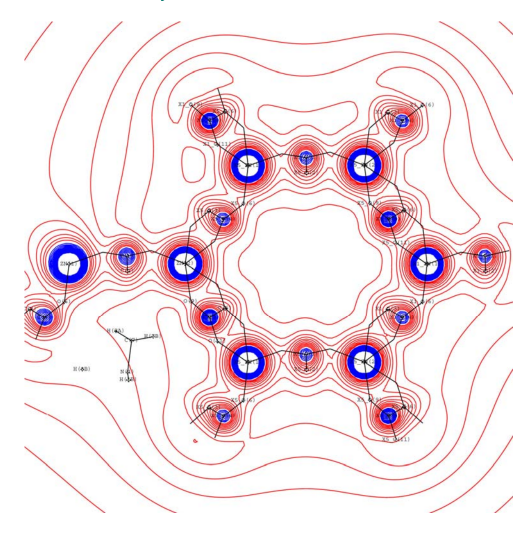


Schweizerische Gesellschaft für Kristallographie Société Suisse de Cristallographie Società Svizzera di Cristallografia Swiss Society for Crystallography

Sektion für Kristallwachstum und Kristalltechnologie Section de Croissance et Technologie des Cristaux

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SGK/SSCr NEWSLETTER

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Dear members of the Swiss crystallographic society,



Today, I would like to address a practical question as it concerns also our society: Online banking versus payment slips - or, to pay cash or not to pay cash – this is the question.

When arriving in Switzerland in 1998, I was surprised by the habit of many people to collect all payments to be done by the end of the month, and to pay all these bills cash at the post office. I adapted to this habit as, since I was a junior scientist, the sums I had to pay were not high. Nevertheless, I was surprised to hear people in front of me in the post office line pay sometimes thousands of Swiss Francs cash.

Today, the online banking makes these payments so much less risky, once you have the account opened and the secret codes to enter your account. Just like an email account. And, one avoids carrying around large sums of cash. Personally, I would like to encourage our members to profit from this opportunity to pay your annual fee. Why? Because the classical postal payments slip makes us loose money! The personal cash transaction at the post counter costs 5,- CHF which is taken off the sum of payment. With an annual membership fee which is guite low, this is a large percentage of the fee which we do not receive. According to our accounts, this seems to arise to more than 150,- CHF. This corresponds to one third of a travel grant which we give to young students!

Online banking is so much safer...

Enjoy the rest of the "summer"

Katharina Fromm

News for and from members

We welcome the following new members of the SGK/SSCr:

Personal members

Mr. Yvens Chérémond (Department of Chemistry, Université de Fribourg, 1700 Fribourg, PhD student)

Missing Members

I would like to thank all for helping me to find our missing members. Presently, our database is up-to-date.

Travel grants for young SGK/SSCr members

The committee will award the grants according to the following rules:

- Preference is given to PhD students
- Proof has to be given that there are no grants available covering the expenses
- A supporting letter by the supervisor of the applicant is necessary
- Applicant MUST be a member of our society

If you wish to apply for a travel grant, please send the above mentioned documents to the president of the SGK/SSCr anytime. You should have been member for at least one year before applying for a grant.

Travel grants are good opportunity allowing young scientist to profit from our society in a period with low income. By becoming afterwards a long-term member of our society, you can return this good-will later to the next generation.

Details for applications are given at:

http://www.sgk-sscr.ch/TravelGrants.pdf

Klaus Yvon, Université de Genève:

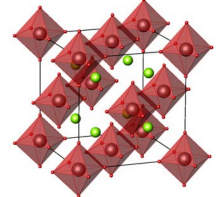
More than 40 Years of Structure Research in Materials Science

Contributed by Peter Fischer (Peter.Fischer@psi.ch)
Laboratory for Neutron Scattering (LNS), Paul Scherrer Institut



On the occasion of Klaus Yvon's retirement, it is an honor for me to congratulate him to his successful career as crystallographer and physicist. 1967 he obtained his master degree at the University of Vienna with structure-chemical investigations on carbides and carbohydrides of transition metals. As professor of the *Laboratoire de Cristallographie, Université de Genève*, his lectures on crystallo-

graphy and crystal chemistry and in particular attractive material science research



motivated over many decades young students to excellent diploma and master work under his supervision.

Thus more than 300 publications resulted. From this impressive list I would like to mention only a few examples illustrating common interests: ,LAZY PULVERIX' [1] got until now the record number of 2050 citations. At that time it had been one of the first general programs to easily calculate x-ray and neutron powder diffraction patterns and was also popular at our laboratory. Often cited (228) is furthermore the review

Fig. 1: Unit cell of Mg₂FeD₆ with Mg shown in green and with the characteristic D₆ octahedra around central Fe in red.

article on the crystal structures of high- T_c super-conductors for the years 1987-1988 [2]. It includes the common neutron investigation on the structural phase transition at 150 K in $La_{1.85}Sr_{0.15}CuO_4$.

Neutron diffraction cooperation in 1979 concerning the hexagonal Laves-phase deuteride $ZrMn_2D_3$ [3] received 101 citations. With high-pressure synthesis developed at the Geneva University new interesting hydrides such as Mg_2FeH_6 [4] were found; see the corresponding neutron diffraction result shown in Fig. 1. The hydrogen density of this compound exceeds the one of liquid hydrogen by a factor of approximately two. Maybe I should also mention our teamwork for a review on ternary metal hydrides published in 1988 [5]. With respect to mobile applications of hydrides Klaus tended later to lighter and more complex hydrides. Thus particular interest obtained the newer synchrotron x-ray work performed both at the Swiss-Norwegian beam line at ESRF and

at SLS by Klaus Yvon's research group on polycrystalline LiBH₄ [6]. Recently it had been to a large extent confirmed at SINQ by neutron scattering on LiBD₄ [7]. In this context we should particularly thank Klaus for his open-mindedness towards new research tools and his readiness to support corresponding efforts. An example is the financial contribution of the University of Geneva promoted by him already for the multidetector neutron powder diffractometer DMC at reactor Saphir. Moreover, he was and partially still is active in SGK/SSCr, SGN/SSDN, in research councils and as Swiss representative at ILL.

Current research topics of Klaus Yvon are metal hydrides, hydrogen technology, superconductors and ferromagnetic compounds, related to energy storage and conversion goals of MaNEP. In particular in the field of metal hydrides Klaus became a worldwide estimated expert and had been invited to many conferences including Gordon and MRS research meetings.

Klaus Yvon also aimed at practical applications of his research. Thus he realized an environmentally friendly lawn mower working already more than 14 years with hydrogen from metal hydrides. In addition he is involved in a privately owned photovoltaic production and storage installation in a one-family house, converting solar energy into hydrogen fuel for seasonal energy storage.

Ten of his former phD students and postdocs are presently in teaching positions as professors, H.F. Braun (Uni Bayreuth), Y.Filinchuk (Uni Louvain), M. François (Uni Nancy), R.Gladyshevskii (Uni L'viv), H.Kohlmann (Uni Nevada, Las Vegas), F.Kubel (Uni Wien), S. Y. MAO (Uni Xiamen), F.Mautner (TU Graz), M.Mendoza-Alvarez (Univ Puebla), P. Selvam (IIT Madras).

Because of all these merits Klaus Yvon has been honored by the triclinic, again hydrogen containing mineral $Cu(AsO_3OH)^{\cdot}2H_2O$ which now is called Yvonite. Congratulations from my colleagues from LNS and myself! Finally we would like to thank you, Klaus, for the good, stimulating cooperation over all the common years. Personally, I wish you a smooth transition into the 'stress state' of retired people, including further research activities –

Peter Fischer

- [1] K. Yvon, W. Jeitschko and E. Parthé, J. Appl. Cryst. **10**, 73 (1977).
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- [6] J. P. Soulie, G. Renaudin et al., J. Alloys Comp. **346**, 200 (2002).
- [7] F. Buchter, Z. Lodziana et al., Phys. Rev. B **78**, 094302 (2008).

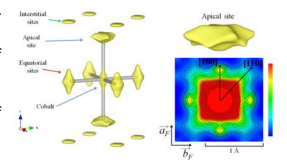
Recent Publications and Thesis Work Accomplished

Loïc Le Dreau, Thesis, Rennes July 2011

Phase transitions and oxygen ordering in $La_2CoO_{4+\delta}$ and (T, T')- La_2CuO_4 : single crystal growth and structural studies using synchrotron and neutron diffraction methods

Oxygen diffusion in this Ruddlesden Popper phases can not explained by a classical Arrhenius law. The latter involves activation of fast ionic diffusion at higher temperatures than what is observed in $La_2CoO_{4+\delta}$ (~430K) and thus do not provide satisfactory explanations of oxygen mobility at moderate temperatures in the Ruddlesden-Popper phases. Furthermore, the common concept of ionic diffusion depicts the phenomena in terms of random push-pull mechanism implying faster diffusion for higher structural disorders. However, in the present study, we clearly demonstrated that ordering plays a prominent role in the phase transitions undergone by oxygen-rich $La_2CoO_{4+\delta}$ phases.

Collaboration between the University of Rennes and LNS/SINQ@PSI, founded by CNRS (France) and ESM (Entwicklungsfonds Seltene Metalle, Zurich).



Nuclear density of CoO_6 octahedra and interstitial sites calculated by Maximum Entropy Method based on neutron diffraction data (TriCS@SINQ/PSI). The strongly anisotropic apical site is show on the right side in 3D (top), and below, the 2D projection on the a-b plane (bottom) of the density within $0.66 \le z/c \le 0.69$ corresponding to apical site.

(PS: Loic is presently looking for a Post-Doc position: loicledreau@hotmail.fr)

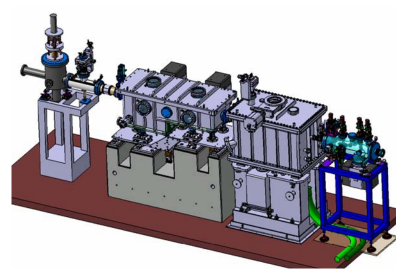
The new MS beamline at SLS/PSI: Building for a brighter future

Contributed by Ph.Willmott, A. Cervellino, D. Meister and M. Lange (SLS@PSI)

Welcome from the MS beamline! The Materials Science beamline was one of the first four to be built at the SLS and first saw light in 2001. The beamline was unique in being served by a 61-mm-period wiggler source (W61), which allowed access to photon energies as high as 40 keV. Since its inception, the MS beamline has provided high-quality synchrotron radiation for both powder diffraction and surface diffraction experiments.

