## 25<sup>th</sup> RAU ANNAL USERS MEETING LNLS/CNPEM September 16<sup>th</sup> and 17<sup>th</sup>, 2015

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Ministério da Ciência, Tecnologia e Inovação



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### PRESENTATION

Dear participant,

The Users Committee welcomes all the participants of the 25<sup>th</sup> Annual Users Meeting of the Brazilian Synchrotron Light Laboratory (RAU/LNLS).

This year's meeting highlights two important dates: the celebration of both the **25th Annual Users Meeting** as well as **the International Year of Light and Light-based Technologies** (IYL2015). The history of RAU started even before the LNLS has become an important international facility open to users since 1996. The laboratory's scientific community has grown fast since then, consisting of more than 1600 researchers from all over the world nowadays.

The importance of LNLS to the global scientific scenario has led to the conception of a new 4th generation machine – Sirius – which will be available to users in the year 2019. This breakthrough situates Brazil as one of the leading countries in synchrotron-based technologies. As preconized by Richard Feynman in 1959, "there is plenty of room at the bottom" and the opportunities provided by the new source will, literally, shed some light on the most challenging today's problems.

The chairperson would like to thank all the scientific committee for its efforts in the indication of the invited speakers as well as on the evaluation of the oral and poster presentations. We also thank the LNLS directors and all the local committee for the valuable support on the meeting organization.

#### Fabio Furlan Ferreira

On behalf of the LNLS Users Committee

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- Prof. Dr. Fabio Furlan Ferreira (Chair) President (UFABC)
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  - Prof. Dr. Manoel Gustavo Petrucelli Homem (UFSCar)
     6. Prof. Dr. Marcelo Alves (USP)
    - 7. Prof. Dr. Mario de Oliveira Neto (UNESP)
    - 8. Prof. Dr. Valmor Roberto Mastelaro (USP)

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    - 12. Tatiane Cortes (Communication CNPEM)
      - 13. Renan Picoreti (LNLS CNPEM)
      - 14. Vilmara Congílio (LNLS CNPEM)

### PROGRAM

	16th September 2015
08:00 - 08:30	Reception/Registrations
08:30 - 08:50	<b>Opening</b> Carlos Américo Pacheco (Director-General of CNPEM) and Antonio José Roque da Silva (Director of the Brazilian Synchrotron Light Laboratory, LNLS)
08:50 - 09:30	<b>RAU 25 years</b> Retrospective – José Antônio Brum
09:30 - 10:30	Plenary 1 Pushing the XAS frontiers: XANES approaches for chemical speciation and structure characterization and cutting-edge features of Balder beam-line at the MAX IV synchrotron radiation facility Ingmar Persson – Swedish University of Agricultural Sciences
10:30 - 11:00	Coffee Break
11:00 - 12:40	Thematic Session /
12:40 - 14:00	Lunch
14:00 - 15:00	<b>Plenary 2</b> Relaxation dynamics of isolated atoms and molecules in the tender x-ray domain (1-12 keV) <b>Marc Simon – CNRS</b>
15:00 - 16:00	Plenary 3 Application of Synchrotron Radiation for Chemical Industry Wolfgang Hoeffken – BASF
16:00 - 16:30	Coffee Break
16:30 - 17:30	Panel discussion: Committee & LNLS with the users
17:30 - 19:30	Posters Session

	17th September 2015
08:00 - 08:30	Reception/Registrations
08:30 - 09:30	Plenary 4 Looking inside materials using synchrotron X-ray diffraction Jon Wright – ESFR
09:30 - 10:30	Oral Communication 1
10:30 - 11:00	Coffee Break
11:00 - 12:40	Oral Communication 2/
12:40 - 14:00	Lunch
14:00 - 15:00	Sirius: Status and Perspectives Antonio José Roque da Silva
15:00 - 15:10	Closing

# 25<sup>th</sup> RAU ANNAL USERS MEETING LNLS/CNPEM September 16<sup>th</sup> and 17<sup>th</sup>, 2015

## ABSTRACTS







Ministério da Ciência, Tecnologia e Inovação



### The use of synchrotron radiation in Astrobiology: Lithopanspermia studies and the Biosun project

Presenter: Abrevaya X. C.

Abrevaya, X. C.<sup>1,2</sup>; Galante, D.<sup>3</sup>; Nóbrega, F.<sup>4</sup>; Tribelli, P.<sup>5</sup>; Rodrigues, F.<sup>6</sup>; Araujo, G.<sup>7</sup>; Gallo, T.<sup>3</sup>; Ribas I.<sup>8</sup>; Sanz-Forcada, J.<sup>9</sup>; Rodler, F.<sup>10,11</sup>; Porto de Mello G.F.<sup>12</sup>; Leitzinger, M.<sup>13</sup>; Odert, P.<sup>13</sup>; Hanslmeier, A.<sup>13</sup>; Horvath, J.<sup>2</sup>

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Panspermia theory assumes that microscopic life forms (e.g.: microorganisms) could survive long interplanetary travels (Arrhenius, 1903; Hoyle and Wickramasinghe, 1979). This has been matter of debate since a long time ago, opening the possibility of an extraterrestrial origin of life on the Earth. One of the open issues is that it is still not known if these life forms could survive these travels because they would be exposed to multiple extreme conditions (e.g.: radiation, vacuum). Abrevaya et al (2011) showed the capacity of some species of microorganisms to survive several doses of VUV radiation and vacuum as those related to the conditions of low Earth orbit (L.E.O.). In a new round of experiments performed at the TGM beamline (LNLS, Campinas, Brazil) we tested the survival of the microorganisms considering Lithopanspermia (interplanetary transfer of life through meteorites) in the context of the BioSun project (Abrevaya et al., 2013). The Martian meteorite "Nakhla" was chosen as model for these studies because it contains halite inclusions (NaCl evaporitic minerals). This is connected to the fact that microorganisms known as haloarchaea were found entrapped inside ancient halites (250 Mya) on Earth (e.g.:McGenity et al., 2000); therefore these organisms were proposed as possible inhabitants of Mars and possible candidates for the interplanetary transfer of life (Stan-Lotter et al., 2004). As the project is focused in the radiation environment of the young Sun, for the simulation experiments we selected as main parameters low pressure (vacuum) and VUV radiation as those we could found in L.E.O. around 3.8 Gyr ago. Two species of haloarchaea and the radioresistant bacteria D. radiodurans were entrapped inside halites and irradiated with VUV (57.5 -124 nm) with different doses up to 40000 J/m2 (eq. to 10 days in L.E.O.). We showed that the survival of of the microorganisms is strongly dependent on the specie and that halites could not offer enough protection.

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#### Application of X-Ray Phase Contrast Microtomography Using Brazilian Synchrotron Light Laboratory to Improve the Visualization of External and Internal Structures of Rhodnius prolixus head

Presenter: De Almeida A. P.

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Phase Contrast Microtomography (PhC-µCT) using synchroton radiation is a non-destructive technique that allows the microanatomical investigations of Rhodnius prolixus head, one of the most important insect vectors of Trypanosoma cruzi the etiologic agent of Chagas' disease, which accounts for about 12,000 deaths per year. The control of insect vector is the most efficient method to prevent this disease and this work is part of a series of articles [1-3] that uses PhC-µCT for the study of R. prolixus morphology. This technique provides anatomical details that could not be seen with others techniques. In this work, nymphs of Rhodnius prolixus were taken from the Laboratory of Biochemistry and Physiology of Insects, Oswaldo Cruz Foundation (FIOCRUZ), Brazil. The micro tomographic images were obtained using the new experimental setup which was recently made available at the Brazilian Synchrotron Light Laboratory (LNLS) with a 2 µm resolution, and the results showed internal and external structures of Rhodnius prolixus head. Understanding the behavior of internal and external structures of Rhodnius prolixus head can help to understand the mechanism of blood digestion by Rhodnius prolixus and its interaction with the agent of Chagas' disease, Trypanosoma cruzi, the parasite that grows within the insect's digestive system. PhC-µCT is clearly one of the best imaging techniques for insect research and has allowed a better documentation of the detailed external and internal morphology of Rhodnius prolixus without dissecting.

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### VUV excitation and SEM analysis in nanofluoride produced by microwave hydrothermal synthesis

Presenter: Andrade A. B.

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Different nanoparticles (NP's) size of barium fluoride (BaF2) has been studied by x-ray powder diffraction (XRPD) and Rietveld refinement method analysis, scanning electron microscopy (SEM), X-ray photoelectron spectroscopy (XPS) and vacuum ultraviolet excitation measures. We show in this study that BaF2 nanoparticles produced by hydrothermal microwave method (HTMW), displayed a structural lattice strain which depend of the NP's size. From Rietveld refinement method, were observed that lattice parameters to nanoparticles are smaller than crystal bulk. These refined lattice parameters increase as function of the median size distribution of the BaF2 NP's. Vacuum ultraviolet excitation spectra were performed at Brazilian Synchrotron Light laboratory (LNLS) in the Toroidal grating monochromator (TGM) beamline for all samples, and was possible observe a distinct initial and final region energy to self-trapped exciton (STE) formation energy around 9.4-9.6 eV. The width band gap was observed in 10.5 eV to all samples, without shift. These results suggest that the binding energy of the self-trapped exciton depends on particle size and that the lattice strain can be induced by lattice compression as function of the nanoparticle size effect.

### Bentonite: Al – Si k xanes characterization for this clay treated at two temperature

Presenter: Andrini L.

