ionic liquids (ILs) fully meet these requirements. Ionic liquids are salts consisting of a large organic cation and a small organic or inorganic anion whose melting point is below 100°C. Most of them are already in a liquid aggregate state at room temperature (room temperature ionic liquids - RTIL). The main advantage of ionic liquids is the possibility of their design because by combining anions and cations, the desired properties are achieved, which makes it possible to overcome problems present in various fields. For example, in the last two years, disinfection has become commonplace in everyday life as a preventive measure against the spread of the new SARS-COV-2 virus. Ionic liquids can be designed so that they are non-volatile and, at the same time, exhibit antimicrobial and antiviral effects, thus ensuring long-term disinfection of the surfaces to which they are applied.

This work aims to synthesize and characterize aqueous solutions of the newly synthesized ionic liquid 1-butyl-3-methylimidazolium 2-hydroxy-2-phenylacetate. In the synthesis of the ionic liquid, mandelic acid, and 1-butyl-3-methylimidazolium chloride were used as starting components. One of the most important properties of mandelic acid is its bacteriostatic properties, while 1-butyl-3-methylimidazolium chloride shows good physicochemical properties derived from the aromatic imidazole ring. The density and viscosity of the ionic liquid at different temperatures were measured, and then the interactions that exist in aqueous solutions of this ionic liquid were considered. Then, the antimicrobial activity of the corresponding ionic liquid was tested, and it was determined whether it could serve as a disinfectant.

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N-methylimidazolium-based ionic liquids as potential modulators of metabolic processes

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N-methylimidazolium-based ionic liquids were synthesized and characterized as potential modulators of betaoxidation of fatty acids in the organism via inhibition of Carnitine Acetyltransferase (CrAT). The inhibitory effect of the synthesized compounds was tested *in vitro* and compared with controls: Meldonium (an approved drug for the treatment of coronary heart disease, under the trade name Mildronate®) and MeGBB (a compound in third-phase clinical trials).

All compounds demonstrate inhibitory effects in low millimolar concentrations, some being more active than the controls. The IC50 values of all compounds were determined. Structure-activity relationships (SAR) were elucidated - a) effect of the length of carbon chain - R (C2-C10), b) presence of different substituents attached to carbon chain - R (e.g. hydroxyl, nitrile, ester group), c) carbon chain branching and d) effect of the anion (e.g. Cl-, Br-, salicylate, caffeate) With the most active compounds of the group, kinetic studies were conducted to determine the mechanism of inhibition of Carnitine Acetyltransferase (CrAT).

The results showed that N-methylimidazolium-based ionic liquids are potential modulators of metabolic processes and drug for treatment of coronary heart decease, diabetes and even some types of cancer.

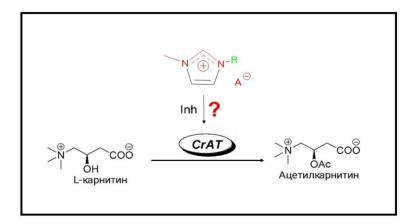


Figure 1. N-methylimidazolium-based ionic liquids are potential inhibitors of fatty acids oxidation as they supress conversion of L-carnitine to AcetylCarnitine.

P10 Sphalerite dissolution in deep eutectic solvents

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Sphalerite (ZnS) is the main resource of zinc. The conventional method of zinc recovery from sphalerite involves roasting, leaching, and electrolysis. However, this process suffers from environmental issues, high energy demands, and the generation of impurities. Such drawbacks, along with the current drive for the minerals processing industries to develop low-energy, low-cost, and more environmentally sustainable processes, have led to the recent invention called ionometallurgy. Ionometallurgy aims to dissolve ores at low temperatures using non-aqueous solvents [1]. Promising results have been reported regarding the use of ionometallurgy for the treatment of sulfide minerals [2].

Here, we report our experimental results on the dissolution of sphalerite in **d**eep **e**utectic **s**olvents (DESs). DESs are binary or ternary mixtures with deep melting point depression. Three well-known DESs, namely Ethaline (1 mole choline chloride: 2 moles ethylene glycol), Reline (1 mole choline chloride: 2 moles urea), and Oxaline (1 mole choline chloride: 1 mole oxalic acid dihydrate), have been used for dissolution experiments. The results revealed that using DESs improves the dissolution of sphalerite under both non-oxidative and oxidative conditions compared to an equivalent acidic aqueous solution (Figure 1). The highest solubilities were found using Oxaline. Oxidants were found to influence the dissolution of sphalerite in both DESs and aqueous media, with cupric ions being most effective in DESs and ferric ions in aqueous media.