In the meantime, undulator and storage ring technologies have made significant steps, especially with regards to magnetic materials and control of the electron beam. It is now possible to obtain similar fluxes to that provided by the W61 and a much improved brilliance using an undulator. It was therefore decided in 2008 that the MS beamline would undergo a comprehensive upgrade, involving both exchange to undulator radiation, and redesign of the optics to best exploit this.

The wiggler was removed at the end of October 2010, and replaced by a short-period (14 mm) in-vacuum, cryogenically cooled, permanent-magnet undulator, (CPMU, U14) at the start of 2011. The front end and optics have been completely redesigned to optimally exploit the characteristics of the U14 source and were installed in the first five months of 2011.



The optics hutch

The main components of the optics hutch are shown in Figure 1.

Figure 1: Components of the optics hutch. 1/right: FSX (Filters – Slits - X-ray quadrant BPM-1) box; 2: double-crystal monochromator (Cinel) and quadrant BPM-2; 3: Mirror chamber; 4/left: Si₃N₄ window and quadrant BPM-3.

The Cinel monochromator and double-mirror chamber were surveyed. The ultimate pressure in these optical components is less than 10^{-8} mbar and the first crystal in the DCM is not only cryogenically cooled, but in addition is surrounded by a cryogenic shield. It is hoped that these measures will mean that the first crystal's surface will remain clean over years, thereby facilitating experiments that exploit coherence, such as coherent diffractive imaging.

Figure 2: Michael Lange, Phil Willmott, Antonio Cervellino, Steven Leake, and Dominik Meister at the MS beamline on July 1, 2011, when the first monochromatic beam was detected in the experimental hutch. The beam is visible on the screen monitor as a white stripe.



First Results

In late June 2011, the undulator gap was closed to 6mm and beam was sought. Pink beam was found in the first experimental hutch on Tuesday 28th June, and monochromatic beam on Friday, July 1, 2011 (see Figure 2).

Gap scans are being performed presently in order to characterize the U14 spectrum. Focussing both vertically using mirror-2 and horizontally using the second DCM crystal were shown to operate effectively, and systematic studies will be carried out shortly (see Figure 3).



Figure 3: X-ray eye image of the U14 beam focused for the first time. The field of view is 6mm (H) x 2 mm (V). The unoptimized beam focus is estimated to have a size of 70 µm (H) x 35 цm (V).

The resolution of the DCM was tested at 9 keV. The theoretical Darwin width of Si(111) at this energy is 0.0017°, while the vertical beam divergence is 0.0018°. The FWHM of the convolution of these two yields a resolution of 0.0025°, which is exactly what was measured.

The energy was also calibrated in this energy range by recording an energy scan across the K-edge of copper. This was achieved simply by synchronized scans of the two DCM crystals. The relative angular positions of the two crystals remained optimized over this narrow angular range. The results are given in Figure 4.

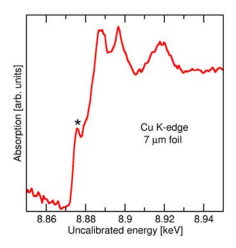


Figure 4: XANES spectrum of the Cu K-edge. The prefeature marked with an asterisk was measured at a Bragg angle of 12.870°, corresponding to an uncalibrated energy of 8.877 keV. The real energy of this feature is 8.982 keV, corresponding to a Bragg angle of 12.718°. The offset was thereby determined to be 0.162°.

Further energy calibrations using both EXAFS foils and Si powder diffraction patterns will be carried out in the next month.

Commissioning and user proposals

Optics commissioning will continue over the summer months. First pilot users are planned for September and a few experienced users will be contacted shortly. If you have experiments which you feel would be suited to the pilot-user phase, feel free to contact either Antonio Cervellino or Phil Willmott to discuss them. The MS beamline will accept proposals both in powder and surface diffraction for the regular proposal deadline of 15th September (call sent out on Aug. 1, 2011). By then, it should be clear whether we will be able to fill the entire first semester of 2012 with regular users, or whether some time will still need to be reserved for commissioning.

New techniques

In addition to the expected increase in performance of powder and surface diffraction at the MS beamline, we are planning to develop the following techniques:

- 1) microbeam Laue diffraction using the relatively broad photon energy range the high-K undulator in pink beam mode.
- 2) coherent diffraction imaging (CDI). This is particularly interesting for the MS beamline, as speckle around Bragg peaks far from the direct beam can be accessed using the surface diffractometer.
- 3) SAXS/WAXS. These techniques are expected to become invaluable adjuncts that can be used in parallel with more conventional powder and single crystal diffraction experiments.

All the above techniques will use the Mythen, Pilatus, and EIGER detectors, the next generation hybrid pixel detectors developed at the PSI. They will be developed by the MS-team but involvement and support by expert users would be highly appreciated.

Other news

After playing an integral and valued role in the MS beamline for the last ten years, Fabia Gozzo will leave us in September to start a new life with her family in Brussels. Everyone at the MS beamline would like to take this opportunity to thank Fabia for all her invaluable work over the last decade and to wish her success and happiness in the future. We are presently searching for a dynamic and talented scientist to replace Fabia as a tenure-track beamline scientist. Interested persons should apply to our advertisement.

A new introductory textbook on synchrotron radiation written by Phil Willmott has been published by Wiley in July 2011 (also available as a Kindle edition). The text is aimed at a broad audience, in particular nonphysicists unfamiliar with synchrotron techniques.



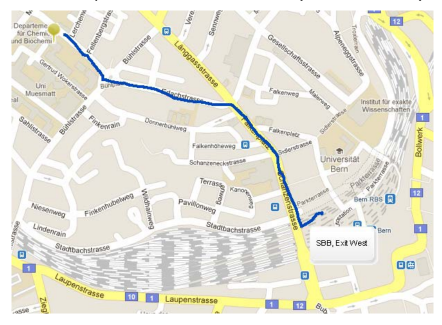
Abstracts and Program

The annual meeting of the SGK/SSCr and general assembly will take place at Department of Chemistry and Biochemistry, University of Bern, Freiestrasse 3, Bern on **Friday September 16, 2011**. The meeting is jointly organized by the groups of Mineralogical Crystallography (Prof. Dr. T. Armbruster) and Chemical Crystallography (PD Dr. P. Macchi) of the University of Bern.

A social dinner (Haus der Universität, http://www.hausderuniversitaet.ch/) is scheduled on Thursday September 15, 2011, 19h (evening before the meeting).

More details on the website: http://www.sgk2011.unibe.ch

Way from the SBB main station (15 minutes walk):



Registration (meeting and dinner): September 1, 2011

The meeting is free of charge (except for dinner and accommodation).

Meeting Title: Modern crystallographic techniques for new materials **Program Scientific Program (16 September, room UG113)**

9.00-10.00		registration /Poster session
10.00-10.05		Welcome message
10.05-10.50 [*]	IT1	at extreme conditions: Innovative techniques and new trends"
10.50-11.15		Coffee break /Poster
11.15-11.40 [*]	01	Session 1 (Inorganic materials and minerals) Pascal Schouwink: "Exciting Crystal Chemistry of Bimetallic Borohydrides"
11.40-12.05 [*]	O2	Ravi Sura: "Oxygen Ion Diffusion in $La_{2-x}SrxCuO_{4\pm\delta}$ Investigated by Oxygen Isotope Back Exchange and Diffraction Methods"
12.05-12.30 [*]	03	Volodymyr Svitlyk: "Influence of temperature and pressure on the crystal structure of the $A_x(Fe_{1-y}Se)_2$ (A = Cs, Rb, K) superconductors"
12.30-13.15		Lunch/Poster
13.15-13.45		General Assembly of the SGK
13.45-14.00		Coffee break /Poster
14.00-14.50	IT2	Dylan Jayatilaka (University of Western Australia): "Charge density refinement of X-ray diffraction data without the multipole model: what else can we squeeze out of the data?"
14.50-15.00		Poster Prize Winners announcement
15.00-15.25 [*]	04	Modeling structure and Properties of materials Riccarda Caputo: "Ab-initio crystal structure prediction"
15.25-15.50 [*]	O5	Anna Krawczuk-Pantula: "Atomic polarizabilities and multipolar model"
15.50-16.15 [*]	06	Bernard Spingler: "Some thoughts about the single crystal growth of small molecules"
16.15-16.45		Coffee Break/Poster
16.45-17.15 [*]	07	Facilities and new instruments Dmitry Chernyshov: "3D Mapping of Reciprocal Space with Synchrotron Light"
17.15-17.35*	08	Olha Sereda: "High Temperature in-situ XRD Studies on New Solder Materials for MEMS Packaging Applications"
17.35-17.55 [*]	09	Eric Hovestreydt:" Latest Results from CMOS detector technology"
17.55-18.00		Closing Ceremony

^{*} includes 5 minutes for discussion

List of Posters:

P01	Precursors for oxide materials
P02	Precursors for ion conducting battery oxide materials
P03	Optimizing the input parameters for powder charge flipping
P04	Behaviour of the ring silicate Benitoite BaTiSi ₃ O ₉ at high pressure
P05	Crystal structure of the uranyl mineral grimselite,
	$(K_{2.43}Na_{0.57})_{\Sigma 3.00}Na[(UO_2)(CO_3)_3](H_2O)$, from Jáchymov, Czech Republic
P06	Crystal structures and microstructures of ordered and disordered
	kalsilites. Is kaliophilite a KAlSiO ₄ polymorph or kalsilite polytype?
P07	Vibrational (FT-IR and Raman) spectral studies, geometrical parameters
	and theoretical calculations of triethylenetetramine and its cyano-bridged
	heteronuclear polymeric complex
P08	Hetero-metallic coordination polymers: Crystal structures of trans-[M(N-
	Meim) $2Ni($ 丟 $-CN)_4]_n$ (M=Cu(II), $Zn(II)$ and $Cd(II)$)
P09	Structure analysis of two open-framework zirconium phosphate materials
P10	The effect of hydration and strain on ceramic proton donductors
P11	Perfect epitaxial growth of germanium on silicon and crystal quality
	Quantification by high-resolution X-ray diffraction
P12	Stepwise in situ dehydration of yugawaralite: a temperature dependent
	single-crystal X-ray study
P13	Spin density distribution of the orbital singlet Mn(II): A polarised and
	non-polarised neutron diffraction study of the[Mn(Imz) ₆] ₂ +(NO ³⁻) ₂
	complex, where Imz = Imidazole"
P14	Intercluster Compounds for Nanosized Materials
P15	High pressure structure of $Mn_2(CO)_{10}$: an off-axis M-M bond
P16	On the purity of multi-layer focused X-ray radiation
P17	Properties of metal oxalate materials from electron density. Modelling the
	building blocks anPd evaluation of framework interaction
P18	Pavlovskyite $Ca_8(SiO_4)_2(Si_3O_{10})$ - a new mineral of altered silicate-
	carbonate xenoliths from the two Russian type localities: Birkhin massif,
	Baikal lake area and upper chegem caldera, northern Caucasus
P19	Lithium self-diffusion in LiAlSi ₂ O ₆ glass and single crystals
P20	Diffuse scattering study of β -pigment red 170
P21	α -"NaLuF ₄ ": 6 fold twinning with modulation and diffuse scattering
P22	Synthesis of new silver compounds with nicotinic acid derivatives