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Bentonite is an absorbent aluminium phyllosilicate (Smatitec 2:1 group) and Montmorillonite is the best known member of the this group [1]. Bentonites are highly valued for their sorptive properties, which stem from their high surface area and their tendency to absorb water in the interlayer sites [2] which are most extensively used are Na-Bentonite and Ca-Bentonite [3] as it are called in the industry. We studied by Al - Si K XANES a predominantly Na-Bentonite before and after thermal treatment at 800°C and 1050°C to correlate the information obtained with this technique with that obtained by conventional Xray Diffraction, and clarify and assign the structural changes. The results from the Si K XANES spectra show no major structural changes attributed to the Si sites, but there are significant changes in the Si 3p-holes density which increases due to thermal treatment. The results from the Al K XANES spectra show significant structural changes. The shoulder associated at AlO4 disappear with the thermal treatment and the first main peak has a broadening for higher temperature. The same goes for the second main peak but more intensely. This is assignable to splits Al 3p-levels due to the effects of crystal ligand field in Al morphological reconfiguration.

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### Small-angle x-ray scattering investigation of asphaltene structures in crude oils

Presenter: Balestrin L. B. da S.

Balestrin L. B. da S., Padula L., Sabadini E., Cardoso M. B. and Loh W.

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Crude oil asphaltenes represent a solubility class, not specific molecules. Crude oil asphaltenes consists of a complex mixture of several distinct molecules whose properties and structures are strongly related to the fractionation method for their isolation and to the geologic source from which they are extracted. [1] A representative asphaltene molecule is described as possessing an aromatic core of several rings, surrounded by alkyl chains. [2] Several problems related with petroleum are associated with asphaltenes stability during many operations. [3] The deposition of solids is reported at various places in the crude oil systems, thereby limiting oil production rates. [1] Crude oil asphaltenes behave, in some cases, like classical colloids. [1-3] They have been studied with a colloidal approach by numerous techniques, at various length scales. [2] They display surface-active properties and a self-assembling tendency. Overall, most of the results seem to agree that there are different levels of aggregation producing structures with varied sizes. However, there are few evidences supporting that this pattern is also present in crude oil samples (as opposed to simpler model systems). This work reports studies employing SAXS measurements on Brazilian crude oil samples with different asphaltene contents. These samples were analyzed by SAXS using synchrotron source (at the Brazilian Synchrotron Laboratory). In all cases, the SAXS curves confirmed that different levels of aggregation are presented: the first one at the nanometer scale, followed by at least two other higher levels of organization. Other experiments were performed with crude oils in the presence of diluent (toluene) or flocculant (heptane). The preliminary results suggest that the second level is affected by heptane even well below the precipitation onset, but less affected by toluene.

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#### Ultrafast charge transfer dynamics and morphological investigation in thermal annealed donor-acceptor copolymer and fullerene: F8T2 and F8T2: PCBM films

Presenter: Basabe Y. G.

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Synopsis Ultrafast charge delocalization dynamics on internal donor-acceptor polymer poly(9,9-dioctylfluorenyl-co-bithiophene) (F8T2) and its blend with the fullerene derivate [6,6]-Phenyl  $C_{61}$  butyric acid methyl ester (PCBM) was studied by the resonant Auger spectroscopy (RAS) measured around sulfur K-edge using the synchrotron-based corehole clock approach. The RAS spectra were measured at the SXS beamline at the Brazilian Synchrotron Light Source (LNLS) using an ultrahigh vacuum chamber with a base pressure of 10<sup>-8</sup> mbar and hemispherical electron energy analyzer. The effect of thermal annealing on the charge transfer delocalization times ( $\tau_{CT}$ ) was also investigated recording RAS spectra while detuning the photon energy around S-1s absorption edge. Poor charge delocalization was observed for as-cast polymeric films at photon energies corresponding to the S 1s  $\rightarrow \pi^*$  transition, which may suggest a weak  $\pi$ -electronic coupling due to weak polymer crystallinity and chain stacking. Atomic Force Microscopy (AFM) topography for as cast F8T2: PCBM shows a top position of PCBM units relative to the polymer, homogeneously distributed in film surface. This configuration improves the charge delocalization through S 1s  $\rightarrow \pi^*$  molecular orbital for as cast blended film, suggesting a strong  $\pi$ -electronic coupling. A new rearrangement of F8T2:PCBM film was found after thermal annealing, leading to a more efficient electron transfer channel through  $\sigma^*$  molecular orbitals.

#### Hybrid nanostructured sulfonated poly (ether ether ketone) / zirconium oxide based membranes comprising protic ionic liquid for PEMFC application

Presenter: Batalha J. A. F. L.

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In view of the lack of fossil fuels in a near future, and aiming, not only to diminish the environmental impacts caused by its gas emissions from combustion, but also better quality of life for the population, alternative energy sources are being each time more studied, which is the case of fuel cells, normally fed by hydrogen. Among the many types of fuel cells, the solid, non-corrosive, non-toxic polymer electrolyte one stands out. The great challenge of this technology nowadays is that its operation occurs in temperatures higher than 100 °C, the limit for Nafion®, reference material for electrolytes. Many efforts are being effected towards this possibility, such as synthesis of composites with inorganic fillers. However, the most promising studies face the employment of ionic liquids, especially the protic ones, as integral replacement to water in the operation of this kind of fuel cell. With this being said, the greatest challenge to the development of this technology is to retain the ionic liquid within membranes, without great decrease in conductivity properties. In this work, hybrid membranes based on sulfonated poly(ether ether ketone) (sPEEK) and zirconium oxide, incorporated via sol-gel, comprising diethylmethylamine triflate ([dema][TfOH]) were explored. The membranes presented good mechanical and chemical stabilities, as well as thermal stability over 300 °C. SAXS curves indicated the possible formation of ionic domains. Moreover, other analysis such as ionic conductivity, DMTA and XRD were also executed.

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#### Structural and Optical Studies of Nanoparticles of CaF\$\_2\$

Presenter: Bezerra C. dos S.

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The development of scintillators materials is an experimental field that has attracted attention from researchers in last years and one interesting class of materials are the fluoride matrices doped with rare earth ions that have attracted interest also due to applications in lasers. Compared with the oxide matrices fluorides have advantages such as high transparency in a wide wavelength region from the VUV to IR and low energy phonons. In this work, the CaF\$ 2\$ pure and doped with erbium were produced by microwave-assisted hydrothermal synthesis, which is a promising method to produce nanostructured powders in a simple, low cost and fast way. The samples were produced using chelating agent (EDA-Ethylenediamine) and synthesized at low temperature and a short time period. The crystalline phases of the samples were identified by X-ray diffraction (XRD) and the morphology of the particles was determined by scanning electron microscopy (FEG-SEM). The results shown that the microwave-assisted hydrothermal synthesis is efficient for production of the CaF\$ 2\$ nanoparticles and the use of the chelating agent in the production of the samples influences the average particles size. The study of the chemical composition in the surface was performed by X-ray Photoelectron Spectroscopy (XPS) and indicated the presence of greater amounts of hydroxyl groups and oxygen ions in the samples produced with EDA. Radioluminescence (RL) measurements showed that the CaF\$ 2\$ presents an intrinsic luminescence with an emission band at approximately 300 nm. Optical study was conducted to understand the formation of band structure and to determine the band gap of the materials. The excitation spectra in the VUV region were measurements in Brazilian Synchrotron Light Laboratory (LNLS) and the results shown evidences that the formation energy of the exciton and band gap energy are smaller than the ones for the single crystal.

### Arsenic speciation in geological samples using X-ray absorption spectroscopy

Presenter: Bia G. L.

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The Chaco Pampean plain is one of the largest regions in the world (1,000,000 Km2) affected by the presence of high concentrations of As in groundwater (Nicolli et al. 2012). The primary source of As is associated with the presence of volcanic ash spread in the loessic sediments that blanket the entire region. Bia et al. (2015) identified by XPS the presence of As (III)-S and As (V)-O nanocoatings, deposited on the surface of volcanic glass particles. The As contained in these coatings is easily released in contact with water, but a higher proportion of As remain within the aluminosilicate glass structure as an impurity. The aim of this work is to determine the oxidation state of arsenic, its local chemical coordination (to a radius of ~4 Å around As) and the relative proportion of the As species in volcanic Andean ashes and loess sediments. To carry out this study, volcanic ash samples of recent eruptions (Hudson, 1991 Chaitén, 2008 and Puyehue, 2011) and samples of loess were analyzed by Xray absorption fine structure spectroscopy (XAFS), using the facilities of the Brazilian Synchrotron Light Laboratory (LNLS, Brazil). XANES analysis allowed to discriminate three oxidation states of arsenic in the studied samples: As5+ is the dominant oxidation state in loess sediments while in volcanic ash samples just the oxidation states As-1 and As3+ were identified. The proposed EXAFS models fit well with the experimental data, suggesting that in loess sediments, As5+ could be in the form of arsenate ions adsorbed onto ferric oxyhydroxides, or precipitated as scorodite (FeAsO4•2H2O). In volcanic ashes, the species As-1 identified is likely associated with arsenopyrite (FeAsS) or arsenical pyrite (FeS2-2xAsx), while As3+ is likely related to As atoms present as impurities within the glass structure.

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#### Nanostructured Ga doped ZnO thin films prepared by sol-gel spin-coating

Presenter: Bojorge C. D.