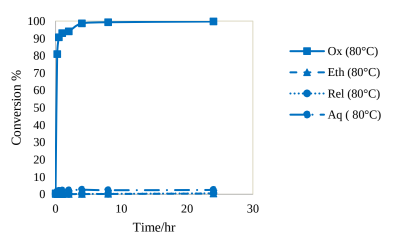


Figure 1. Conversion of sphalerite in Oxaline (Ox), Ethaline (Eth), Reline (Rel), and aqueous acidic (Aq) solutions, solid to liquid ratio = 1:100.

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Influence of the ionic liquids based electrolytes on the tomato (Solanum lycopersicum L.) and cucumber (Cucumis sativus L.) growth, development and oxidative stress

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Ionic liquids have been identified as possible alternatives to traditional organic solvents in lithium-ion battery electrolytes due to their distinct properties, including high thermal stability, low volatility, and wide electrochemical windows. These features could make them safer and more efficient. Despite the benefits, there are concerns about the environmental consequences of using ionic liquids in lithium-ion battery electrolytes. One significant concern is the threat of soil and plant contamination. If these liquids spill or leak from used lithium-ion batteries [1,2], they could enter the soil through various pathways and contaminate crops such as cucumber and tomato plants, which have extensive root systems that may facilitate the absorption of ionic liquids. After absorption, some electrolyte components could accumulate inside the plants and have toxic effects, potentially harming plant growth and crop yields.

This study investigated how spilling electrolytes with varying combinations of ionic liquids, organic solvents, and lithium salts in different concentrations affects the growth and development of tomatoes and cucumbers. The focus is on ionic liquids in mixture with propylene-carbonate. Special attention was paid to examining the influence of electrolyte components on aerial parts and/or fruits of these plants and the levels of metabolites involved in antioxidant protection under stressful conditions, such as malonyldialdehyde and photosynthetic pigments like chlorophyll a, chlorophyll b, and carotenoids.

Acknowledgements: The authors are grateful for the financial support of Provincial Secretariat for Higher Education and Scientific Research, grant number: 142-451-2545/2021-01.

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Quantitative precipitation of cobalt based oxide material from battery recycling solutions

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Lithium-ion batteries are composed of a graphite anode and a mixed metal oxide cathode (Li MO_2 ; M = x Ni, y Co, z Mn; x + y + z = 1), which are separated and recycled to reuse their valuable components when reaching their end of life [1].

In this study we investigate the reaction of cobalt(III) amines in lithium hydroxide solution to produce lithium cobalt oxide compounds by precipitation, as a possible step to produce the active material $LiCoO_2$ directly from black mass leachates.

It is well known that hexammine cobalt(III) ions form cobalt oxyhydroxide precipitates in alkaline media (Eq. 1). Due to similar crystal structures of CoO(OH) and LiCoO₂, an ion exchange of hydroxide protons and lithium ions takes place in solution (Eq. 2)[2,3].

$$[Co(NH_3)_6]^{3+} + 3 OH^- \rightarrow CoO(OH) + 6 NH_3 + H_2O$$
 (Eq. 1)

$$CoO(OH) + Li^+ + OH^- \rightarrow LiCoO_2 + H_2O$$
 (Eq. 2)

In our research, identification of phases via P-XRD, Raman spectroscopy as well as investigation of chemical composition by thermoanalytical methods and wet chemical analysis confirmed the presence of $\text{Li}_x\text{CoO}_{1+x}(\text{OH})_{1-x}$ (0.48 < x < 0.93). The phase composition is adjustable by modifying the reaction temperature (70 - 200 °C), reaction time (2 - 48 h), and lithium concentration up to saturation of LiOH·H₂O.

Analysis of the supernatant shows cobalt recovery of >99.9 %, which was confirmed by UV-Vis spectroscopy. Lithium and hydroxide concentrations decrease by precipitating hydroxidic lithium cobalt(III) oxide. Counterions, e.g. chloride and sulfate, do not take part in the precipitation process. In reactions under reflux only 20 % of the maximum amount of ammonia was detected. Contrarily, ammonia remains up to 90 % in hydrothermal reactions performed in autoclaves, which can be recovered as starting material to resynthesize cobalt(III) ammine compounds.