Meeting Sponsors











Abstracts of oral contributions

IT1: Single-crystal diffraction at extreme conditions - Innovative techniques and new trends

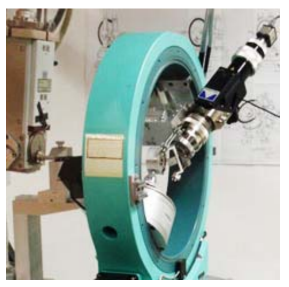
R. Miletich¹, T. Pippinger¹, D.G. Gatta², N. Rotiroti², M. Merlini², C. Hejny³

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The demand for obtaining knowledge on the structure-property relationships and hence the crystallography of solids at even more and more extreme conditions has lead to a continuous development in techniques applied to in-situ studies at non-ambient sample environments. In order to achieve even more extreme conditions it inevitably involves scaling down available sample sizes and thus giving a renaissance to single-crystal techniques. In particular modern high-pressure research nowadays aims for the multi-megabar pressure regime or the combination of variable pressure settings at e.g. high/low temperatures, electric/magnetic fields, or the sample being simultaneously exposed to any kind of irradiation.

Apart from persistent improvements on the sample environment [1], in particular new heating techniques such as mobile lasersystems [2] (see Fig.) or heatable gaskets [3], new diffractometer components and available sources allow higher resolutions in reciprocal space, a higher sensitivity for faint features (e.g. weak superstructure reflections), and the measurement on shorter time scales. Moreover, timeoptimized experiments now allow retrieve structural information even from samples, which are instable over time. Several examples of recent measurements on various phases will be provided in order to demonstrate the quality from state-ofthe-art experimental setups and the current limitations with available techniques.



High-pressure diamond-anvil cell and the mobile laserhead optics mounted on the χ -circle of the Huber four-circle diffractometer in Vienna

- [1] T. Pippinger, R. Miletich, M. Burchard, Rev. Sci. Instrum. (in press)
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 - N. Dubrovinskaia, J. Synchr. Rad. 16, (2009)
- [3] R. Miletich, D. Cinato, S. Johänntgen, High Press. Res. 29, 290 (2009)

- P. Schouwink¹, D. Ravensbæk², Vincenza d'Anna³, Radovan Cerny¹
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- ³ Département de chimie physique, Université de Genève

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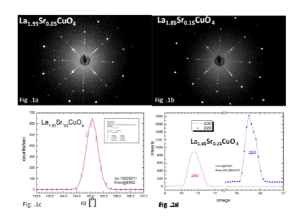
The fascinating diversity in the Crystal Chemistry of novel compounds based on the BH₄ group continues to proliferate. In this study we contribute with the structural characterization and classification of recently found novel ternary borohydrides, e.g. $K_2M(BH_4)_4$ (M=Mn, Mg, Zn) or KZn(BH₄)₃, based on bivalent Metals, which are brought into a crystal-chemical context and discussed in terms of possible differences concerning the bonding scheme. Currently planned studies on high-pressure polymorphs will also be presented. The examination of the present results is in good agreement with the established and ongoing trend of this compound group to form ionic as well as framework structures closely related to sulfates and oxides as well as halides in general, as observed in numerous recent works. It is shown by Synchrotron Powder Diffraction, backed up by spectroscopical methods, that structural similarities, e.g. between Mn- and Mg-modifications observed for binary borohydrides^[1,2] can be extended to corresponding ternary compounds. The compounds reported here all follow very similar building principles, the most prominent structural unit being isolated complex anions $[M(BH_4)_4]^{2-}$ or $[Zn(BH_4)_3]^-$, which are charge-compensated by the larger alkaline cation, resulting in packed structures derived from the Sulfates and Halides. Thus, while evidence for the covalent bonding contribution in the binary compounds of the respective M metal lies intrinsically in the connectivity of their distinct frameworks, the big complex anion in the ternary ones has various bonding-schemes as a consequence; the interaction between ligands and the central atom of $[M(BH_4)_x]^{n-}$ being of partially covalent character, as deduced by the directionality of hydrogen bonding reported in many works as well as the integrity of the tetrahedron, while on a larger scale the compound crystallizes in well known ionic structure types and their distorted derivatives. The ongoing reports of such structural similarities, culminating in solid solutions, not only extends the promising prospects concerning the precise compositional and structural tunability of this compound class regarding their hydrogen storage properties, but also is a clear hint of structural flexibility with regard to the more complex quaternary structures, which may show greater dependency on structural parameters such as site mixing and occupancies with respect to their physical properties, e.g. Li⁺ conductivity.

- [1] Ya. Filinchuk, R. Cerny, H. Hagemann, Chem. Mater, **21**, 925 (2009)
- [2] R. Cerny, N. Penin, H. Hagemann, Ya. Filinchuk,
 - J. Phys Chem. C, **113**, 9003 (2009)

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- 4 Chimie et Cristallochimie des Matériaux (C2M), Université de Montpellier, France

The understanding of oxygen ion conduction in solids at moderate temperatures is a key issue for the development of oxygen membranes in solid oxide fuel cells (SOFCs) [1-3]. The oxygen diffusion in $La_{2-x}Sr_xCuO_{4-\delta}$ can be decreased by Sr doping as reported by Opila [4]. Sr doping can therefore be used to investigate deeper how



oss of weight Temperature (°C)

Figure 1a, 1b: Back scattering Laue pictures of as grown single crystals. Figure 1c, 1d: High quality single crystal of La_{1.95}Sr_{0.05}CuO₄ and (200), (220) collected at ORION@SINQ.

Figure 2. Oxygen Isotope Back Exchange (OIBE) in La_{2-x}Sr_xCuO₄ showing the suppression of oxygen diffusion for high Sr doping (x).

diffusion and structure correlate. For this we need to examine high quality single crystal to avoid porosity and grain boundaries present in polycrystalline samples and influencing therefore the equilibrium. We have succeeded to grow $La_{2-x}Sr_xCuO_{4\pm\delta}$ single grain crystals by using a mirror furnace and tested them with X-ray Laue (Fig. 1a, 1b) and neutron diffraction (Fig. 1c, 1d). We did Oxygen Isotope Back Exchange (OIBE) experiments (Fig 2) between 40 °C to 1000 °C we proved that free oxygen mobility can be realized for x = 0, 0.05 already below 500 $^{\circ}$ C but is depressed for x = 0.10 and 0.15. This gives evidence that low temperature oxygen mobility relies on Sr doping in La₂CuO₄.

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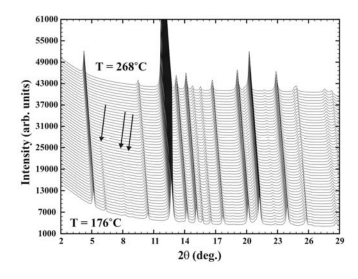
O3: Influence of temperature and pressure on the crystal structure of the $A_x(Fe_{1-y}Se)_2$ (A = Cs, Rb, K) superconductors

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Synchrotron radiation was used to study temperature $(Cs_x(Fe_{1-y}Se)_2)$ and pressure $(Cs_x(Fe_{1-y}Se)_2)$, $Rb_x(Fe_{1-y}Se)_2$ and $K_x(Fe_{1-y}Se)_2$ evolution of crystal structures of the new layered Fe-based superconductors.

Cs_{0.83}(Fe_{0.86}Se)₂ undergoes a firstorder structural trans-formation on heating associated with the disorder of the iron vacancies in the FeSe layer, as we have shown before [1] and in agreement with



other reports [2]. Irreversibility of the temperature dependence of the lattice dimensions and Bragg intensities is most probably related to a mobility of the intercalated alkali ions. First order of the transition is manifested by the hysteresis in the lattice behavior and also confirmed by the differential scanning calorimentry [1].

No clear pressure-dependent anomalies for $Cs_{0.83}(Fe_{1-y}Se)_2$, $Rb_{0.85}(Fe_{1-y}Se)_2$ and $K_{0.8}(Fe_{1-y}Se)_2$ ($y \sim 0.14$) were observed around 80 kbar, where transport experiments indicate an offset of superconductivity [3]. The Bragg reflections indicative of the vacancies ordering in the FeSe layer do not disappear up to applied pressures of ~120 kbar. This indicates that the ordering of the vacancies in the FeSe layer may be present in both superconducting and non-superconducting states.

Thermal expansion coefficients calculated from the temperature dependent diffraction and bulk moduli obtained from the pressure dependent diffraction may serve as constrains for theoretical models.

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Charge density refinement of X-ray diffraction data without the multipole model: what else can we squeeze out of the data?

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The philosophy and practice of deriving a reasonable single-determinant wavefunction from experimental charge density X-ray diffraction data will be reviewed.

This hybrid "experimental" wavefunction can be used to produce many properties not normally associated with the X-ray charge density technique; for example, electron localization functions (ELF) and Roby bond indices. The message here is that charge density measurements can be used to do far more.

As well as improved qualitative chemical information, results from these "experimental" wavefunctions are presented for some linear and non-linear optical (NLO) properties, namely refractive indices and second order NLO coefficients.

The results from the "experimental" wavefunctions are in better agreement with experimental measurements than those from wavefunctions where no X-ray data is used as input. They are also in better agreement with multi-determinant wavefunction calculations.

As further evidence of the usefulness of wavefunctions which incorporate experimental data, results for geometrical parameters and ADPs for a small didpeptide gly-ala will be presented and compared with corresponding neutron data.

The agreement between such measurements, particularly for hydrogen atoms, has been previously thought to be unachievable from X-ray diffraction data alone.