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ZnO based films have been actively studied because of their applications as solar cells, gas sensors, piezoelectric transducers, ultrasonic oscillators and for different optoelectronic applications. Besides their interesting optical, electrical and piezoelectrical properties, this material exhibits a high chemical and mechanical stability. ZnO presents novel properties and potential applications in optoelectronic fields because its nonlinear optical properties, excitonic emission at room temperature and quantum size effect. In the present work we studied pure and Ga-doped ZnO thin films prepared by the sol-gel spin coating technique by grazing incidence small-angle X-ray scattering (GISAXS) and X-ray reflectivity (XR) methods. We also compare the present spin coating pore distribution results with those obtained in previous works.

#### **Photoionization studies on Thionitrites**

Presenter: Canneva A.

Nitric oxide (NO) has been recognized as a fundamental molecule in biology and medicine. Among them, S-nitrosothiols, RSNO, have been shown to store, transport, and release NO within the mammalian body(1). It has been suggested that the formation and decay of low molecular weight RSNOs, such as S-nitrosoglutathione and S-nitrosocysteine, also represent a mechanism for the storage or transport of NO.(2) The determination of the electronic properties of simple thionitrites is an important task, because their serve as a model compounds for more complexes species, especially in connection with the previously The main objective was the determination of their described biological importance. molecular and electronic properties. Thus, depending on their chemical stability, different approaches have been applied, including Gas Electron Diffraction and vibrational spectroscopy techniques. Moreover, the gas phase He(I) photoelectron spectrum (PES) of t-BuSNO has been measured. Here, we present the results of studies on the shallow-core electronic levels of thionitrites species (CH3)2CHSNO and (CH3)3CSNO. The dissociation process that dominates the RS-NO photolysis is mostly influenced by the electronic distribution around the sulfur atom as we conclude of the analysis to the sulfur 2p level, which could be studied by using the TGM beamline.

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#### X-ray absorption spectroscopy characterization of Zn1-xCoxO thin films applied as ozone gas sensors

Presenter: Catto A. C.

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Zinc oxide (ZnO) pure or doped are one of the most promising metal oxide semiconductors for gas sensing applications due the well-known high surface-to-volume area and surface conductivity. It was show that ZnO is an excellent gas-sensing material for different gases such as CO, O2, NO2 and ethanol. In this context, pure and doped ZnO exhibiting different morphologies and a high surface/volume ratio can be a good option regarding the limitations of the current commercial sensors. Different studies showed that the sensitivity of metaldoped ZnO (e.g. Co, Fe, Mn,) enhanced its gas sensing properties. Motivated by these considerations, the aim of this study consisted on the investigation of the role of Co ions on structural, morphological and the gas sensing properties of nanostructured ZnO samples. In this work, we will present the results concerning the short-range order study of the Zn1xCoxO (0 < x < 5 wt%) thin films deposited by means RF-magnetron sputtering. Their gas sensing properties were investigated by exposing the samples to different ozone levels (0.06-2.5 ppm). Electrical measurements indicated that the Co addition into the nanocrystalline ZnO samples favors the ozone gas sensor performance of the Zn1-xCoxO thin films. The Zn and Co K-edge X-ray absorption spectra of Zn1-xCoxO thin films were collected at the XAFS2 LNLS beam line in order to investigate the electronic structure around Zn and Co atoms. In the as-prepared samples, Zn and Co K-edge XANES measurements pointed out the presence of metallic Zn and Co state. The annealing treatment in oxygen-rich atmosphere improved the thin film oxidation with XANES reveling the presence of Zn(II) and Co(III) Acknowledgements: This work was supported by Brazilian financing agencies ions. FAPESP and CNPq.

- T. Jiang; D. E. Ellis. X-ray absorption near edge structures in cobalt oxides. J. Mater. Res., Vol. 11, n. 9, p. 2242-2256, 1996. - H. S. Al-Salman, et al. Structural, optical, and electrical properties of Schottky diodes based on undoped and cobalt-do

### Iron altered oxidation energy due to Yttrium doping in Fe3O4 spinel ferrites

Presenter: Coelho L. N.

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Doping of spinel ferrite nanoparticles (NPs) alters significantly their magnetic response and other characteristics REF. Furthermore magnetic fluids made with yttrium (Y) doped NPs show high stability, remaining years without settling or flocculating. The precise character of Y doping effects on the NPs is yet to be fully understood. EXAFS measurements showed that Y occupy the tetrahedral site, entering the crystalline structure without causing much structural damage, as can be readily verified by X-Ray diffraction data. One hypothesis for enhanced colloidal stability in these ferrites revolves around the fact that these systems, in particular, are ionic colloids, without any surfactant molecule on the particle surface. The stability, in these cases, comes from electrostatic repulsion between particles, coated with charge of equal sign that repel each other efficiently. The charge is highly influenced by the system pH. In this work we will present X-Ray Photoemission Electrons spectroscopy measurements which strongly indicates that the Y-doping of the NPs favors Fe2+ oxidation into Fe3+ ions, consequently increasing the particle surface charge. This is readily perceived by Zeta-potential measurements of doped and undoped particles at the same pH. The Ydoped NPs presented a Zeta potential about 38% higher than the undoped sample. Since the zeta potential is derived from surface charge which is directly associated with colloidal stability this different oxidation energy detected by XPS may be the main cause of low flocculation and virtually no sedimentation in Y-doped ferrites.

#### XAFS characterization of Sr1-xCuxTiO3 and SrTi1-xCuxO3 perovskites applied to water-gas shift reaction

Presenter: Coletta V. C.

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Strontium titanate (SrTiO3) has a wide range of possible applications, including catalysis, due to its high stability and the possibility of incorporation of active metals into the perovskite structure by the partial substitution of strontium or titanium sites. Coppercontaining catalysts are a low-cost alternative to noble metals, such as gold and platinum for the water-gas shift (WGS) reaction. However, there are no reports of the Sr1-xCuxTiO3 and SrTi1-xCuxO3 systems as catalysts for the WGS reaction to our knowledge. Thus, our research work consists in the synthesis and characterization of nanoparticles of Sr1xCuxTiO3 and SrTi1-xCuxO3 aiming for the application of these materials as catalysts for the water-gas shift reaction. The synthesis was carried out through a modified polymeric precursors method, with calcination in an inert atmosphere followed by a treatment in oxygen to remove carbon, which resulted in particles with large area in comparison to the conventional calcination in air. We performed in situ X-ray diffraction and absorption experiments at Cu and Ti K-edge and fitted the EXAFS spectra to study the reducibility of copper. Our results showed the specific reduction of copper from 2+ to metallic state after a reducing treatment until 350°C, which was kept during the WGS reaction. The presence of metallic copper is related to the activity, which reached 74% of CO conversion for SrTi0.80Cu0.20O3.

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#### Structural characterization of kraft lignin under different pH

Presenter: Dias O. A. T.

Dias O. A. T., Negrão D. R., Gandin C. A., Leão A. L., Neto M. de O.

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Lignin is a biopolymer with high molecular weight which can be totally solubilized on alkaline pH. After that, its molecular structure became very reactive occurring rearrangement till its stabilization. Because of this, is appropriate to wait a time for its molecular stability, approximately one week (Maziero et al., 2012). In this assay, lignin was recovered by acidification of the Kraft black liquor (LNK) at two pH 3.0 and 5.0. 5 mg of each type of Kraft lignin were transferred to 1.5 mL microtubes and daily was added 1 mL of NaOH solution (0.1M) in each tube. Samples were prepared to be diluted for 10 days in duplicate. SAXS analyses provided different patterns in fractal order of lignin due to its structural reorganization mainly till 6th d. After 7th d of dilution, it was observed an increase of fractal complex these molecules for both lignins obtained at pH 3.0 and 5.0, indicating molecular stabilization. Both lignins suffered an increase in the Rg according to the dilution days, stabilizing at 8th d; same behaviour was observed with polydispersity. The size of Rg from lignin recovered at pH 3.0 was lower than lignin recovered at pH 5.0. Based on the obtained data, SAXS analysis contributed to the structural characterization of lignin obtained at two pH regarding to the size, shape and polydispersity.

MAZIERO, P., OLIVEIRA NETO, M., MACHADO, D., BATISTA, T., CAVALHEIRO, C. C. S., NEUMANN, M. G., CRAIEVICH, A. F., ROCHA, G. J. M., POLIKARPOV, I., GONÇALVES, A.R. Structural features of lignin obtained at different alkaline oxidation conditions from sugarcane bagasse. Industrial Crops and Production, v. 35, 61-69, 2012.