We aim to use these precipitates as potential cathode materials in lithium-ion batteries.

Acknowledgements: The authors would like to thank K-UTEC Salt Technologies for funding.

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Studying the impact of converting creatine and guanidinoacetic acid from zwitterionic to cationic form on their water solubility

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Creatine (CR) is an endogenous amino acid naturally synthesized by the human body to provide energy to muscles. It facilitates the regeneration of adenosine triphosphate (ATP) utilized during muscular exertion, resulting in enhanced strength and endurance [1]. While creatine monohydrate is the most extensively researched and widely utilized form of creatine, it is associated with limitations such as poor water solubility, spontaneous conversion to creatinine, and the requirement for specific transporters for cellular uptake. Consequently, there is a need to explore novel formulations to mitigate these drawbacks [2].

Guanidinoacetic acid (GAA) serves as a natural metabolic precursor to creatine, undergoing biological conversion into creatine via hepatic methylation. Recent scientific literature suggests that the synergistic combination of creatine and GAA is more efficacious in elevating creatine levels in serum, muscle tissue, and the brain compared to the use of pure creatine [3]. However, both creatine and GAA exhibit relatively low water solubility, underscoring the desirability of converting them into more soluble hydrochloride salts. While several advanced formulations in the sports supplement market incorporate creatine hydrochloride salts, the solubility of creatine, creatine hydrochloride, and GAA hydrochloride remains inadequately studied. In contrast, the solubility of GAA in water has been previously investigated [4].

Consequently, this study seeks to first convert creatine and GAA into hydrochloride salts and subsequently ascertain the solubility of creatine, creatinine, creatine hydrochloride, and GAA hydrochloride in water within the temperature range of T = (293.15-313.15) K at atmospheric pressure. The findings of this investigation will facilitate a comprehensive analysis of the thermodynamics governing the dissolution of the compounds.

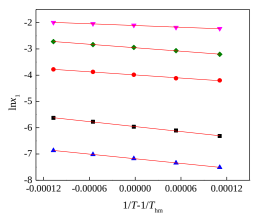


Figure 1. The plot of natural logarithm of the mole fraction of (■) creatine, (●) creatinine, (△) GAA, (\checkmark) creatine hydrochloride, (♦) GAA hydrochloride in a saturated aqueous solution, $\ln x_1$, against 1/T- $1/T_{hm}$.

Acknowledgements: The authors are grateful for the financial support of Provincial Secretariat for Higher Education and Scientific Research, grant number: 142-451-2545/2021-01.

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Influence of differently structured aqueous solutions on organic reactions

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To make organic reactions more environmentally friendly and to reduce waste caused by organic solvents, more and more reactions are tried to be carried out in aqueous systems. Water has the advantage of being nontoxic, environmentally friendly, and non-flammable. However, the challenges of using aqueous systems are the poor solubility of the reactants and the possible decomposition of these substances. In the literature, the yield of the reactions is often increased by the addition of surfactants, like TPGS-750-M. On the one hand, the formation of micelles can increase the solubility of the substances due to the more hydrophobic area of the micelles, and on the other hand, according to literature, the reactions are improved by the compartmentation of the solvent.

The physical background, such as solubility, structuring, and specific interactions with the solvent that influence different organic reactions has been investigated.

To investigate the influence of different aqueous systems, a cascade reaction consisting of a Heck reaction followed by an enzyme catalysis from the work of Cortes-Clergent *et al.* was carried out [1]. The reaction was performed in water, a surfactant-based and surfactant-free microemulsion, and in a non-structured binary mixture. The highest yield was obtained in the surfactant-free microemulsion. However, the higher yield compared to the reaction in pure water, cannot be attributed to the structuring of the system but to the solubility of the reaction components, as the reaction proceeds better in the non-structured binary system than in the micellar solution [2].

Furthermore, the synthesis of dihydroquinolinones with a rhodium catalyst from the work of Linsenmeier *et al.* was analysed [3]. The reaction was also performed in water, a surfactant-based and surfactant-free microemulsion, and in binary systems without structuring. In this reaction, the highest yields were obtained in the structured solvent systems. The increased yields can possibly be explained by the incorporation of the reactants into the interface. In the case of the surfactant-containing system, interactions occurring between the reactants and the surfactant were determined with NOESY NMR. The solubility of the reactants seems to play a subordinate role in this reaction, as the solubility in the binary non-structured mixtures is similar to the solubility in the surfactant-free microemulsion. However, it should be mentioned that the differences in the yields between the structured and unstructured systems is only about 12%.