O4: Ab-initio crystal structure prediction

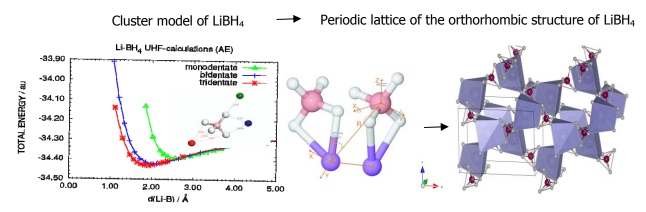
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Ab-initio crystal structure prediction is still one of the most challenging and interesting issue in condensed matter science. The first difficult task is the determination of the irreducible descriptors required to search for the global minima among different conformational geometries, a molecular system can adopt when visiting the many local minima of the potential energy surface.

By presenting some case studies, it will be discussed an ab-initio computational methodology of crystal structure modeling, which combines cluster modeling, via global minimization techniques, and periodic lattice calculations, based on Density Functional Theory. Whereas known experimental structure are available, the comparison with them helps to validate the predictability power of the computational approach and provides a molecular insight of stable and metastable structures of the particular system under study. The examples are represented by the first-principles ground-state structure determination of selected complex borohydrides, metal borides and transition metal oxides of interest in materials science for energy applications.



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O5: Atomic polarizabilities and multipolar model

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When a molecule is placed in an external electric field, its electron distribution changes in response. A useful measure of this effect is the molecular dipole moment together with its first and second derivatives (polarizability, hyperpolarizability). Using the QTAIM theory [1] it is possible to decompose the dipole moments and the polarizabilities into the atomic components. It was previously discussed [2] that each atomic contribution can be expressed as the sum of "atomic polarization" and "charge transfer" contributions. The transfer contribution is given by the weighted translation vector from atom's nucleus to the corresponding BCP, its sign and value strongly depend on the nature and number of bonded groups to the selected atom.

Here we present the relationship between the charge density distribution, obtained with the use of multipole model of Hansen and Coppens [3], and electric properties such as electric dipole moment, as well as the response of a molecule to the external electric field in simple organic molecules. It is well established that the multipolar reconstructed electron density (XD2006 [4]) can provide sufficiently accurate electric moments of a molecule in the crystal, however our results show that this does not hold for derivatives of the electric moments. In particular, while the bond polarity term is sufficiently well reconstructed (at least at zero field), the atomic polarization term is more problematic, even without the application of an external electric field. Empirical methods to estimate the polarizability based on the zero field electron density distribution will be tested, in order to allow estimation of linear optic properties from a multipolar model.

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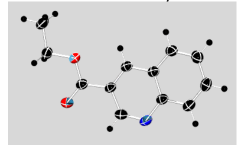
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Crystallisation of suitable single crystals undoubtedly is the most decisive step of a successful X-ray structure determination. Quite often, only an X-ray analysis will definitively reveal the composition and three-dimensional arrangement of an unknown compound. However, this powerful technique fully relies upon the ability to grow single crystals of sufficient quality and size. Although crystallisation is an important purification technique which is taught and widely used in every Chemistry undergraduate curriculum, different techniques may have to be employed for the growth of diffraction-quality single crystals.

One of the useful descriptions about growing single crystals is that of Jones.[1] Kroon and co-worker have summarized the different techniques for growing crystals of organic molecules, though their methods can be extended to many small molecules.[2] Hulliger has described in great detail techniques that are suitable for various types of compounds.[3] On the internet, many pages are dedicated to the description of how to grow single crystals [4, 5]. The crystallisation of active pharmaceutical ingredients has been reviewed.[6] Not surprisingly, several publications have studied the influence of solvents upon the crystallisation of polymorphs.[7, 8] Of course, the use of different solvents often results in solvate crystals as shown for organic crystals[9], metal complexes[10], or pharmaceuticals.[11-13] Since the number of possible solvents is very large, solvents were grouped after statistical analysis of selected solvent parameters.[14-16] However, the knowledge gained from these investigations was not used to tackle an open crystallisation challenge.

Nevertheless, it has been our experience in the last fifteen years, that beginners are often lacking the knowledge of how to setup crystallisation trials. Secondly, even experienced researchers are grateful for hints how to optimize crystallisation setups that hitherto have failed to produce suitable single crystals. We have assumed for the following studies that the compound of interest is available in small amounts of not more than 50 milligrams only. This quite common situation has severe consequences: a quantitative determination of the solubility in common solvents is not possible. Furthermore, some crystallisation techniques cannot be used, because they require more material and they do not allow the recovery of the compound of interest.



The presentation has a two-fold aim: For beginners in the field of single crystal growth, we want to describe and critically compare the most commonly used basic techniques for the growth of single crystals: a) vapour diffusion (including solvent evaporation) and b) solvent layering. We briefly also discuss the unconventional technique of crystal growth in gels. At the same time, we want to give the more advanced

researchers some ideas, in case their standard methods fail to work. In particular, we will show that a careful screening of the solvents, including some uncommon ones, increases the chances of getting single crystals. This is demonstrated by the example of

3-carbethoxyquinoline (**1**, see ORTEP representation of one out of two molecules in the asymmetric unit). In addition, we will share our experience about various aspects of crystallisation that we have accumulated over the last fifteen years.

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In a diffraction experiment correlated static or dynamic disorder manifest itself in structured diffuse scattering. Thanks to the availability of bright synchrotron radiation and fast area detectors with high dynamic range, diffuse intensity can readily be observed even for weakly scattering compounds. Moreover, for many cases diffuse intensity can be represented in form of 3D distributions in reciprocal space. These distributions can be now measured with a high level of sensitivity and in a relatively short time. As illustrations, we show examples of reciprocal mapping for the Kohn anomaly in zinc, chemical disorder in a Prussian Blue analogue and a perovskite, thermal diffuse scattering in quartz, in an Invar alloy, and in hexagonal Ice, as well as diffuse scattering in a relaxor ferroelectric.

We will also present a project of new diffractometer, to be installed at beamline BM1A (SNBL at ESRF), that combines a fast pixel detector PILATUS 2M with a flexible 3-axis kappa goniometer. Funding for this instrument has now been received from the Swiss National Science Foundation and the Norwegian Research Council. The diffractometer is designed as a fast mapper of reciprocal space; it will allow us to combine accurate and complete Bragg data with a full sphere 3D map of diffuse scattering measured within a few minutes. Some basic parameters of this diffractometer will be given, and possible modes of data collection will be discussed.

O8: High Temperature in-situ XRD Studies on New Solder Materials for MEMS Packaging Applications

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In the fabrication of Micro Electro Mechanical Systems (MEMS), the application of advanced packaging methods involves the development of new solder materials with improved properties [1]. The joining of advanced metallic alloys is limited by high brazing or soldering temperatures. The nanoscale effect of melting point depression (MPD) offers a possibility of lowering the filler metal melting temperature without the compromises resulting from alloying [2].

In-situ high temperature XRD studies have been conducted in order to follow the melting behavior of mixed metal nanolayer systems (Figure 1) and to have a direct observation not only on the materials melting points but also on the formation of intermetallic phases. The influence of internal and external interfaces in the material is crucial and has also been studied.

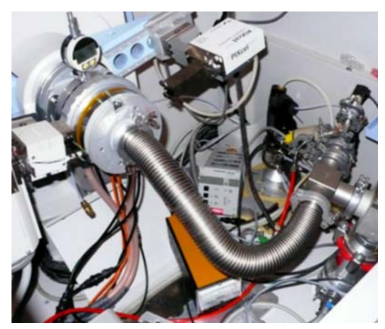


Figure 1.

The MPD effect has been studied for nanolayer brazing fillers in a multilayer system in which the layers of the brazing filler are separated by diffusion barrier layers (DBLs). The example of the AgCu/AlN nano-layer system is shown in Figure 2. The analysis shows that for the 7.7 nm thick AgCu layers, a decrease of 25°C for the melting point was observed for the AgCu/AlN nanolayer configuration. In this case, the eutectic AgCu brazing filler separates in two single phases and the local material interfaces Ag/AlN, Cu/AlN are formed.

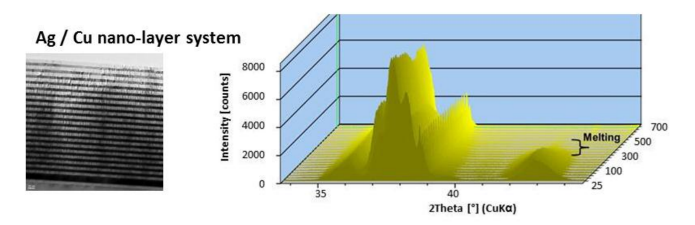


Figure 2.

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O9: Latest Results from CMOS detector technology

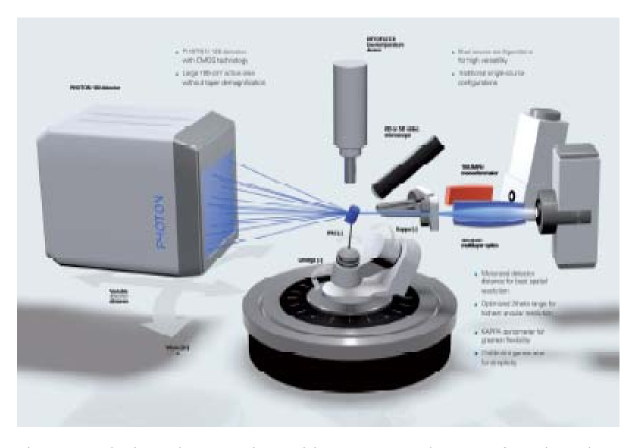
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Detectors for crystallographic applications require a unique combination of sensitivity, speed, dynamic range, resolution and detector size. CMOS (Complementary Metal Oxide Sensor) technology has shown great potential to provide sensors for the next detector platform of choice and has incited a significant wave of advancement in high-performance crystallographic imaging. CMOS technology provides large high-performance sensors that were previously not achievable with CCD manufacturing processes.



Results, as recently obtained in our application laboratory, on a wide variety of samples and applications, from high-pressure to charge density measurements will be discussed

Abstracts of Posters

P01: Precursors for oxide materials

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Heterometallic compounds can be used in different applications, for instance as precursors for photo-electronic devices or mixed metal oxides in high- T_c superconductor (e.g. YBCO) [1]. A synthetic strategy to obtain mixed metal oxides is to use decomposition techniques like (metal-organic) chemical vapour deposition (MO-CVD) to combust metal-containing volatile complexes. For the drastic decomposition process, it is suggested to use simple ligand systems which can easily be prepared in large scale. Complexes should then be easily obtained.