#### Photoionization dynamic of O-methyl dithiocarbonate (dimethyl xanthate), CH<sub>3</sub>OC(S)SCH<sub>3</sub>, on the sulfur 2p absorption edge

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Xanthates (from the Greek "golden") are the esters of xanthic acid, with general formula ROC(S)SR'. These are important organosulfur compounds used in the production of cellophane and as flotation agents for extraction of metals from many ores. As an example, sodium ethyl xanthate is added as a collector in the selective flotation of PbS (galena) to separate it from ZnS (sphalerite). A series of xanthate and xanthogen derivatives are being the subject of investigation in our group, including synthetic aspects together with the elucidation of structural, conformational and vibrational properties.<sup>1</sup> Following our general project aimed at elucidating the shallow and inner-shell core electronic properties and photoionization dynamics of sulfur-containing derivatives,<sup>2,3</sup> we became now interested in the simplest xanthate compound, with R=R'=CH<sub>3</sub>. Here we report a study of the photon impact excitation and dissociation dynamics of CH<sub>3</sub>OC(S)SCH<sub>3</sub> exited at the S 2p level by using synchrotron radiation. Interestingly, the photoelectron spectrum of CH<sub>3</sub>OC(S)SCH<sub>3</sub> in the valence region (7.5-13 eV energy range) was reported by Guimon et al. in 1974 and interpreted in terms of ionization of non-bonded orbital mostly localized on the sulfur atoms. <sup>4</sup> The S 2p core XPS and KLL Auger spectra of CH<sub>3</sub>OC(S)SCH<sub>3</sub> were early reported by Summinen et al. <sup>5</sup> showing that the ionization energies of the =S and -S- atoms within the xanthate group differ from each other by 1.5 eV, this difference assigned to the initial-state charge distribution rather than to final-state relaxation. In this presentation, the TIY spectra of CH<sub>3</sub>OC(S)SCH<sub>3</sub> following S 2p excitations is reported and compared with the previous data. The dissociative photoionizaiton is discussed in term of PEPICO and PEPIPICO spectra and possible fragmentations mechanisms are deduced from the interpretation of the PEPIPICO spectra. Thus, the advantage of energy tunability offered by synchrotron radiation allowed us for a clear understanding of the resonant electronic processes occurring after excitation of each of the sulfur atoms present in this molecule and how the ionic dissociation is affected.

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#### Core-shell aggregates formed by polyion-surfactant complex salts: study of the internal liquid crystalline structures and stability

Presenter: Ferreira G. A.

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We have studied the self-assembly of complex salts, formed by ionic surfactants and polyions, in aqueous solution, to vield different liquid crystalline phases. Our group investigated the phase behavior of hexadecyl (C16TA+) and dodecyltrimethylammonium (C12TA+) surfactants counterbalanced by poly(acrylate) homopolymer in water and in the presence of different alcohols [1,2]. It was possible to achieve different liquid crystalline phases (cubic, hexagonal and lamellar), depending on the surfactant chain length and the cosolute used. Recently, we have reported colloidal dispersions of core-shell aggregates formed by complex salts of poly(acrylate-b-acrylamide), PAA-b-Pam, polymers in which the cores displayed liquid crystalline structure with the same geometry as the ones observed for complex salts of the poly(acrylate) homopolymer [3]. We are now using the same approach of employing a complex salt of the block copolymer PAA-b-Pam and cationic C12TA+ and C16TA+ surfactants to prepare systems with adequate compositions to reproduce the abovementioned phase diagrams attempting to obtain stable core-shell dispersions with the cores reproducing the phase structure observed in the previous studies. Preliminary results showed the presence of different liquid crystalline cores by adding different amounts of n-alcohols, in agreement with our previous reports using the complex salts prepared with the homopolymer. The poly(acrylic acid-b-isopropylacrylamide) (PAA-b-PNIPAm) polymer is also being used to analyze the effect of temperature, since PNIPAm has T LCST~32°C. This system holding PNIPAm shows similar crystalline phases at 25°C and 45°C, although there is a small contraction of the core size related to the increase in the hydrophobicity of the PNIPAm chains and segregation of a concentrated phase. Attempts to stabilize the dispersion in water using a second complex salt of PAA-b-PEO are promising, once PEO has more affinity to water and no further changes were observed.

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### Analysis of the concentration of heavy metals in the polychaete using the TXRF technique on sandy beaches of the coast of São Paulo

Presenter: Freitas M. C. S.

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The sediment tends to embody and accumulate contaminants poured in the water, in which become available for associated organisms. In this regard, benthic invertebrates such as polychaetes, which often use the sediment to obtain food or to construct protective tubes, are susceptible to the contamination by elements present in this environment. Thus, the present study examined the concentration of heavy metals in the species of polychaete Dispio remanei Friedrich, 1956 found at the beaches of Itararé (São Vicente) and Boracéia (Bertioga) and Orbinia cf. johnsoni Moore, 1909 in Araçá bay and Guaecá beach, both located in São Sebastião. The Aracá and Itararé are located very close to port areas, characterized by high urbanization and industrialization, thus existing possibilities of coastal marine environment being contaminated by heavy metals, that can have negative impacts on the biological, social and economic aspects. During periods of low tide, on each site, was delimitated a quadrat area of 100 m<sup>2</sup>, in which three points were randomly selected to obtain individuals D. remanei and O. cf. johnsoni. The material collected was analyzed by using the X-Ray Fluorescence by Total Reflection (TXRF) technique; it is possible to perform the correlation between essential and toxic elements in the body, through the issue of spectrum with energies characteristics of each metal. From procurement of different metal concentrations, comparisons of the mean concentrations between the beaches were conducted. Significant differences were identified for concentrations of Zinc with a higher concentration of the beach Boraceia regarding to the Itararé beach. This result was different than expected, because the beach Boraceia lies far from the places where this metal is used. For species O. cf. johnsoni, we identified a significant difference to the Fe element in the Bay of Aracá, which had a higher concentration than Guaecá Beach. This result was expected, since the Bay of Araça is a receiving area of effluent contents, and it is also very close to the port of São Sebastião, as well as to the industries of the region. Both beaches showed significant differences for the element Potassium.
#### Present and future developments for UV-VUV science at LNLS

Presenter: Galante D.

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In this work, recent advancements and future perspectives for the science done at the UV-VUV range at LNLS will be discussed. Currently, the TGM beamline of LNLS can operate from 3 to 330 eV (ca. 400 to 4 nm), and it is coupled to LNLS D05 (4°) bending magnet, which is one of the two bending magnets providing photons with energies below 100 eV. Its design aims to provide photon fluxes of up to 1E10 photons per second per mm2 with moderate resolving power. Three toroidal gratings are used to efficiently cover the large energy range with the following specifications: A Pt coated grating with 75 grooves/mm and two 200 and 1800 grooves/mm. Their optimal energies range are from 3-13 eV, 12-100 eV, 100-330 eV respectively. TGM was the first beamline of LNLS to be built and installed, and it has been upgraded on the last few years to ensure competitive characteristics on its energy range. In fact, it is currently the sole synchrotron beamline operating at the UV-VUV on the Americas. Recent upgrades have included the change on the control system, installation of a new grating for low energies, automation of its gas filter for harmonics' suppression and improvement of monitoring and stability, as ongoing projects. Near-future developments include the installation of a photoemission microscope (PEEM), and the development of new experimental stations for circular dichroism (SRCD), measurement of optical properties of materials (luminescence, fluorescence, absorption) and gas-phase reactions. These new stations are being developed in collaboration with the users' community. With Sirius being built, we will also discuss the perspective of the installation of a UV-VUV beamline at it, in order to ensure that the current and future needs for experiments at this energy range are fulfilled on the best possible way.

### Degradation of carotenoid of poly-resistant bacterium Deinococcus radiodurans on simulated environments with applications in astrobiology

Presenter: Gallo T. M.

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From the moment that an organism dies and is buried the original organic material tends to be lost being transformed into disorganized forms of carbon or in some cases mineralized. Organic molecules derived from the buried body can also be fossilized sometimes being the only trace of their existence. These are known as chemofossils or biomolecules that after burial and changes imposed by different geological processes exhibit changes in their structure, but also bringing characteristics of the original molecule which in some cases may contribute for the identification of the organism that originated the molecular signature. Chemofossils are also of interest in astrobiology since the conditions of fossilization in extraterrestrial environments similar to what eventually happens on Earth may have not favored the preservation of the body but only of its molecules. We used different mineral substrates mixed with the poly-resistant bacterium Deinococcus radiodurans for the study of the interaction of one chosen biosignature (carotenoids, present on this bacteria) for the preservation process. We prepared pellets of substrate with bacterium and irradiated them in three different environmental conditions with different doses: with a solar simulator (UVA/UVB), in Martian simulation with a specially designed chamber (AstroCam), and with UV-VUV for space simulation at the TGM beamline of LNLS. After these different simulations we used Raman spectroscopy with 532 nm and 785 nm excitations to analyze the degradation of the carotenoid. It was observed there is a reduction of the carotenoid Raman signal as function of the dose and there are significant differences when the sample is irradiated with different wavelengths and in different substrates. These results can contribute for a better interpretation of future data from Mars explorations missions, such as ExoMars (2018, ESA) and Mars2020 (2020, NASA), which will carry Raman spectrometers for the search for signals of life.

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## Grazing-incidence X-ray scattering studies of myelin membranes at air/water interface

Presenter: Gasperini A. A. M.

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Thin and ultrathin films in liquid surfaces and interfaces have been intensively studied in different domains such as pharmaceutics, electronics, and food industry. These materials represent a model case to study the dynamical and structural properties of 2D confined soft matter. A very interesting class of ultrathin films is given by the Langmuir monolayers, which are monomolecular layers of amphiphilic molecules spread on air/water interfaces, since they can represent a model for biological systems. Langmuir monolayers formed by myelin lipids or full myelin membranes offer advantageous features to approach the exploration of the surface properties of a complex lipid-protein interface since they contain all the components of natural membranes[1]. Surface-sensitive synchrotron-based X-ray scattering techniques such as grazing-incidence diffraction (GID) and scattering (GISAXS/GIXOS) are powerful tools for investigating the molecular structure of organic films on liquid surfaces due to the penetrating properties of the X-rays and the ability to couple structural and dynamical properties of interfaces with nanoscale resolution[2]. We present in this work the first results of a surface-sensitive X-ray scattering experiment performed at LNLS/XRD-2 beamline[3]. We used this technique to study Langmuir monolayers formed by myelin lipids and full myelin membranes as a function of the monolayer surface pressure. The new experiment opens new possibilities for the soft condensed matter community in Brazil. The structural study as a function of the change of preparation conditions and thermodynamical parameters make this kind of experiment ideal for many technological and industrial applications. AAMG, RGO and LCP acknowledge grants from FAPESP. XPM thanks to CNPq.