Due to the mentioned small differences in the yields between structured and unstructured systems of the last reaction, the esterification by lipases in aqueous systems will be investigated in future work (from the work of Singhania *et al.*) [4]. Normally, lipases would catalyse the reverse reaction, i.e., the cleavage of the ester, at increased water content. This reaction was also carried out in various solvents. However, up to this point, only measurable yields were obtained in surfactant-containing systems. This allows the assumption that in this case the structuring and the interaction of the surfactants with the enzyme and the reaction partners play a decisive role.

The yields of the products are determined with NMR measurements.

In summary, it can be said that the influence of solvents, especially surfactant-based and surfactant-free microemulsions, can vary depending on the reaction. In some cases, it is sufficient to increase the solubility of substances, for which no surfactant is necessary, while in other reactions the micellar structuring actually has an influence on the reaction.

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Critical evaluation of volume properties of the zirconium-based fluorides melts $MF-K_2ZrF_6$ (M = Li, Na, and K)

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MF- K_2ZrF_6 (M = Li, Na, and K) melts are important systems in metallurgy applications. Based on the experimentally measured data of density (determined by the Archimedean method) molar volumes, partial molar volumes and the excess molar volumes were calculated. Partial molar volumes of K_2ZrF_6 were obtained by using two different approaches - simple polynomial regression analysis and multicomponent polynomial regression in the form of the Redlich-Kister equation [1]. The excess molar volume possesses negative values throughout the entire concentration range for all three investigated systems. These negative values of the excess partial molar volumes of K_2ZrF_6 are similar in the case of the LiF and NaF and are slightly suppressed in the KF system. The excess partial molar volume of K_2ZrF_6 in KF shows smaller overall volume contraction that adopts 60% of volume contraction in LiF or NaF systems.

Systems MF- K_2 ZrF₆ (M = Li, Na and K) were converted to ZrF₄ coordinate and compared with the literature data. The partial molar volumes of ZrF₄ of the systems MF-ZrF₄ (M = Li, Na and K) are also significantly different in KF system in contrast to LiF and NaF systems.

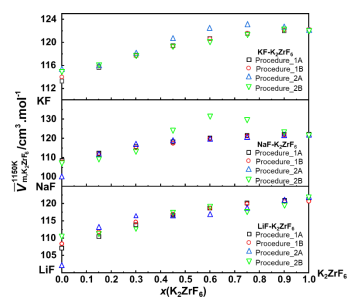


Figure 1. Concentration dependence of partial molar volumes of K₂ZrF₆ at 1150K.

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Solubility data of basic magnesium chloride hydrates (Sorel phases) and modelling with THEREDA in the oceanic salt system

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Basic magnesium chloride hydrates are the solid phases (Sorel phases) in the aqueous system Mg-Cl-OH-H₂O and the binder phases of magnesia building material (Sorel cement or concrete). They have special importance for the construction of geotechnical barriers (plug and sealing systems) in salt formations as well as for the adjustment of geotechnical environments to prevent transport processes of radionuclides.

The general composition of Sorel phases is $Mg_kCl(OH)_m \cdot nH_2O$, beyond that they can be described as x-y-z phases according to the double salt hydrate notation $xMg(OH)_2 \cdot yMgCl_2 \cdot zH_2O$, too. Various Sorel phases (x-y-z = 3-1-8, 5-1-8, 9-1-4, 2-1-2, 3-1-0) are known and all of them have been well characterized.

To ensure the long-term stability against salt solutions containing Na^+ , K^+ , Mg^{2+} , Ca^{2+} , Cl^- and SO_4^{2-} the solubility diagrams with respect to crystallization branches of the basic magnesium chloride phases must be known at the temperature of interest. Solubility data in the ternary subsystem Mg-Cl-OH-H₂O have been published at 25°C to 120°C by [1]-[4]. In the quaternary subsystem Na-Mg-Cl-OH-H₂O solubility data are available only at 25°C ([3], [5], [6]).

Here we present our latest experimental solubility data of Sorel phases in terms of OH⁻ molalities and H⁺ molalities in the ternary system and quaternary system at NaCl saturation depending on temperature [7]. In addition, our extended and adjusted data set, to include much higher temperatures of solubility constants of the Sorel phases (3-1-8, 9-1-4, 2-1-4) and Pitzer parameters, is implemented to the thermodynamic reference database THEREDA [8] and is discussed in the context of experimental data.