In previous work [1], a mixed metal complex of barium and copper was obtained. To obtain a tri-metallic complex with barium, copper and yttrium, the idea of a co-ligand for yttrium came to mind. The beta-diketonates seems to be good co-ligand because yttrium is a hard ion and this kind of ligand are also hard (even more when deprotonated) [figure 1].

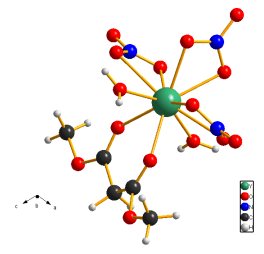


Figure 1: Yttrium complex with dimethylmalonate

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Ion conducting materials are of increasing interest predominantly due to the commercial interest in solid oxide fuel cells and Li batteries. The layered oxide phases of LiM_xO_y (M = Co, Ni, Mn, Fe) are often used as cathode materials for these batteries [1-3]. While the search for new materials continues to attract considerable attention there has been a noticeable increase in research of new precursors for those materials. Metal alkoxides have been proposed like good precursors due to their high solubility, low decomposition temperatures, cross linking ability, ease of modification and commercial availability [1, 4, 6]. Moreover, recent research [5] showed the use of binary iron/lithium organometallic complexes as single source precursors to solid state cathode materials for potential Li-Ion Battery applications.

For this purpose, we developed new metal alkoxides (e. g. Fig.1) using the halides of iron, nickel, cobalt, and manganese as starting materials. We will present the first results of this work using the Schlenk techniques to synthesize those precursors, which can come as molecular species or as coordination polymer networks (or metal organic frameworks, MOFs).

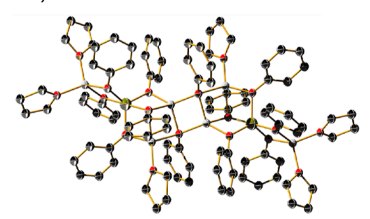


Figure 1: Single Fe-Li-compound with structure formula: ((THF)₆Li₆Fe₂(OPh)₁₀)₄

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A variety of structures have been solved using the Powder Charge Flipping algorithm [1, 2] incorporated in program *Superflip* [3, 4]. But, as the program is relatively new, the selection of the values for the input parameters has been rather arbitrary. Therefore, a systematic study answering the questions about the effect of the different input parameters on structure solution; minimum data resolution that can be used and use of the information obtained from the other sources was undertaken.

The data collected for one inorganic (1) and one organic (2) compound were used for these tests.

- (1) The original solution (*Superflip*) of zirconium phosphate structure did not have the correct symmetry (P-1), and a few of the atoms in Zr-O-P layers and the quinolinium ions between the layers were missing [5]. Combination of input parameters that deal with: defining the groups of overlapped reflections, repartitioning the overlapped reflections, and the isotropic displacement factor were tested within a sensible range, and an optimal parameter set could be established. With this set, a complete layer structure exhibiting the correct symmetry was produced.
- (2) For ribose [6], extensive tests on *Superflip* input parameters did not result in a satisfactory structure solution. Reasoning that starting with non-random phases might help, chemically feasible structures were generated using the direct-space program FOX [7]. None of these were correct. But, using their phases as seeds for generating phase sets in *Superflip* resulted in slightly better solutions. It was only when threshold for flipping was reduced, and establishing the definition of the groups of the overlapped reflections was reoptimized that fully interpretable solutions with the correct symmetry were obtained.

In both cases, the input parameter set was reoptimized for the lower resolution data. For the zirconium phosphate, the structure could be solved even with 2.4 Å resolution data, and the quality of the solutions was insensitive to the input parameters. For ribose, complete solutions were obtained down to the resolution of 1.18 Å when the input parameters were reoptimized.

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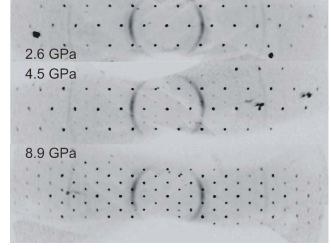
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The mineral benitoite, $BaTiSi_3O_9$, is remarkable in many respects: Not only is it a well-priced gemstone with strong pleochroism, but it is also used in electron microprobe laboratories as standard material and for alignment of the electron beam due to its fluorescence. The basis of the structure is a mixed framework of $\{Ti[Si_3O_9]\}^{2^-}$ composed of discrete Ti octahedra and $[Si_3O_9]$ triple rings. Ba atoms lie in columns between the triple rings giving the overall symmetry of the very rare space group P-6c2 1,2 .

In-situ high-pressure single-crystal diffraction of BaTiSi $_3$ O $_9$ shows additional Bragg peaks above 3.70 GPa indicating a non-isomorphous transition from P-6c2 (no.188) to P31c (no.159) space group symmetry with a'=a $_3$.

Fig 1: Reconstructed *hk4* reciprocal layer in Benitoite, BaTiSi₃O₉ with increasing pressure.



The crystal structure investigations reveal the off-centre displacement of the Ba atoms on release of symmetry constraint to be responsible for the high compressibility of the high-pressure form. The triple ring in the Benitoite structure loses the mirror symmetry resulting in an increasing deviation from 90° of the angle spanning between the connecting line between the terminal oxygen atoms of the silicon tetrahedra and the central ring plane from 0° at the phase transition to a maximum of 15° at 8,88 GPa. Similar behaviour with pressure has also been observed in closely related $K_2ZrSi_3O_9$, wadeite, as well as other cyclosilicates with triple rings of silicate tetrahedra³.

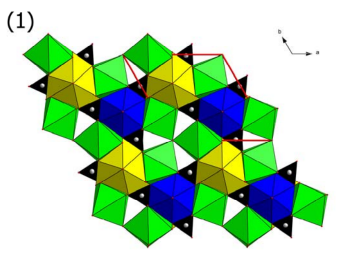
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P05: Crystal structure of the uranyl mineral grimselite, $(K_{2,43}Na_{0.57})_{53,00}Na[(UO_2)(CO_3)_3](H_2O)$, from Jáchymov, Czech Republic

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The crystal chemistry of uranium compounds is of high importance due environmental impacts to connected with the long-term underground depositories of "spent nuclear fuel" (SNF) [1]. Despite of considerable improvement of our knowledge of crystal chemistry of U⁶⁺, studies carried out on natural minerals are important. Here we present the refinement of the crystal structure of the uranyl mineral



grimselite, determined for the first time by single-crystal X-ray diffraction on a natural specimen, found at Jáchymov, Western Bohemia, Czech Republic. The crystal structure of the synthetic analogue is known [2]. Grimselite was firstly reported from a cable tunnel between Gerstenegg and Sommerloch in the Grimsel area, Aarmassif, Switzerland [3].

The starting model of the synthetic analogue [2] refined with JANA2006 [4] to R_1 = 0.0082, $wR_1 = 0.0185$ with GOF = 1.33. Natural grimselite is hexagonal, of the space group P-62c, a = 6.7050(3), c = 13.2857(5) Å, V = 595.74(3) Å; the structural formula derived from the refinement is $(K_{2,43}Na_{0.57})_{53.00}Na[(UO_2)(CO_3)_3](H_2O)$, Z = 2. Crystal structure of natural grimselite is built upon heteropolyhedral sheets (1), consisting of uranyl hexagonal bipyramids (blue), linked by trigonal planar CO₃²⁻ groups (black) to NaO₈ (yellow) and (K, Na)O₈ (green) polyhedra. In contrast to the synthetic analogue, natural grimselite has a mixed Na, K occupation of KO₈ polyhedra. Results of the structure refinement are supported by chemical analysis (EPMA-WDS) and vibration spectroscopy (IR and Raman).

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P06: Crystal structures and microstructures of ordered and disordered kalsilites. Is kaliophilite a KAlSiO₄ polymorph or kalsilite polytype?

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Despite of numerous investigations using various experimental techniques, the problem of polymorphism in feldspathoid KAISiO₄ is still not solved. Recent discovery of a new polymorph called magakalsilite [1] or investigations of possible phase transitions in kalsilite [2] significant steps in the right direction. However, existing results about the KAISiO₄ system are extremely complex and often controversial.

Literature suggests that a completely satisfying structure model of kalsilite does not exist and that the crystal structure of kaliophilite is still unknown.

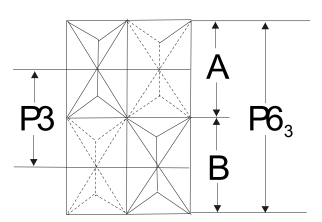


Fig.1 Schematic representation of two types of layers in disordered kalsilite in P63 and P3 groups

As a part of on-going research in this system, we have investigated two X-ray powder diffraction patterns that correspond to ordered and disordered states of kalsilite (KAlSiO₄). Rietveld refinement of both experimental patterns in two different space groups $P6_3$ and P3, inspection of electron density maps and microstructural characteristics revealed disordered layers of type A and B (Fig1) i.e. stacking faults characteristics of disordered kalsilite. Based on the unit cell values and analysis of the layer disorder, we postulate that the here described disordered kalsilite structure corresponds to synthetic kaliophilite. Therefore, kaliophilite is not a KAlSiO₄ polymorph but a kalsilite polytype characterized by random disorder for A and B layers stacked along the *c*-axis.

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P07: Vibrational (FT-IR and Raman) spectral studies, geometrical parameters and theoretical calculations of triethylenetetramine and its cyano-bridged heteronuclear polymeric complex

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In our previous studies [1, 2], the experimental results of the vibrational spectroscopic, thermal and structural analysis of the cyano-bridged heteronuclear polymeric $[Cd(teta)Ni(\mu-CN)_2(CN)_2]\cdot 2H_2O$ and $[Zn(teta)Ni(\mu-CN)_2(CN)_2]_n$ investigated. In this study we report on the molecular geometry and vibrational frequencies of triethylenetetramine (teta) and its cyano-bridged heteronuclear polymeric complex, $[Zn(teta)Ni(\mu-CN)_2(CN)_2]_n$. The molecular geometry and vibrational frequencies of triethylenetetramine (teta) in the ground state has been calculated using the density functional method (B3LYP) with 6-311++G(d,p) basis set. The optimized geometric bond lengths and bond angles were obtained by DFT (B3LYP) had provided the best agreement with experimental data. Furthermore, we attempt a theoretical investigation of the metal-ligand vibrational modes, which will be discussed in detail in the second part of the study. The molecular geometry was optimized from Density Functional Theory (B3LYP method) using LANL2DZ basis set on the ground state and the infrared wavenumbers and intensities were predicted by using this geometry. Calculated wavenumbers and intensities were compared with FT-IR and Raman spectra of triethylenetetramine and its cyano-bridged heteronuclear polymeric complex. Additionally, time depended density functional theory (TD-DFT) method using 6-311++G(d,p) basis set was used to determine the minimum energy structure of the complex. According to the calculated results, the vibrational wavenumbers, geometric parameters and excitation energies show an excellent agreement with the experimental data.