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### Magnetic moment of Fe3O4 films with thicknesses near the unit-cell size

Presenter: Gomes G. F. de M.

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There is a large scientific and technological interest in magnetic materials in ultrathin film form due to the anisotropy created by the low dimensionality which can change physical properties. Magnetic and structural properties of ultrathin iron oxides composed of less than 10 atomic layers are poorly understood [1-4]. The magnetic spin moments for ultrathin magnetite as determined experimentally by various groups [1-3,5] present much lower values than the bulk spin moment, which is very intriguing. Another open question is what is the lowest thickness required to maintain the bulk ferrimagnetic order at room temperature. We have followed the evolution of the magnetic moment of ultrathin Fe3O4 as a function of the thickness (8–45  $A^{\circ}$ ) in the [100] and [111] crystallographic directions. The stoichiometry and surface quality of the films have been attested by X-Ray absorption spectroscopy (XAS), X-Ray magnetic circular dichroism (XMCD), and low-energy electron diffraction (LEED). The XMCD study were performed at the PGM beam line at the Laboratório Nacional de Luz Síncrotron (LNLS). The results show that at low thicknesses, FeO has probably been formed and evolved to Fe3O4 above the thickness of 1 unit cell. For both the [100] and the [111] orientations we observe a characteristic dichroic signature of magnetite around the unitcell thickness, however, with a lower spin moment that evolves to the bulk value for thicker films. A spin moment of 3.6µB/f.u. was found at 35 °A thickness for Fe3O4 [111] [6]. These results are direct evidence that both the ferromagnetic order and the bulk moment value are preserved at room temperature around the thickness of 2 unit cells. We have also shown that even 10-A° -thick magnetite already presents a significant magnetic moment. We believe that the conclusions of this work are of great importance, especially if magnetite layers down to subnanoscale thickness are employed in spin electronics devices.

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## X-ray crystallographic structure of a transcriptional activator of virulence factor in enterococcus faecalis, ELRR.

Presenter: De Groote M. C. R.

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RNPP family of transcriptional regulators is present exclusively in Gram positive bacteria and regulates factors involved in virulence, invasion, biofilm formation, etc. That family shares structure, functions and also the regulation by peptides. ElrR protein is a transcriptional regulator of Rggfamily, that positively regulates ElrA, which is a virulence factor of Enterococcus faecalis. Rgg familly was proposed to belong to RNPP family. Here we present the crystallographic structure of APO ElrR obtained at 2.15Å resolution. We solved the phase problem using diffraction data at 2.55 Å resolution of Se-Met-ElrR crystals measured on the absorption edge ( $\lambda$ =0.979096 Å) of Selenium atoms, collected at LNLS Syncrhotron, Brazil. We then improved the resolution using a dataset obtained at home  $(\lambda = 1.5418 \text{ Å})$  which diffracted at 2.15 Å resolution. This structure was refined to Rwork 0,2024 e Rfree 0,258. Crystals presented space group P1 with crystallographic cell a 70.800 b 75.330 c 85.280  $\alpha$  112.99  $\beta$  89.93  $\gamma$  103.88 which contains two homodimers. The crystallographic model has shown the same fold in ElrR as PgrX and RGG2sd structures, in spite of the low sequence identity (under 13% in all cases). The structure presents a helixturn-helix XRE-type domain on the N-terminal, a 5-helix central domain and a C-terminal domain build of tetratricopepitides (TPR)-like repetitions. The interface between the Central and the C-terminal domains contains the alosteric site, like in distant homologous proteins of the RNPP family, but the cleft on the alosteric site in ElrR is wider. Then, ElrR ligand can be bigger than those found in these homologous structures.

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### Electronic Structure of SrTi<sub>1-x</sub>Ru<sub>x</sub>O<sub>3</sub>

Presenter: Guedes E. B.

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Metal-insulator transition (MIT) in an intriguing phenomenon in transition metal oxides that can have different driving forces, such as electron correlation, disorder or percolation. The  $SrTi_{1-x}Ru_xO_3$  (STRO) series undergoes a MIT for  $x_c \sim 0.35$ . The end members of the series show completely different physical properties: SrTiO3 (STO) is a diamagnetic band insulator with cubic structure, whereas SrRuO<sub>3</sub> (SRO) is an itinerant ferromagnet with an orthorhombic structure. The electronic structure of the STRO series has been studied via electrical conductivity measurements [Phys. Rev. B 62, 10785 (2000)], X-ray photoemission and x-ray absorption spectroscopies [Eur. Phys. J. B 25, 203 (2002), Phys. Rev. B 73, 235109 (2006)], band structure calculations [Phys. Rev. B 77, 085118 (2008), Phys. Rev. B 77, 212407 (2008)] and other techniques. Despite the efforts, the mechanism leading to the transition in this system is still under debate. Previously, a study on SRO where several spectroscopic measurements were interpreted via an extended cluster model [Phys. Rev. B 86, 235127 (2012)] showed that the ground state of this material is dominated by configurations with holes in the ligand band. Now, in order to study the electronic structure of the STRO series, we developed and implemented a double cluster model, in which two octahedra are connected via an oxygen atom. The model includes correlation, covalence and multiplet effects, as well as charge fluctuations between the two octahedra. The results are compared with X-ray Photoemission Spectroscopy from the literature, and all the spectra were reproduced with a single set of parameters.

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# Structure, activity and reaction mechanism of NahK and its complex with NahL

Presenter: Guimarães S. L.

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Among all types of toxic Polycyclic Aromatic Hydrocarbons (PAHs), naphthalene is the most studied and a model compound. Its degradation by the bacteria Pseudomonas putida is the focus of numerous studies. In this organism, two consecutive pathways are able to convert naphthalene to pyruvate and acetyl-CoA. In the lower pathway, the enzymes NahK (4oxalocrotonate decarboxylase) and NahL (vinylpyruvate hydratase) catalyze an interesting sequence of reactions that raise a number of mechanistic and structural questions. In this work we aim to characterize kinetically and structurally the NahK enzyme expressed without NahL, and also both enzymes co-expressed as a complex. The nahK gene was subcloned into pET28a-TEV vector for expression, and also along with nahL gene into pETDuet-1 vector for co-expression in Escherichia coli BL21(DE3) cells. The protein samples were analyzed by Dynamic light scattering, Circular dichroism, and the steady-state kinetic values for their natural substrates were obtained. The NahK/NahL complex was also analyzed by Small-angle X-ray scattering. NahK and NahK/NahL samples in their apo or ligated forms with different ligands were crystallized and X-Ray diffraction data were collected. NahK expressed alone and co-expressed with NahL was purified to homogeneity and both enzymes show kcat/Km values up to 10<sup>7</sup> M-1 s-1. The crystalline structures of NahK in apo and ligated forms with Mg2+ and substrate analogues allowed us to propose a dynamic reaction mechanism involving a lid domain. SAXS measurements suggested a particle of 240 kDa for NahK/NahL complex, and the crystalline structure of this complex confirmed a heterodecamer assembly. NahK and NahL face their active site cavities to each other into a large, elegant and functional quaternary structure. Our work provided a deep structural and functional understanding of this interesting protein complex.

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#### Au / Ag nanowires atomic distribution revealed by XAFS

Presenter: Herrera F. C.

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Nanowires which were defined as having at least two dimensions between 1 and 100 nm, have received a great interest due to their unique optical, electrical, magnetic, and thermal properties with dimensionality and size confinement [1-4]. The intrinsic properties of nanowires are mainly determined by its size and composition. To study the size dependent properties, it is the crucial task to synthesize size-controlled nanowires. In order to investigate the local and electronic structure of Ag-Au(1-D) nanowires we perform a comprehensive study by XAFS in the Au-L3, Ag-L23 and Ag-K edges of isolated Ag-Au(1-D) nanowires with different Au:Ag molar ratio. Ag-Au(1-D) nanowires were synthesized by a method of colloid chemistry in a first stage (Ag), followed by a galvanic reaction to incorporate the second metal in the structure (Au). Au L3 EXAFS data shows that every Au atom, independently of the Au:Ag molar ratio, are incorporated in the Ag lattice surrounded by 8 and 4 Au and Ag atoms respectively. Ag K EXAFS data analysis shows that depending on the Au:Ag molar ratio the Ag atoms change their environment. This result is also supported by Ag L23 XANES data that shows different hole density in the Ag 4d level depending on the Ag-Au(1-D) nanowires stoichiometry.

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#### Python as a tool for analyze X ray small angle scattering data

Presenter: Huck-Iriart C.