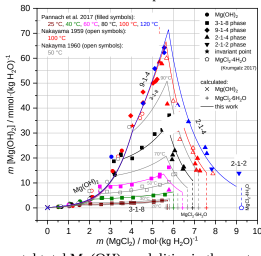


Figure 1. Calculated and experimental total Mg(OH)₂ molalities in the system Mg-Cl-OH-H₂O between 25°C and 120°C [9].

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N-hydroxyamides' p K_a determination by NMR spectroscopy

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The N-hydroxyamides are siderophores present in nature with impact on agriculture and medicine[1-2]. They form metal complexes, mainly with iron(III), which modify the overall concentration of such cations in the aqueous systems. For the building of predictive models where N-hydroxyamides are present, their acid behaviour in water, expressed by the pK_a value, must be considered.

We have measured the aqueous pK_a values of five different *N*-hydroxyamides, two *N*-hydroxyimides, and three reference compounds (succinimide, benzotriazole and vanillic acid). Some of them were synthesized in the laboratory, others had been purchased. Due to the impurities or degradation products in some compounds, the 1H and ${}^{19}F$ NMR spectroscopy were the chosen pK_a measurement techniques. Measurements were carried out at 298 K. The NMR enabled the identification of the peaks corresponding to the actual measured compounds, even in the presence of impurities. The observed changes in the chemical shifts $(\Delta\delta)$ with pH were very close to theoretical relations between $\Delta\delta$ and pH, both for monoprotic and diprotic acids. Both absolute (pK_a) and relative (ΔpK_a) between an acid and a reference acid) values were calculated. The ΔpK_a values between different acids were compiled into a ladder scale and, together with the absolute pK_a , used for achieving higher self-consistency and accuracy of the results. The activity coefficients were calculated using the Debye-Hückel theory.

Because of the multiple repetitions and the inclusion of relative measurements against reference pK_a values, we concluded that our pK_a values were more reliable than some literature data. New experimental data were also acquired, such as for vanillic acid, 2,2,2-trifluoro-N-hydroxyacetamide and the second acidity constant of 2-(hydroxyamino)-2-oxoethan-1-aminium.

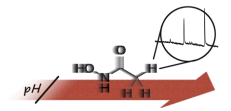


Figure 1. Graphical Abstract – The NMR heartbeat of *N*-hydroxyacetamide running on a p*H* scale.

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Supersaturation of curcumin in plant protein-based carrier system

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Curcumin is a polyphenol extracted from the plant Curcuma Longa L. (C. Longa). Besides its bright yellow colour, the compound is an intensively studied phytochemical with a variety of highly praised health beneficial properties. Due to the low bioavailability and low chemical stability of curcumin in physiological conditions, it remains challenging to access these properties to such an extent that it has a proven positive effect on human health. [1]

The solubility in aqueous systems and thus in a broader sense, the bioavailability of curcumin, can be increased with protein-based carrier systems. [2] A non-equilibrium state of solubilised curcumin is achieved by binding the polyphenol to pea protein. To the best of our knowledge, this is not reported in the literature so far. Within 24 hours, precipitation occurs in the initially translucent sample. Considering the non-problematic nature of the pea protein and the magnitude of the supersaturation (factor of 5-10), a non-equilibrium-formulation of curcumin and pea protein is interesting for pharmaceutical applications.

For a deeper understanding of the solubilisation process and the supersaturation, the solutions of curcumin in aqueous pea protein (PPI-Cur) were analysed according to their (chemical) stability, the particle sizes, the role of the protein in the solution and variation in the parameters of the preparation process. These analyses are fundamental and important steps in evaluating the suitability of curcumin as a health-promoting component.

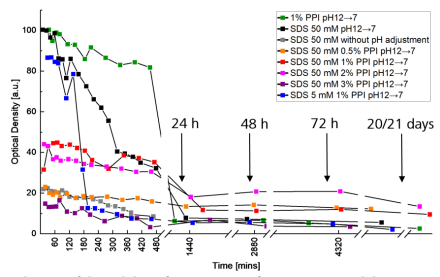


Figure 1. Evaluation of the stability of PPI-Cur (+surfactant) via optical density measurements.