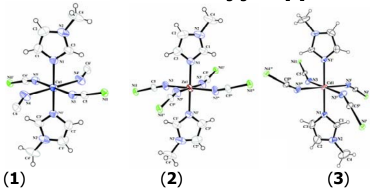
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Cyano complexes were among the first coordination compounds ever prepared, studied, described and used [1, 2] and they are still under investigation in coordination and organometallic chemistry at present [3]. These compounds are interesting as ion exchangers, molecular sieves or materials for storing gases [4].



In this study, three cyano bridged heterometallic complexes of general formula, trans- $[M(N-Meim)_2Ni(\mu-CN)_4]_n$ (N-Meim = N-methylimidazole and M = Cu(II), Zn(II) and Cd(II)) have been synthesized. The crystallographic analyses reveal that the complexes, $[Cu(N-Meim)_2Ni(\mu-CN)_4]_n$ (1), $[Zn(N-Meim)_2Ni(\mu-CN)_4]_n$ (2) and $[Cd(N-Meim)_2Ni(\mu-CN)_4]_n$ $CN)_4]_n$ (3), have polymeric 2D networks. In the complexes, four cyanide groups of [Ni(CN)₄]²⁻ coordinated to the adjacent M(II) ions and distorted octahedral geometries of complexes are completed by two nitrogen atoms of trans *N-Meim* ligands. Although the literature lists many reports on the binding modes of tetracyanonickelate(II) complexes, from the X-ray crystal structure point of view only a limited number of structures with corrugated and four cyanide-bridged polymeric networks have been obtained until now. The Cu(II) and Zn(II) complexes are the first examples showing coordinated 2D layered structures with four cyanides.

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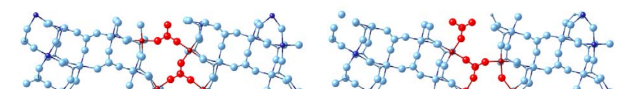
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The structures of the ZrPOF materials synthesized in the presence of ethylamine (EA) and diethylamine (DEA) have been analyzed. As the synthesis conditions and diffraction patterns were similar, few structural differences were expected. However, while the ZrPOF-EA structure is solved and refined [1], the ZrPOF-DEA structure displays some ambiguities.

The powder pattern of ZrPOF-DEA could be indexed with an orthorhombic unit cell comparable to that of ZrPOF-EA. An initial structural model was derived using the powder charge flipping algorithm [2], implemented in the program Superflip [3]. Rietveld refinement revealed that ZrPOF-DEA has adopted the symmetry $Pmc2_1$ (a = 6.623, b = 37.171 and c = 20.033 Å) whereas ZrPOF-EA crystallizes in the higher space group Pbam (a = 19.957, b = 37.067 and c = 6.617 Å). Thus, ZrPOF-DEA contains a total of 124 framework atoms (16 Zr, 24 P, 84 O/F) in the asymmetric unit, which is double that of ZrPOF-EA. Even though the two structures display common features, the refinement of ZrPOF-DEA has not yet been successfully completed. Only one of the diethylamine ions has been located and there appears to be some additional disorder in the framework that still needs to be addressed.



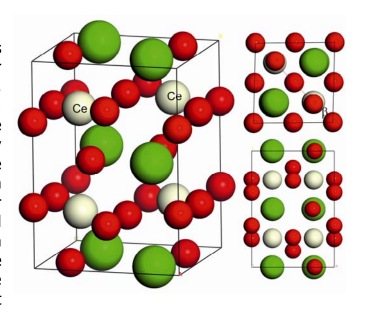
The model obtained for ZrPOF-DEA from charge flipping (left) deviates slightly from the ZrPOF-EA model (right). The differences in connectivity are indicated in red.

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The proton conducting perovskites are extensively studied for their potential use in fuel cells, electrolysers and gas sensors. Recent Empa data show that the proton transport activation energy scales linear with the lattice constant, suggesting that enlarged lattice volume example thin films with epitaxial can promote proton conductivity. To comprehend the interaction of water with the oxide lattice, and the underlying transport



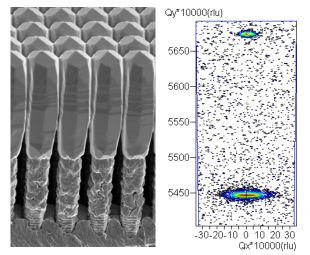
mechanisms, we combine electrochemical transport measurements *in-situ* with synchrotron and neutron spectroscopy and scattering methods. Optical Raman spectroscopy and XRD studies on the compressively strained $BaCe_{0.8}Y_{0.2}O_{3-\delta}$ proton conductors indicated that the proton transport may be anisotropic. The effect of hydration was studied by *in-situ* XPS, where the reactions of waster on its surface were revealed.

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The epitaxial growth of germanium on silicon allows for the integration of germanium into silicon CMOS technology, availing in standard ICT devices the superior properties of germanium as high electron density and high electron mobility. Nevertheless, layer stresses have been observed being responsible for wafer bowing and in some cases layer cracking. The observed layer stress is a result of the lattice mismatch between the germanium and the silicon and the high mismatch of thermal expansion coefficients of germanium and silicon [1].



A new approach for the germanium deposition on silicon has been introduced with the use of a patterned silicon substrate. This innovative approach results in a complete relaxation of the germanium being epitaxially grown on the Si pillar substrate. The threading dislocation density is also reduced to close to zero and a perfect crystal quality is obtained despite the large lattice mismatch. No layer cracking and a minimized wafer bowing is observed which presents an innovative key technology to wafer handling and post-growth processing.

Fig. 1. Left: SEM view of 8 μm Ge towers on Si pillars. The high crystal quality is apparent by the growth along the crystal planes and is confirmed and quantified by HRXRD angular scans and RSMs. Right: RSM of the (004) reflections of Si (top) and Ge (bottom). The FWHM of the Germanium (004) ω/2θ-reflection is 16 arcsec.

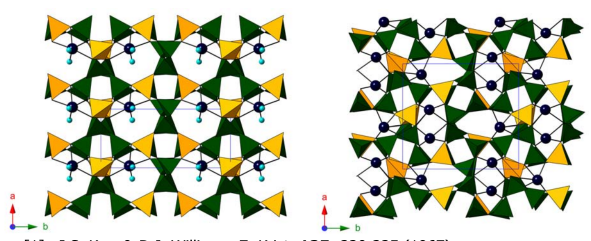
The crystalline quality was investigated by High Resolution X-Ray Diffraction (HRXRD) using symmetrical (004) and asymmetrical (224) reflections and measuring angular scans and Reciprocal Space Maps (RSMs). These measurements allowed for a quantification of the crystal mismatch, the layer tilt and the degree of relaxation. It was shown that layers up to 50 µm thicknesses could be grown showing 100% relaxed and perfectly parallel germanium layers with respect to the silicon substrate crystal.

This is a large step towards the integration of germanium into the silicon technology which is particularly interesting for future solid-state X-ray detectors where germanium is highly superior to silicon having an X-ray absorption coefficient being ~25 higher than silicon.

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- P12: Stepwise in situ dehydration of yugawaralite: a temperature dependent singlecrystal X-ray study
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Yugawaralite CaAl₂Si₆O₁₆·4H₂O, space group *P*c, is a natural zeolite. The tetrahedral framework ([1] -[4]) consists of well ordered Si (green) and Al (yellow) tetrahedra (Figs.). At room temperature 8-coordinated Ca is located at the intersection of the channels and is bound to four framework O and four H₂O molecules. Single-crystal dehydration was performed under a dry nitrogen atmosphere in steps of 25°C up to 325°C. According to structure refinement yugawaralite looses 1H₂O pfu below 50°C triggering a small change in the until cell parameters. At 300 °C Ca is reduced to sixfold coordination, bonding to W1 and W2 and four framework O with little change in the cell parameters (left Fig.). Above 300°C dehydration proceeds resulting in a decrease of coordination of Ca, which now bonds to five framework O [5, 6]. The structure transforms to space group *P*n resulting in a doubling of the *a*-axis (right Fig.) leading to complicated intergrowth and twinning. The partially occupied Ca sites compress the channels yielding a 7.11% reduction of (normalized) volume.



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P13: Spin Density Distribution of the orbital singlet Mn(II):

"A polarised and non-polarised neutron diffraction study of the
[Mn(Imz)₆]²⁺(NO₃-)₂ complex, where Imz = Imidazole"

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The degree of covalency of the Mn – N bond in manganese(II)-histidine interactions, has been of interest due to the vital role manganese(II) play in biological systems. In this work, polarised neutron diffraction (PND) and unpolarised neutron diffraction, multipolar least square analysis (MPLSQ), Density Functional Theory (DFT) calculations and assessment, will be used to understand the Mn2+ spin density distribution in the complex [Mn(Imz)₆]²⁺(NO₃-)₂. Firstly, within the trigonal unit cell for the complex, while correlating these results to the local anisotropic axis of the unit cell. And, lastly, using these results in understanding the spin density distribution, and n-anisotropy, effect about the Mn – N bond.

For the Mn(II) ion, where the orbital angular momentum L=0, and which is represented by the 6A1g ground state, the spin density distribution of the unpaired delectrons is defined by a superposition of one-center density population orbital product functions. Multipolar expansion provides a representation of these orbital functions in the spin density form factor description.

Through multipolar expansion computation,³ the spin density structure factors, derived from the flipping ratios of polarised neutron diffraction, can be correlated to the population of respective multipoles, and thus orbitals. We have performed a PND experiment on the D3, and a non-polarised neutron diffraction experiment on the D9, instruments at the ILL in France, on a $2\times2.5\times5$ mm3 (40 mg) crystal of the deuterated [Mn(Imz)₆]₂+(NO₃-)₂ complex. We have also performed DFT calculations and modelling on this system.

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The term "intercluster compound" refers to assemblies that are constituted of at least two different large, inorganic building blocks [1, 2]. A few intercluster compounds already exist and it has been possible to determine their crystal structures by single crystal X-ray diffraction.

These results deliver insights into the arrangement of large building units in the solidstate material and the underlying intermolecular forces [3, 4].

These first results show that intercluster compounds are very promising for the study of structural and physical properties of nanosized particles.