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Small angle x-ray scattering (SAXS) analysis requires several numerical steps to be done, starting from data reduction to data interpretation (physical models, invariants calculations, etc). All of these steps must be done in order to squeeze the information obtained by [1]. In the last decade they began to appear few user friendly programs available which allow doing a full data analysis [2]. Nevertheless, some of those software packages are useful to analyze simple systems or to perform a set of predetermined routines for SAXS data analysis. Other programs let users to write new functions or adding in-house written plug-ins. Those types of packages allow a complete and flexible analysis of the experimental data but at the expense of learning a specific programming language. Python is a general purpose, high level programming language with a coding syntax very easy to learn and is not restricted only to scientific computation. Language features and a small set of core packages includes: NumPy (mathematical arrays), SciPy (linear algebra, differential equations, signal processing and more), SymPy (symbolic mathematics), matplotlib (graph plotting), etc; turns Python into a very popular programming choice in the scientific community [3]. The main aim of this contribution is to introduce Phyton to the general user and to show some of its capabilities applied to the particular case of SAXS data analysis including a full pixel GISAXS image analysis and a few tricks to speed up computer calculations.

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### Real Time Monitoring Nanoparticles Distance and Structure upon Different Variables.

Presenter: Ibañez F. J.

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The nanoparticles (NPs) separation, order, and structure play a crucial role not only in in fundamental science but also, in the applied field such as in chemiresistors, localized surface plasmon resonance (LSPR), and surface-enhanced Raman scattering (SERS) for just mention a few. For instance, as long as a metal nanoparticle (Ag or Au) approaches another metallic center, electromagnetic waves are enhanced leading to an improved SERS or LSPR effect. We recently showed, via GISAXS experiments, that organic-coated NPs change their interparticle distance and film structure upon the exposure to volatile organic compounds (VOCs) and dithiols (nonanedithiol) place-exchange.[1] These results can be correlated with changes in current in chemiresistors. In this presentation I will show the importance of the film (NPs film) deposition method and type of organic coating needed for obtaining correlation. I will also discuss how inter-nanoparticles distance and film correlation changes upon different organic coatings (surrounding NPs), substrate type (graphene vs. silicon) increasing temperature, and addition of dithiols.[2]

<sup>[1]</sup> Dalfovo, et al, and Francisco J. Ibañez\*. Real-time Monitoring Distance Changes in Surfactant-coated Au Nanoparticle Films upon Volatile Organic Compounds (VOCs). J. Phys. Chem. C 2015, 119, 5098–5106 [2] Proposal 17155-2014: Dithiols as molecular rulers: the importance of metal nanoparticles separation and graphene towards surface enhanced Raman scattering (SERS).

### Spectroscopic Techniques on the Study of Biosignatures: Degradation of the Heme Group under Environmental Stress

Presenter: Junior J. C. S.

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This study aims to examine preservation and degradation processes of biologically important molecules by spectroscopic techniques. The biomolecule model adopted as biosignature is the class of the porphyrins in their native as well degraded form (particularly the heme group), as this is an ubiquitous molecule on life as we know, and very resilient to degradation on the environment. The analysis of the degradation of heme group is mainly made by the study of the central metal ion behavior, through XANES, and neighboring chemical groups, by molecular spectroscopy techniques - Raman and Infrared. Our initial goal is to better understand the alterations suffered by the central metal ion after being exposed to simulated environmental stress, especially in which conditions the ion substitution happens. This may provide a diagnostic tool to infer the paleoenvironment from a measured substitution rate on fossil porphyrins. Simulations of extreme conditions (such as of the surface of Mars and extreme terrestrial environments), with varying pressure, temperature and incidence of electromagnetic radiation in different energy ranges are being performed in order to improve our understanding of the behavior of the molecules and to test their endurance. An interesting characteristic of porphyrins is the diversity of metal ions that can bind at its center, as in the heme (Fe) and chlorophyll (Mg), which can be replaced by geochemical processes. An example is the change of Fe by V in the heme group. The initial experiments were performed in aqueous media, in which the heme is not soluble, but some vanadium salts are, such as vanadium(III) chloride -VCl3, which is soluble in hydrochloric acid, in which heme is not. Using X-ray Absorption Spectroscopy - XAS (in particular XANES), it was possible to assess the substitution grade, the oxidation state and chemical neighborhood of the central metal ion of heme in the native or degraded forms, as well as in contact with the vanadium solution. Within this scope, synchrotron based XAS is a powerful tool for the characterization of the results of the simulations, as well as for direct measurement of environmental samples containing preserved porphyrin groups. Practically without any need of special sample preparation or environmental simulations, in the first case due to the high brightness of the source and the second energy range and because of flow that can subjecting the sample.

# X-ray back-diffraction pointing to a target soft inelastic X-ray scattering spectrometer

Presenter: Kakuno E. M.

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Soft X-ray back-diffraction (SXBD), X-ray diffraction at angles near and exactly equal to 90° at low energies (~ 3.2 keV) was carried out at Soft X-rays Spectroscopy (SXS) beamline at LaboratorioNacional de Luz Sincrotron (LNLS) [1]. A high-resolution Si(220) multi-bounce back-diffraction monochromator was designed and constructed for this experiment. An ultrathin Si(220) crystal (5 µm thick) was used as the sample. This ultra-thin crystal was characterized by profilometry, rocking-curve measurements and X-ray topography prior to the XBD measurements. It is shown that the measured forward-diffracted beam (o-beam) profiles, taken at different temperatures, are in close agreement with profiles predicted by the extended dynamical theory of X-ray diffraction (2-beam case), with the absence of multiplebeam diffraction (MBD). This is an important result for future studies on the basic properties of back-diffracted X-ray beams at energies slightly above the exact XBD condition (extreme condition where XBD is almost extinguished). Also, the results presented here indicate that stressed crystals behave like ideal strain-free crystals when used for low-energy XBD. This is mainly due to the large widths of SXBD profiles, which lead to a low strain sensitivity in the detection of defects. This result opens up new possibilities for mounting spherical analyzer crystals without degrading the energy resolution, at least for low energies. With these results we aspire to build a soft inelastic X-ray scattering spectrometer, where experiments such as element-specific magnetic imaging tools [2] could be explored.

Hönnicke M.G., Conley, R., Cusatis, C, Kakuno, E. M., Zhou, J., Bouet, N., Marques, J. B., Vicentin, F. C. J. Appl. Cryst. 47 16581665, 2014.
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### Influence of the nanoparticle concentration on the magnetic and structural properties in Fe3O4-PVA nanocomposites

Presenter: Londoño O. M.

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Our aim is to design magnetic nanocomposites in which the dipolar inter-nanoparticles interactions can be tuned. To achieve this, a set of magnetic gels were synthesized, with different nanoparticle concentrations. Polyvinyl alcohol (PVA) was chosen as a containing non-magnetic matrix due to its biocompatible properties. To ensure a correct dispersion of the fine particles within the PVA matrix, these were coated with citric acid (CA), thus the hydrogen bonds formed between the carboxylic (CA) and hydroxyl (PVA) groups promotes a better dispersion and prevent percolation of the nanoparticles [1]. Magnetic results indicated that when the nanoparticles concentration rises, a shift in the blocking temperature (TB) towards higher values is observed. According to the superparamagnetic theory, such behavior is related to an increase in the magnetic anisotropy of the system [2] or an increase in dipolar magnetic interaction between nanoparticles [3]. To discard any structural distortion in the internal structure of the nanoparticles, which could lead to changes in the magnetic anisotropy of the studied systems, we decided to carry out measures of X-ray absorption fine structure (EXAFS and XANES). XAFS results are indicating that the internal structure of the NPs is not affected by the increase in nanoparticle concentration (or inter-nanoparticles interactions). Changes in TB, and in magnetic properties in general are associated with the increase in magnetic interactions, possibly due to the formation of non-compact aggregates. In order to confirm this hypothesis, were performed measures of small angle X-ray scattering (SAXS). SAXS experimental data were fitted using the expression postulated in fractal aggregate model [4]. Fitting results suggest that the increase in the nanoparticle concentration promotes the formation of denser and compact aggregates. We acknowledge the support of CONICET and ANCPyT (Argentina); LNLS, FAPESP and UNICAMP (Brasil); and IFW (Germany).

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### Solving the structure of bimetallic particles with EXAFS

Presenter: López J. M. R.

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We resolved the structure of core-shell bimetallic Co-Pt nanoparticles using Extended X-ray Absorption Fine Structure (EXAFS) spectroscopy at the Co-K and Pt-L3 edges. Using the results of the EXAFS fits at both edges and theoretical models we could determine the core and shell composition as well as the inner and outer radius of two different bimetallic nanoparticles obtained varying the synthesis conditions. The only compatible structure with our results is a CoPt3@Pt core-shell nanoparticle. Our models are supported by X-ray fluorescence, Transmition Electron Microscopy and Small Angle X-ray Scattering results.

# XANES and micro-XRF spectroscopies for chemical characterization of fossil samples

Presenter: Maldanis L.