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Green electrochemical sensor based on biochar for quantification of selected pesticides in aqueous solutions

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The sustainability of materials for developing the electrochemical sensor is crucial in the framework of a circular economy due to the need for environmentally friendly and greener analytical chemistry. The decrease in the impact of waste on the environment requires innovative approaches for biochar (BC) production and usage [1-2]. As a highly porous and carbon-rich material, BC has a desirable role as a catalytic material to enhance the analytical performance of BC-based sensors [3]. There has been a growing interest in the development of rapid and cost-effective techniques for detecting pesticides using electrochemical sensors with the possibility of their modification with sustainable materials [4]. Since the modification of carbon paste electrode (CPE) could improve the selectivity and sensitivity of trace level analysis of various electroactive analytes [5], the application of CPE modified with BC was investigated.

A sensitive and selective analytical method is developed regarding the voltammetric determination of fungicides maneb (MAN) and mancozeb (MCZ) using BC-CPE in an aqueous solution. The experimental conditions including pH of the supporting electrolyte, amount of BC in CPE, and differential pulse adsorptive stripping (DPAdSV) parameters were optimized. Under the optimal working conditions, the determination of MAN and MCZ was performed by applying the following DPAdSV parameters: $E_{acc} = -0.2 \text{ V}$, $t_{acc} = 90 \text{ s}$, and $E_{acc} = -0.2 \text{ V}$, $t_{acc} = 30 \text{ s}$, respectively. The linear increase of pesticides oxidation peak was recognized in a concentration range from $0.049-1.84 \ \mu g \ mL^{-1} \ MAN$ and $0.025-2.78 \ \mu g \ mL^{-1} \ MCZ$ in an aqueous Britton-Robinson buffer pH 7.0 using CPE modified with 10% BC. The relative standard deviation of six replicate measurements of MAN and MCZ was 3.2% and 2.9%, respectively, indicating a good repeatability of the developed DPAdSV method. The evaluated limit of detection of 0.015 $\mu g \ mL^{-1} \ MAN$ and 0.0075 $\mu g \ mL^{-1} \ MCZ$ indicated that the modification of CPE by BC provides a fast and sensitive determination of target analytes in an aqueous solution.

By exploring innovative solutions, electrochemical sensors based on BCs could provide monitoring of aquatic environmental samples to the control detrimental effects of pesticide residues.

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Chromatographically Examination of thiocarbohydrazone's lipophilicity

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Thiocarbohydrazone derivatives have become popular in contemporary scientific research due to their exceptionally antioxidant, antimicrobial and antitumor activity. Thanks to the presence of nitrogen and sulfur atoms, they easily form coordination bonds and build complexes with many metals, which have shown significant antiviral activity.

Knowledge of the new compound's lipophilicity is of crucial importance to assess the existence of its biological activity. In the first phase, a theoretical assessment of the thiocarbohydrazone derivatives' bioavailability and their ecotoxicity was performed. Also, their lipophilicity was determined *in silico*, as well as by using reversed phase thin-layer chromatography (RPTLC18F254s) in the presence of two organic modifiers (ethanol and dioxane). The effect of the substituent's nature and position, and the impact applied organic modifier on the chromatographic behavior of the analyzed derivatives were examined. The possibility of applying the chromatographic parameters ($R_{\rm M}^0$ and m) of the tested thiocarbohydrazone derivatives as alternative measures of their lipophilicity and ecotoxicity was studied by their correlation with partition coefficient (logP) and effective concentration (EC_{50}). As a result, satisfactory mathematical models were obtained.

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Examination of caffeine taste, hydration properties, and solubility in the presence of different additives

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Caffeine, the most widely used psychoactive substance, is classified as a xanthine alkaloid. It is not only commonly found in coffee, black and green tea, soft drinks, and energy drinks, but also in medicinal products, making it a regular part of the daily routine for approximately 80% of the global population [1]. The prevalence of hydrophobic in water-poorly soluble drugs and supplements is a major problem in drug design, food science, delivery, drug effect, and bioavailability.

The solubility of caffeine in water is relatively low, approximately 16 mg/mL at room temperature. This limited solubility is attributed to the self-association and aggregation of caffeine molecules through hydrophobic interactions [2]. The low solubility of caffeine in water presents a significant challenge for products stored or consumed at low temperatures, such as certain beverages, supplements, pharmaceuticals, and cosmetics. The addition of some biocompatible molecules, excipients or hydrotropes can lead to better caffeine solubility in water. To this day, only the effects of several compound classes on enhancing solubility and preventing self-aggregation have been studied. However, a systematic and comprehensive investigation is required to derive definitive conclusions.