They also open a wide range for new applications (lithography techniques, electronic or optical devices, study of quantum effects and also of orbital bands). Since only a few compounds have been studied, there is a huge potential for new discovery in this field. We would like to present new building blocks [5] and their corresponding intercluster compounds.

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P015: High pressure structure of $Mn_2(CO)_{10}$: an off-axis M-M bond

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The bond between transition metals has always attracted structural chemists, because of the intriguing features that characterize this interaction. In particular, Mn-Mn bond in $Mn_2(CO)_{10}$ is one of the most characterized, because of the historical importance of this molecule [1]. Spectroscopic characterizations at high pressure have also been reported, suggesting the occurrence of a phase transition above 0.8 GPa producing a conformational change (from staggered to eclipsed) [2]. This transformation would be very similar to that we observed on species like $Co_2(CO)_6L_2$ (L=AsPh₃, PPh₃) [3].

However, periodic DFT calculations and X-ray powder diffraction are not in agreement with the reported interpretation of the Raman spectra. Indeed, no conformational change is observed on increasing the pressure, but a curious rotation of the two $Mn(CO)_5$ moieties, leading to an unprecedented Mn-Mn off-axis bond (the Mn-Mn bond does not coincide with the Mn-CO apical axis, as in the gas phase molecular structure, see the Figure). Interestingly, this molecular geometry can be anticipated also from diffraction at ambient pressure, because the molecule shows an incipient distortion. Single crystal data at high pressure are not possible, because the samples easily break upon application of the pressure.

Theoretical calculations within the interacting quantum atom approach are also used to discuss the new geometry.



Figure 1. Left: the molecular conformation of $Mn_2(CO)_{10}$ at ambient condition (staggered). Centre: the molecular conformation at 3GPa as hypothesized from Raman spectroscopy (eclipsed). Right: the conformation of $Mn_2(CO)_{10}$ at 3GPa from P-DFT theoretical predictions and XRPD experiments.

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P16: On the purity of multi-layer focused X-ray radiation

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Multilayer optics are currently adopted in many laboratories for single crystal diffraction, using very low power X-ray generated by micro-sources. The high-brilliance and the micro-focusing are the main advantages of these radiation sources, that allow high performance experiments also on a laboratory scale.

However, we recently discovered [1] a fundamental defect of this technology, namely the significant contamination of the characteristic radiation by low energy photons which are reflected by the mirrors because of the small incidence angle. Simple experiments show that the contamination can significantly reduce the accuracy of measured intensities, especially when Mo Ka radiation is used.

We have therefore proposed a simple and economic solution to the problem [1]: an aluminium filter of adequate thickness efficiently removes the low energy contaminant photons. Performances of Al-filtered data collections are reported and alternative solutions are discussed.

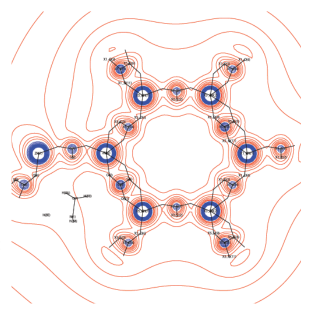
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Extensive research work has been carried out in last few decades on the synthesis and characterisation of several metal organic materials including carboxyalates, phosphates, and arsenates [1]. To a large extent, the interest is due to the interesting properties possessed by the materials and potential applications, such as electrical conductivity, magnetism, photo mechanism, host-quest chemistry etc. We focused our attention to metal oxalates, testing the possibility to model electron density of building blocks and obtain at approximate least evaluation properties. In these materials, the oxalate ion often acts as a rigid bidentate ligand



which bridge metal centres [2] therefore facilitate the formation of extended structures with dimensionalities ranging from zero to three [3].

New inorganic–organic hybrid structures based on Zinc oxalate structures, which show 1D linear, 2D honeycomb and 3D structures were studied. In order to model the building blocks of these frameworks, we used as benchmark some simple structures like $Zn(C_2O_4)$, $Zn(C_2O_4)(H_2O)$, $Zn(C_2O_4)(C_4N_2H_{10})$, $(HC_2O_4)_2(C_4N_2H_{12})$. All compounds were obtained through hydrothermal synthesis. Electron density distribution was studied through X-Ray diffraction and through density functional theory. Once the modelling was refined and tested, the electron densities of 1D-2D-3D framework were computed using multipoles restricted to optimized theoretical building blocks. This allows to reconstruct the electron density of more complex structures, often not available in the form of good quality single crystals. The results are utilised for evaluation of material properties such as electrostatic potential (a Zn oxalate honeycomb is plotted in the Figure), the interaction energies between the framework and neutral guest molecules or counter-ions, and the calculation of the active surface areas of the framework [4].

- [1] A. K. Cheetham, G.FBrey, T.Loiseau, Angew. Chem. **111**, 3466 (1999) Angew. Chem. Int. Ed. 38, 3268 (1999).
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- [3] M. Dan, C.N.R Rao, Angew. Chem. **45**, 281-285 (2006).
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P18: Pavlovskyite Ca₈(SiO₄)₂(Si₃O₁₀) - a new mineral of altered silicate-carbonate xenoliths from the two Russian type localities: Birkhin massif, Baikal Lake area and Upper Chegem caldera, Northern Caucasus

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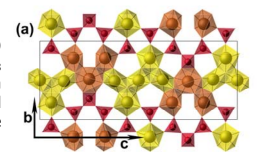
Institute of Geochemistry, Mineralogy and Petrology, Warsaw University, al. Żwirki i Wigury 93, 02-089 Warszawa, Poland

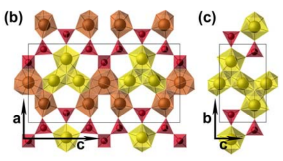
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The new mineral pavlovskyite Ca₈(SiO₄)₂(Si₃O₁₀) occurs at the rims of galuskinite Ca₇(SiO₄)₃CO₃ veins cutting calcio-olivine skarns in the Birkhin gabbro-massif (Eastern Siberia). The second type-locality are xenoliths in ignimbrites of the Upper Chegem caldera (Northern Caucasus).

The structure of pavlovskyite (a) is a 1:1 member of a polysomatic series with kilchoanite $Ca_6(SiO_4)(Si_3O_{10})$ (**b**) and calcio-olivine y-Ca₂SiO₄ (c) as end members. The structure consist of calcio-olivine slices interstratified with trisilicate modules of Ca₄(Si₃O₁₀) composition. The synthetic analog of pavlovskyite is known from cement like materials [1], but no structure refinement has been done so far.





The crystal structure of pavlovskyite, space group *Pbcn*, a = 5.0849(1), b = 11.4116(2), c = 28.6304(8) Å, V = 1661.33(7) Å³, Z = 4 has been refined from X-ray single crystal data to R1 = 3.99%. For comparison with pavlovskyite, the crystal structure of kilchoanite form the Birkhin massif, space group Ω cm, a = 11.4525(2), b = 5.0867(1), c = 21.996(3) Å, V = 1281.40(4) Å³, Z = 4 has been refined from single-crystal X-ray [1] Bennet, J.M., Gard, J.A., Speakman, K. and Taylor, H.F.W. (1966) Ca₈Si₅O₁₈ and

the nature of " γ -dicalcium silicate hydrate". *Nature*, **12**, 1127. data to R1 = 2.00%.

P19: Lithium self-diffusion in LiAlSi₂O₆ glass and single crystals

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Understanding the mechanisms of lithium diffusion is of great interest for geo- and material sciences. Optimizing the performance of Li-bearing solid media has a significant impact in developing new technologies. Knowledge of kinetic Li-isotopic fractionation leads to better understanding of geological processes in which lithium geochemistry plays a major role.

Our ongoing research is aimed to investigate Li diffusion in aluminosilicate media. In the scope of this study, spodumene (LiAlSi $_2$ O $_6$) -like materials were selected as representative model system since lithium, as the only mobile species, migrates through a static aluminosilicate network. Crystalline and glassy materials are compared in order to determine the effect of structural order on Li-diffusion. Glasses were produced by melting of oxide and carbonate mixtures as well as by melting natural spodumene. Natural crystals are from different pegmatites worldwide. Synthetic single crystals were obtained in a slow crystallization process using a flux method. The samples were tested by impedance spectroscopy for ionic conductivity in the range between 1 Hz to 10 MHz at temperatures up to 940 K. Additionally, lithium self-diffusion coefficients were determined by diffusion couple experiments using two halves with same base composition but different Li isotopic abundancies. Li isotope profiles were measured using UV fs laser ablation coupled with ICP-MS. Raman spectroscopy aided in better understanding the local structural features which coordinate lithium migration.

Ionic conductivity was found to be 6 - 7 orders of magnitude slower in natural spodumene crystals than in the glasses while the activation energy for Li conduction is about the same for both materials (0.66 kJ/mol for the glass, 0.76. kJ/mol for the crystal). This implies that the barrier for Li-migration is not sensitive to structural order in aluminosilicate materials. Comparison of Li isotope diffusion data and dc ionic conductivity yields a correlation factor of 0.5 for Li-diffusion in LiAlSi $_2$ O $_6$ -glasses.

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Pigment Red 170 is an economically important automotive coating in the paint industry, but subject to fading. Structurally it has four polymorphic forms namely the α , β , γ and δ -polymorph. Among them the γ -polymorph shows superior pigment properties [1]. It has been found that modifications of some of the molecule's substituents can affect the durability, but a rational approach to such modifications requires an understanding of the crystal structure of the pigment. Despite the importance of Pigment Red 170 in many industrial applications only a few structural studies have been carried out [1,2]. Only recently, Schmidt *et at.*, have determined the structures of the α - and γ -phases from a combination of X-ray powder diffraction and crystal structure prediction algorithms [1-2]. According to the description given, both phases have layer-like structures.

Single crystal data of the β -phase have been collected by Schmidt on a sealed tube diffractometer. They show diffuse scattering. We have collected Synchrotron data from the same crystal and found that the observed rods of diffuse scattering are typical of faulted layer stacking. There is currently some uncertainty concerning the correct unit cell that should be used to describe the average structure. The Synchrotron data suggest a larger unit cell than that derived by Schmidt. We will describe our results from the attempts to derive the correct unit cell and solve and refine the average structure.

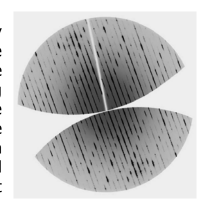


Fig. 1. The h2l reciprocal lattice plane from the β -phase of Pigment Red 170 collected on the BM01A beamline at SNBL (ESRF).