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The application of novel analytical techniques to study fossils is enabling new and more sophisticated scientific problems to be attacked in paleobiology, both in morphology and chemical composition of the specimens and of the surrounding rock matrix, with none or minimum damage to the sample. Particularly, micro-X-Ray Fluorescence spectroscopy can be applied onto chemical mapping of paleontological specimens, allowing the correlation of chemical findings with morphological features, which can provide a range of information about biochemical preservation of the specimen structures and also about the fossilization process. We applied micro-X-Ray Fluorescence in fossil fishes from the Cretaceous Santana Formation of North-East Brazil and in fixed and resin-included Zebrafish samples, to analyze and correlate the distribution of the elements, that could be associated with biological or diagenetic origin. We also applied XANES spectroscopy to better characterize the iron minerals of the fossil and its matrix, besides look for possible remains of hemoglobin preservation. Our results show the association of elements as iron and zinc with specific structures as bones, gastrointestinal tract and scales, both in the fossil and Zebrafish samples. Other elements, as manganese, appears as a diagenetic marker, distributed nonspecifically in the fossil. We can also observe rare-earth elements as neodymium associated to geochemically modified apatite. XANES spectroscopy of iron showed a clear contribution of pyrite in the matrix and some regions of the fossil sample, but could not identify hemoglobin traces in the fossil sample, probably because of the very low (if present) concentration of this kind of iron among the iron minerals of the sample.

### Trace elements alterations in mammary cells exposed to doses used in mammograms – an investigation using TXRF

#### Presenter: Mantuano A.

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The developments in the diagnosis, and treatment techniques for the breast cancer (BC), which include chemotherapy and/or radiotherapy, increased the patients survival rates for this type of cancer, increasing the chances of late effects due to the treatement. A mammogram is an exam that uses ionizing radiation and aims to obtain radiographic images of the breast for the detection of early stages breast cancer. Considering that during interaction of the radiation with the mammary tissue, important biomoleculesionization and subsequent destabilization may occur, the use of ionizing radiation in health care must be justified (1-5). Most studies concerning the effects using low dose radiation focus, mainly, the alterations in the DNA molecule (6-8). Elemental alterations may be associated with changes in the enzymatic activities of the cell. Our purpose was to quantify the trace elements in the two different cell lines: human breast tumor cells MCF-7 and non-tumor cells MCF-10 after the exposure of doses used for mammograms, considering low breast density (9 mGy) and high breast density (18 Gy), subjected to doses used in mammography and compare the results between those groups. Our study indicated some changes in the concentration of certain elements such as Ca, Fe, Cr and Zn in different tumor cell lines, after different treatments, using the TXRF method. We are trying to identify the potential mechanism of how ionizing radiation can activate cellular targets associated with tumor progression (9-12). MCF-7 and MCF-10 cells were irradiated with doses of 17 mGy and 2Gy in a conventional mammography equipment (Siemens). From the resulting solutions, 5 µL were pipetted on the quartz reflectors used for TXRF analysis. These results can help us to better understand the mechanisms involved in the neoplastic transformations, enzymatic alterations, and biomolecules destabilization that may occur after a breast screening.

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### Core level and valence band electronic structure of Sr2FeMoO6

Presenter: Martins H. P.

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The Sr2FeMoO6 material is a half-metallic ferromagnet with  $\mu \approx 4.0 \mu B$ , and presents a lowfield and high-temperature tunneling magnetoresistance (TMR) effect. Despite extensive studies, there are still open questions about the electronic structure of this compound. The electronic structure of Sr2FeMoO6 was studied using X-ray photoemission spectroscopy (XPS) at the SXS beamline in the LNLS (Campinas, Brazil). The sample was confirmed to be in a pure single state phase and the concentration of antisites was less than 3%. The sample was promptly measured to prevent aging effects and repeatedly scraped with a diamond file to remove surface contamination. The valence band spectrum was taken with a photon energy of 1840 eV. Measurements of the Fe 2p and Mo 3p core levels were also taken. The experimental results were interpreted using cluster model calculations. This model expands the ground state beyond the ionic approximation Fe 3d5 - Mo 4d1 [1], treating the correlation on the Fe and Mo sites in a better suited way, as well as the hybridization between the metal ions and the ligand. The Fe 2p and Mo 3p core level XPS spectra were used to determine the parameter set of the cluster model, which were then fine tuned to describe the valence band spectrum. These results help to better understand the electric and magnetic properties of this compound.

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### Study of a highly crystalline Y2O3 sample by Rietveld and Pair Distribution Function Analysis

Presenter: Martinez, L. G. Martinez, L. G.; Ichikawa, R. U.; Turrillas, X.

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An Y2O3 highly crystalline sample, used as standard reference material for X-ray diffraction experiments, was studied by Rietveld and Pair Distribution Function analysis. The structural parameters obtained using Rietveld refinement, such as cell parameters, atomic positions and isotropic atomic displacement factors were compared previous studies and literature data. Besides that, the high-energy x-rays scattering data, collected at the LNLS-XDS beamline, allowed perform pair distribution function (PDF) analysis. The data was collected in Bragg-Brentano geometry and using a scintillation punctual detector, with 20 keV energy radiation, reaching a Qmax resolution of 20.05 Å-1. The PDF analysis showed that the SRM exhibits a very large structural coherence length, no structural disorder and can be considered perfectly ordered over very large interatomic distances, as expected from a standard reference material.

### Temperature and high-pressure dependent X-ray absorption of SmNiO3 at the Ni K- and Sm-L3 edges

Presenter: Massa N. E.

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The primary consequence of applying quasihydrostatic pressure P is the reduction of interatomic distances, the progressive bond shortening, and the modification of superexchange M-O-M angles. Pressure change as thermodynamic external dynamical variable is a unique tool for probing the relationship between structural sublattice distortions and electronic properties by triggering increments in band hybridization and the eventual emerging into a metallic state by valence and conduction band superposition. SmNiO3 is a member of the family of undoped compounds RNiO3 (R=Rare Earth) in which a sharp temperature driven insulator to metal phase transition takes place. The transition from high temperature orthorhombic metallic to an insulating charge disproportionate  $(2Ni3+ \rightarrow$ Ni3+ $\delta$ +Ni3- $\delta$ ) monoclinic distorted phase is consequence of the Ni-O bond length increase and a simultaneous decrease of the Ni-O-Ni angle. SmNiO3, TIM = 400 K, also has a paramagnetic insulating phase, and orders magnetically below TN~205 K A negative slope, -TMI/dP, is shared by all RNiO3 (R=Rare Earth≠ La.). We report XANES and EXAFS measurements of SmNiO3 from 20 K to 600 K and up to 38 GPa at the Ni K- and Sm L3edges. They were performed at the LNLS in the DXAS energy dispersive beamline. Increasing pressure induces Ni-O-Ni angle increments toward more symmetric Ni3+ octahedra of the rhombohedral R<sup>-3</sup>c space group (metallic LaNiO3). A multiple component pre-edge tail accounts for 1s transitions to 3d-4p states while a post-edge shoulder raises distinctively smoother due to the reduction of electron-phonon interactions as Ni 3d and O 2p orbitals overlap. Pressure dependent room temperature Ni white line peak energies show an abrupt 2.4 GPa valence discontinuity due to non-equivalent Ni sites with Ni3+ $\delta$  + Ni3- $\delta$ charge disproportionate in the monoclinic distortion turning at TMI into Ni3+ of the perovskite distorted orthorhombic Pbnm metal oxide phase. At 20 K, still distinctive, this turns smoother.

### Short-range order study around Iron atoms: Crystallization Process in glassy samples followed by X-ray absorption spectroscopy

Presenter: Mastelaro V.

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Transition metals (TM) and metallic particles are widely incorporated within glass-ceramics (GC) due to two distinct roles: they can affect the nucleation rate dramatically, acting as nucleating agents, or give specific properties to the GC, acting as "active" agents [1-3]. The nucleating agents can be used together or separately to promote simultaneous and related phenomena such as bulk nucleation, decrease in crystallization temperatures and increase in nucleation kinetics, either by accelerating phase separation or by lowering the energy barrier of nucleation. To understand the processes governing the amorphous- to-crystal transformation at the atomic scale, we need to follow the first stages of the nucleation and to identify the structural rearrangements that occur around the nucleating agents. These two aspects can be measured simultaneously by recording the X-ray absorption spectra (XANES, EXAFS) and the X-ray diffraction pattern. However, when the amount of nucleating agent is lower, only the measurement of the XANES and EXAFS spectra is possible. XAS technique was applied to study the nucleating process in a CaMgSi2O6 (diopside) glassy sample which contain iron as nucleating agent [4-5]. The purpose of this study is to better understand the potential of iron as nucleating agent for this particular glass using XAS technique. Samples with 2, 4, 7 and 9% Fe2O3 were nucleated for 8 h and crystallized at Tc (crystallization peak onset) for 10, 20, 30 and 40 min. XANES and EXAFS spectra were collected ex-situ and insitu conditions. The ex-situ experiment was done with samples previously submitted to a controlled heat-treatment. The in-situ experiment was done with the 7% and 9% samples submitted to a heat treatment at the beam line. The analysis of XANES and EXAFS spectra confirms the role of iron in the nucleating process. In-situ experiments allowed to determine the temperature where the crystallization around iron takes place.

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# In situ XPD study of structural changes in iron-cerium mixed oxides under reducing conditions

Presenter: Mazan M. O.