In addition to caffeine's low solubility in water, another concern is that caffeine can induce bitterness at high concentrations. Solving this problem is very important from the food and beverage industry aspect.

Our research aims to examine the solubility, structural organization and taste of caffeine molecules in the presence of seven different additives suitable for wide use in the pharmaceutical, food and cosmetic industries.

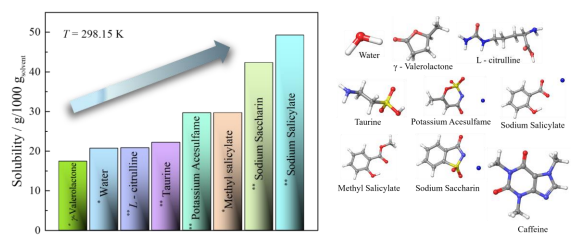


Figure 1. Compared the solubility of caffeine in the presence of different additives: * pure solvents, ** water solutions, at T = 298.15 K, along with their chemical structures.

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P22 CsF-Al₂O₃ system The phase and structural analysis

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A critical evaluation of the chemistry and the thermal stability of cesium fluoro-, oxo- and oxo-fluoro-aluminates in the CsF–AlF₃–Al₂O₃–Cs₂O system were carried out. Precise research of the binary CsF–Al₂O₃ sub-system was done by means of combining X-ray powder diffraction, high field solid state NMR spectroscopy, and thermal analysis methods. Cs₃AlF₆, CsAlO₂, Cs₂Al₂₂O₃₄ and new phase, Cs₂Al₂O₃F₂, were identified in the system. The structure of this new cesium oxo-fluoro-aluminate was determined. It is build up from single layers of oxygen connected (AlO₃F) tetrahedra, those layers being separated into by fluorine atoms. The thermal analysis of CsF–Al₂O₃ system revealed that it can be more defined as a pseudo-binary sub-system of the more general quaternary CsF–AlF₃–Al₂O₃–Cs₂O system. From phase analysis of individual phase fields, the mutual metastable behavior of all founded phases can be considered. Fluoro- and oxo-aluminates are metastable precursors of thermodynamically more stable structure of cesium oxo-fluoro-aluminates.

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Optimizing the extraction of parthenolide from feverfew: an experimental design approach

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Parthenolide (PAR), a sesquiterpene lactone isolated from the feverfew (*Tanacetum parthenium* (L.) Sch. Bip.), has gained significant attention due to its anticancer activity. Research has shown that PAR induces apoptosis in cancer cells and inhibits nuclear factor-kappa B (NF-κB) [1]. Feverfew contains a high amount of PAR, and its extraction from natural source is essential to preserving its full bioactivity.

The complex molecular structure of PAR poses significant challenges for synthesis, as it requires meticulous control over reaction conditions and stereochemistry. This often leads to numerous synthetic steps with low yields [2]. On the other hand, PAR extraction from natural sources is important, as it is generally more cost-effective, especially when improved extraction processes are applied through experimental design.

Experimental design is crucial in the green extraction process, optimizing procedures to maximize extraction efficiency while minimizing environmental impact. By carefully selecting and optimizing variables such as solvent type, temperature, and extraction time, green extractions aim to use environmentally friendly solvents, consume less energy, and generate less waste [3]. A promising category of solvents in this content is polymers, particularly Pluronics – block copolymers composed of ethylene oxide and propylene oxide. These amphiphilic molecules can self-assemble into micelles in aqueous solutions, effectively encapsulating hydrophobic compounds like PAR, thereby enhancing its solubility and stability in aqueous solutions. This feature is particularly beneficial for green extractions, as it reduces the need for hazardous organic solvents while maintaining high efficiency. Integrating Pluronics into extraction processes supports sustainable practices and development [4].

The current study's objectives were to ascertain the experimental variables that maximize extraction of PAR from feverfew by using the Box-Behnken experimental design and to assess the effect of temperature, time, and solid/liquid ratio. This study successfully optimized the extraction process of PAR from feverfew, demonstrating significant improvements in both efficiency and sustainability.