- [1] M. U. Schmidt, D. W. M. Hofmann, C. Buchsbaum, H. J. Metz, Angew. Chem. Int. Ed. **45**, 1313 (2006)
- [2] W. D. S. Motherwell, H. L. Ammom, J. D. Dunitz, A. Dzyabchenko, P. Erk, A. Gavezzotti, D. W. M. Hofmann, F. J. J. Leusen, J. P. M. Lommerse, W. T. M. Mooij, S. L. Price, H. Scheraga, B. Schweizer, M. U. Schmidt, B. P. van Eijck, P. Verwer, D. E. Williams, Acta Cryst. **B58**, 647 (2002)

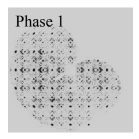
P. P. Das¹, K. W. Krämer², L. Palatinus³, H.-B. Bürgi^{1,2} and A. Linden¹

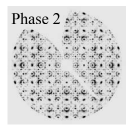
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Hexagonal β -NaLnF₄ (Ln=Y,La-Lu) compounds are a family of up-conversion materials which emit visible light upon IR excitation. Understanding of the properties of these technologically important materials requires a knowledge of their structures at an atomic level [1]. We are presently studying two phases of so-called "cubic α -NaLuF₄" obtained from the melt with a likely composition of Na₅Lu₉F₃₂.

Phase 1 show strong Bragg scattering, commensurate satellite reflections, significant diffuse scattering, while phase 2 shows just strong Bragg reflections with diffuse scattering. The strong, apparently cubic, reflections in reciprocal lattice rows are not collinear and they are split at high angles. This suggests that the crystal is a multiple twin of a structure of lower symmetry with near overlap of reflections. If the satellites are treated as





Bragg peaks, an orthorhombic supercell and six fold twinning follow with likely a space group of *Cmmm*. Considering only the positions of heavy atoms in the asymmetric unit of the small cell (based on the apparently cubic reflections), the average structure may be described equally well in two different ways for both the phases. The presence of residual electron density in the difference Fourier map of both descriptions was interpreted in terms of disordered fluorine atoms. Their positions are chemically more meaningful for one of the two heavy-atom models.

For the phase 1 structure, the phases of the superstructure reflections were determined by band flipping [2] implemented in the program Superflip [3]. The reconstructed difference electron density map shows two distinct commensurately modulated parallel columns of cations: one with varying Na⁺/Lu³⁺/occupancy and one with positional displacements of the ions from the average structure positions. Interestingly, two different solutions result from the band flipping with equal probability. These two solutions differ only by the details of positional and occupational modulation. The two solutions are distinct and the correct one can be identified by subsequent structure refinements.

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- [2] G. Oszlányi, A. Sütő, Acta Cryst. A 63, 156-163, (2007)
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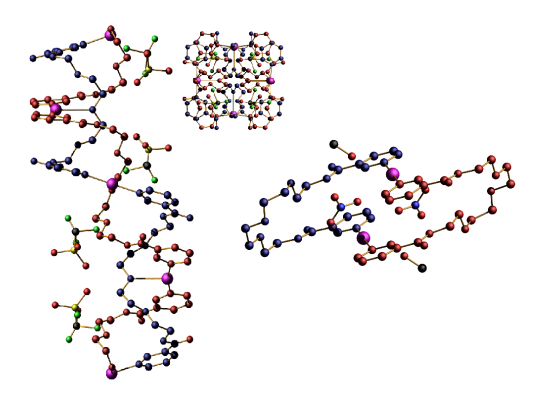
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Complexes with silver ions have a great potential for applications in medical uses. Those complexes with ligands derived from nicotinic acid and with polyethylene oxide groups as spacer are synthesized.

Upon coordination to silver ions, polymorphism can be observed: symmetric or asymmetric metallacycles, single or double helix and, under special conditions, linear chain structures are obtained.



[1] Adeline Y. Robin, Jorge L. Sagué and Katharina M. Fromm, CrytEngComm, **8**, 403-416 (2006)

Agenda of the SGK/SSCr General Assembly 2011

September 16, 2011 13.15-13.45h

Freiestrasse 3, Bern

The minutes of our last General Assembly (2010) are published on page 12 and 13 of the SGK/SSCr newsletter No. 81, Dec. 2010, available at http://www.sgk-sscr.ch/Newsletters/SGK-News-81.pdf.

- a) Jahresbericht le rapport annuel
- b) Jahresrechnung les comptes annuels
- c) Aufstellung des Budgets für das kommende Jahr/ le budget proposé pour l'année suivante
- d) Festsetzung des jährlichen Mitgliederbeitrages/ le montant de la cotisation annuelle
- e) Wahlen/Elections:
 - 1) keine
- Anträge von Mitgliedern 1) ECM Conference in Switzerland The board is proposing a Swiss bidding for the European Conference on Crystalography for 2015 and - in case of rejection - for a re-bidding for 2017 2) other Motions of members

Quorum for final decisions: \geq 10%, out of 166 members = 17 members (148 regular, 18 students, status August 9, 2011)

Patenschaften für Matura-Arbeiten



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The SCNat, which is our mother society, seeks experts for "Maturity works" (travaux de maturité / Maturaarbeiten) for ca. 4 half days per year during which pupils can come to your groups. All areas of natural sciences are welcome. More information is available here: www.maturitywork.scnat.ch. You can also fill in the attached documents

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Ohne Ihren Gegenbericht erlauben wir uns, diese Angaben in einer Broschüre und auf unseren Webseiten zu veröffentlichen

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Calls for proposals

Beside normal proposals, most facilities allow urgent beam time requests. Please check directly with the facility.

Facility	Deadline(s)	Link
SLS: Swiss Light Source		www.psi.ch /useroffice
All except PX lines	March 15, Sept. 15	/ doctoffice
Protein beam lines (PX)	Feb. 15, June 15, Oct. 15	
	,	
SINQ: Swiss Spallation Neutron	Source	
All instruments (regular calls)	May 15, Nov. 15	www.psi.ch
		/useroffice
		www.psi.ch
SµS: Swiss Muon Source		/useroffice
All instruments	Dec. 5	
ESRF: European Synchrotron		
All instruments,	lan 15	
long term proposals All instruments,	Jan. 15	www.esrf.eu
All instruments, short term proposals	March 1, Sept. 1	www.esrf.eu
Short term proposals	Маген 1, Зерс. 1	www.csii.eu
SNBL: Swiss Norwegian Beam Line	March 1, Sept. 1	www.esrf.eu/ UsersAndScience/ Experiments/ CRG/BM01/
ILL: Institut Laue Langevin		
All instruments	Mid February/September	www.ill.eu
FDM TT. Haire Maine Lail 11		
FRM II: Heinz Maier-Leibnitz	lan 2012, postpoped	ucar frm? tum da
All instruments	Jan. 2012: postponed	user.frm2.tum.de

Calendar of forthcoming meetings (Please mail missing information on meetings of interest to Jurg.Schefer@psi.ch)

Aug. 22-29	Madrid Spain	IUCr-2011, 22 nd General Assembly and Congress of IUCr http://www.iucr.org/iucr/cong/iucr-xxii	April 15, 2011
Aug. 30 – Sept. 4	Jaca Spain	2011 International School on Charge Density http://www.iucr2011madrid.es/index.php/program/satellite-meetings	May 15, 2011
Sept. 12-15	Montpellier France	Euromat 2011 – Advanced Materials and Processes http://euromatt2011.fems.eu	closed
Sept. 15-16	Villigen CH	JUM@P 11: Second Joint Users Meeting at PSI http://indico.psi.ch/conferenceDisplay.py?confId=42	to be announced
Sept. 16	Bern CH	SGK/SSCr Annual Meeting http://www.sgk2011.unibe.ch	To be announced
Sept. 20-24	Salzburg Austria	Joint Meeting of the German Crystallographic Society (DGK), German Mineralogical Society (DMG) and Austrian Mineralogical Society (ÖMG) http://www.salzburg2011.org	May 30, 2011
Nov. 10-11	Pfinztal Germany	Fraunhofer ICT: Zeit- und temperaturaufgelöste Röntgen- Pulverdiffraktometrie (X), <i>In-situ</i> -Untersuchungen und moderne Auswerteverfahren, http://www.ict.fraunhofer.de	July 31, 2011

2012

Gif-sur-Yvette France	Jan 18-19	7th SOLEIL Users' Meeting http://www.synchrotron-soleil.fr/Soleil	Dec. 1, 2011
Stoos CH	Jan 22-27	6th International Symposium Hydrogen and http://www.empa.ch	Oct. 14, 2011
April 29 – May 4	Vancouver Canada	ARRS 2012: Meeting of the American Röntgen Ray Societyhttp://www.acr.org	to be announced
cancelled	Villigen CH	PSI Powder Diffraction Summer School	Next school: 2014
July 15-18	Hamburg Germany	Science at FEL's: SRI 2012 Satellite Meeting http://science-at-fels-2012.desy.de/	Jan. 30, 2012
Aug. 7-11	Bergen Norway	ECM-27 http://ecm27.ecanews.org	to be announced
Nov. 18-23	Sidney Australia	SAS2012: International Small Angle Scattering Conference	May 2012

July 7-12	Moscow Russia	17 th International Zeolithe Conference http://www.izc17.com	to be announced
April 14-19	Washington DC, USA	ARRS 2013 – Meeting of the American Roentgen Ray Society	to be announced
Aug. 25-29	Warwick UK	European Crystallographic Association, ECM-28 http://ecm28.org	to be announced

August	Montreal Canada	IUCr-2011, 23 ^{rrd} General Assembly and Congress of IUCr http://www.iucr.org/iucr/cong/iucr-xxiii	to be announced
to be decided	Villigen CH	PSI Powder Diffraction Summer School	to be announced

to be decided	To be	European Crystallographic Association, ECM-29	to be announced
	decided	http://ecm29.org	

to be decided	To be decided	European Crystallographic Association, ECM-30 http://ecm30.org	to be announced
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Become a member of SGK/SSCr

If you are working in the field of crystallography, you will be interested to become a member of our society. For more information as well as online registration, please have a look on our website (http://www.sgk-sscr.ch). Presently, the yearly membership fee is sfr. 30 (sfr. 10 for students). For new members, the membership is free until the end of 2007. Please note: SGK/SSCr members can also apply to be a member of the subsection crystal growth (no additional charge) or for individual membership of the European Crystallographic Association, ECA (additional charge: 10 Euro).

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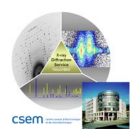






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