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Hydration and deliquescence behavior of calcium chloride hydrates

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Calcium chloride (CaCl₂) has gathered significant attention due to its presence in atmospheric aerosols [1], Martian soils [2], fluid inclusions [3] and applications in the storage and transformation of thermal energy [4]. Its various hydrates, their hygroscopic properties and the low eutectic temperature affect the water cycle on Earth and Mars and cloud condensation in the atmosphere [5,6]. Thermal engineering applications [4] include its use in sorption processes, e.g. for dehumidification, desiccant cooling, drying, refrigeration, water recovery from the atmosphere, the use as phase change material and for thermochemical heat storage.

A peculiarity of $CaCl_2 \cdot 6H_2O$ is that, depending on the experimental conditions, different deliquescence humidities (DRH) are observed if the salt is dehydrated and subsequently subjected to an increasing relative humidity [7]. Some measurements yield a higher value which is consistent with the commonly accepted DRH of 29.1 ± 0.4 % for $CaCl_2 \cdot 6H_2O$ based on the available water activities and solubilities [8,9]. In other measurements a lower DRH of about 19 % RH is observed. The lower value was assigned to the direct deliquescence of $CaCl_2 \cdot 2H_2O$ which did not undergo hydration to the hexahydrate state [7].

In this study, we used a consistent modeling approach to establish the phase diagram of the $CaCl_2$ – H_2O system including metastable equilibria (Fig. 1a). As can be derived from its metastable solubility, the DRH of the dihydrate is significantly lower than the observed DRH of 19 %. Therefore, we carried out a series of water vapor sorption, XRD and *in situ* Raman microscopy experiments to investigate the hydration and deliquescence behavior of $CaCl_2 \cdot xH_2O$. The results reveal two possible hydration sequences from lower hydrated states to deliquescence at 298.15 K (Fig. 1b): (1) Hydration of the monohydrate to the dihydrate, followed by the formation of metastable β - $CaCl_2 \cdot 4H_2O$, ending with its deliquescence at 18.5 % RH; (2) Hydration of the monohydrate to the dihydrate, followed by the formation of the stable polymorph α - $CaCl_2 \cdot 4H_2O$ and of the hexahydrate, ending with the deliquescence at 29 % RH. The two reaction pathways will be discussed in detail.

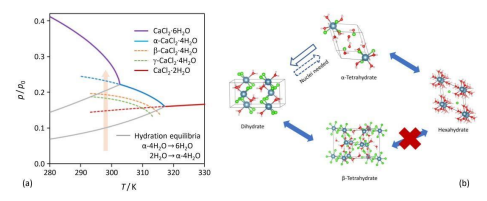


Figure 1. (a) Phase diagram, DRH (metastable dashed) and hydration equilibria (gray) (b) hydration—dehydration pathways

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Optimizing solubility of spirulina pigments using natural deep eutectic solvent

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This study focuses on isolating pigments from *Spirulina*, a blue-green microalga renowned for its rich nutritional profile and potent bioactive compounds. *Spirulina* is a source of phycocyanin, a vibrant blue pigment with strong antioxidant, anti-inflammatory, and potential anticancer properties, making it highly valuable in the food, pharmaceutical, and cosmetic industries. Traditional methods for extracting these pigments typically rely on organic solvents, which pose significant environmental concerns due to their toxicity and non-renewable nature [1,2].

The primary objective of this research is to extract *Spirulina* pigments using Natural Deep Eutectic Solvents (NADES) as eco-friendly alternatives to harmful organic solvents. NADES are a new generation of green solvents, composed of natural, biodegradable components such as choline chloride, betaine, glycerol, and sugars. These solvents are designed to meet the principles of green chemistry, offering low toxicity, biodegradability, and the ability to be tailored for specific extraction needs [3].

In this study, ten different NADESs were synthesized with a strong emphasis on optimizing solubility, a critical factor in maximizing the extraction efficiency of bioactive pigments. The antioxidant capacity of the extracted pigments was assessed using the DPPH test, a widely used method in chemistry for evaluating free radical scavenging activity. The DPPH test measures the ability of antioxidants to reduce the DPPH radical, a stable free radical with a deep violet colour, which fades upon reduction [4].

Among the NADESs tested, those based on betaine and glycerol demonstrated the highest solubility and antioxidant capacity in the extracted pigments from *Spirulina*. This research highlights the potential of NADES as a sustainable and effective solvent system for the extraction of valuable bioactive compounds from *Spirulina*, offering a greener alternative to conventional extraction methods [3].

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