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Although cerium oxide presents good oxygen storage capacity (OSC) and good redox properties, these properties could be improved by doping ceria with trivalent ions like Fe. Doping with alliovalent cations potentially increases the number of anion vacancies improving the oxygen mobility and the reducibility of the material. The resultant solid would be a better material for catalytic applications. Samples with different iron content (Fe:[Ce+Fe] molar ratios= 0, 0.05, 0.1, 0.15, 0.20, 0.30 and 1) were synthesized by Citrates and Liquid-mix methods and calcined at 623K for 2h. The samples were named 0Fe, 5Fe, 10Fe, 15Fe, 20Fe, 30Fe and 100Fe, respectively. The structural changes during the reduction of the \ce{CeFeO {\delta}} system was studied by "in situ" X-ray powder diffraction (XPD), at the D10B-XPD beamline of the Brazilian Synchrotron Light Laboratory, at different temperatures, in an atmosphere containing 5%/ce{H 2}, using a high-intensity and lowresolution configuration. The diffraction patterns were well resolved with high intensity thus allowing the use of Rietveld refinement method. At room temperature all cerium containing samples showed only the peaks corresponding to the fluorite structure of  $ce{CeO 2}$ . However, sample 30Fe presented asymmetric peaks indicating thatsome kind of inhomogeneity could exists. Sample 100Fe presented two phases with hematite-like and At 623K sample 100Fe reduced to pure magnetite while no magnetite-like structures. changes were observed for cerium containing samples. At 923K, all samples excepting pure cerium oxide (0Fe) showed the presence of FeO (wustite). At 1073K, sample 100Fe showed only peaks associated to wustite phase while the mixed oxides presented the peaks corresponding to \ce{CeFeO 3} superimposed with those corresponding to the fluorite structure and very small peaks associated to wustite. No other cerium containing phases were observed during the experiments.

### In situ SR µXRF analysis of Pb in plants used for phytoextraction of soil pollutants

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The phytoextraction technology uses plants to extract toxics metals from contaminated soils and accumulate them in the harvestable parts of the plants, which can then be removed from site. SR µXRF technique offers a powerful approach for probing and mapping the *in situ* distribution of a wide range of elements in plant tissues. In order to understand how Pb and others elements are incorporated into the plant from the soil, we consider essential to use SRmXRF analysis to investigate the spatial distribution of these elements in selected sections of the studied plant. The measurements were carried out at the D09B XRF Fluorescence beamline of the LNLS. The experiments were conducted in Lolium perenne sp. plants, grown in soil contaminated with Pb, and in hydroponics crops exposed to lead at industrial and basal levels. SR µXRF measurements were performed in situ on different parts of the plant (roots and leaves) and in living conditions. Plants specimen were sandwiched between two Ultralene thin films to prevent sample drying or oxidation. The capability for Pb phytoextraction in hydroponics crops versus lead-contaminated soil was compared. The results showed the hydroponics crops of L. perenne sp. can extract and translocate Pb from the ground to the leaves more effectively than plants grown in contaminated soil, where lead mainly stayed in the root. In addition, a spatial correlation between Pb, S and P distributions was observed. The results suggest that further investigations should be done in order to show whether other plants species can offer better efficacy for Pb phytoextraction than the L. perenne sp.

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### Application of XANES spectroscopy to investigate Sb species in corroded bullets crust material oriented to evaluate the potential toxic effects in the environment

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Pb, Sb and others toxic metals from pellets alloy are disperse in the soil of the shooting fields. As long as the corroding bullets are present in soil, secondary Pb and Sb phases in the weathering crusts appears being an important source of bioavailable Pb and Sb. Knowledge on the corrosion mechanism of Sb from the bullet is limited and reports on Sb speciation in soils are still scarce. Preliminary studies by SR-µXRF were performed at the D09B XRF Fluorescence beamline of the LNLS (proposal XAFS1-15234). A positive correlation between Sb and Fe was detected in crust material measured in the outer rim of the weathered bullets due to Sb adsorption to Fe oxyhydroxides of soil. It was also observed a spatial correlation between Sb and Cu and between Sb and Zn in crust. Considering that Sb species have different toxicological properties on the environment, our survey continued performing XANES measurements at the Sb L-edges in order to identify its oxidation states in crust  $(Sb^0)$ , Sb<sup>3+</sup> or Sb<sup>5+</sup>). The measurements were carried out at the D04A SXS Soft X-ray Spectroscopy beamline of the LNLS. Samples consisted of dust crust taken from physically deformed and strongly corroded metallic bullets retained in soil samples sieving from shooting fields of North and East region of Córdoba, Argentina. The results showed that the main species found in all samples was Sb<sup>5+</sup> (Sb2O5) followed by Sb<sup>0</sup> (metallic). Sb<sup>3+</sup> was not detected, and it is known that  $Sb^{3+}$  is more toxic than  $Sb^{5+}$ . The results suggested that in these environmental conditions, pentavalent Sb was the predominant species after weathering of metallic Sb from the corroding bullets.

Acknowledgements: This work was developed at Brazilian National Synchrotron Light Laboratory under the proposal SXS-16953. The authors would like to thank the LNLS staff for its technical support.

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# Amine-alcohol-silicate hybrid matrix as efficient adsorbents for water cleaning

Presenter: Molina E. F.

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Pollution from dye wastewater is becoming a major environmental problem [1-3]. For the removal of anionic dyes, here we employed epoxy-silicate hybrid matrix which are based on polyethers chains covalently linked to silica framework via an amino-alcohol bridge. To prepare the hybrid matrices from 3-glycidoxypropyl-trimethoxysilane and PEO ((O,O'-bis(2aminopropyl)-poly(ethylene oxide)) of molecular weight 500 and 1900 g mol-1, we used the one-pot sol-gel route. The diameter of the xerogels augmented, because the free volume increased as a result of the swollen network. Next, we immersed the PEO xerogels in a solution containing the cationic dye Methylene Blue (MB). However, PEO remained transparent after contact with the MB solution, showing that the xerogels did not adsorb MB. The xerogel efficiently removed the anionic RB dye from the starting solution within a contact time of 120 min; in contrast, the MB concentration in this same solution remained constant throughout this period. The kinetic experiment reveal that equilibrium was reached after 2h with RB dye removal efficiency of 97%). These matrices were able to easily and rapidly separate anionic dyes from a mixture containing anionic along with cationic dyes. The active site of the RB dye and the site groups present in the PEO hybrid matrices played an important role in the capacity of the xerogel to adsorb anionic species. The synthesis of the PEO hybrid is easy to reproduce in any industrial or academic lab. This matrices is potentially applicable as an efficient, fast, selective, and convenient device in water treatment for removal of dyes and metal ions. Financial support: Fapesp (2013/20455-2).

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# Spectroscopic characterization of the interface semiconductor/active layer in sensors based on DNA

Presenter: De Moraes M. O. S.

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The biosensors based on ion-sensitive field-effect transistor (ISFET), as well as the implementation of biosensors based on nanoparticles of TiO2, carbon nanotubes, or graphene occupying a large part of contemporary scientific production(1) having a particular interest for the study of antioxidants compounds(2,5). Sensors based on DNA molecule are high sensitivity and selectivity for the detection of pathogenic and genetic diseases. NEXAFS and XPS techniques were used for the determination which of three kinds of DNA molecules, this is genomic, plasmid and PCR, is more suitable as active layer in biosensors devices. NEXAFS experiments were carried out in the SGM beam line (250-1000 eV) by measuring TEY and the photon flux by Au grid monitor at the LNLS, Campinas-SP. The XPS spectra were obtained at the LaQuiS-IQ, UFRJ. Spectra were obtained in the edges at 1sC, 1sN, 1sO, 2pTi and 2sP before and after the anchorage of the active layer for each type of DNA. For genomic DNA/TiO2, all NEXAFS spectra showed a low intensities and a slight chemical shifts for low energies in relation to others samples. The plasmid DNA/TiO2 and PCR/TiO2 are very similar showing slight differences. In the XPS spectra, genomic DNA presented an opposite behavior, high intensities and shifts to higher energies in relation to others samples. These results indicate a resistive character in charge mobility in the sample genomic DNA/TiO2. Were attributed this resistive characteristics to the presence of histone proteins in structure of genomic DNA even after the removal treatment. A comparative study between the different kinds of anchored DNA was performed to deepen in to the mechanisms of anchoring of the biologically active layers on semiconductor substrate through the NEXAFS and XPS. It's possible to conclude that the genomic DNA is not suitable for the use as the biologically active layer in electrochemical sensors if compared with the anchored mechanism of the plasmid and the PCR DNA molecule.

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### In-situ and in-operando studies of cobalt doped titanates by XRD, XAS and electrochemical impedance spectroscopy simulating working conditions as SOFC anodes and cathodes

Presenter: Napolitano F. R.

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Perovskites with Ti on their B-site has recently been subject of study on the Solid Oxide Fuel Cell (SOFC) field since these materials presents good properties that enables their use as a possible anode materials[1]. At the SOFC cathode side, the (La,Sr)CoO3 is one the most studied cathode material [2]. In this work we present the study of the La0.4Sr0.6(Ti,Co)O3 family (LSTC) for their use as electrode material for new devices with symmetric designs (S-SOFC)[3,4]. In a first step, we completely characterized, for first time, their crystal and electronic structures at room temperature through X-ray diffraction and X-ray Absorption Spectroscopy. This study was followed by non-ambient experiments, simulating in-situ conditions as S-SOFC electrode (oxidizing and reducing atmospheres up to 750°C)[5-7]. We found that cobalt doping increases the electrical conductivity by 5 to 8 orders of magnitude under an oxidizing condition respect to the undoped sample (La0.4Sr0.6TiO3-d), making these materials suitable to be used as cathodes but higher Co doping levels ( $x \le 0.7$ ) decrease the material stability as anode under reducing atmospheres. Besides, Co doping induces a slight rhombohedral distortion in the crystal symmetry at room temperature, favoring a transformation to a cubic phase at higher temperatures (100-500°C, depending of the Co/Ti ratio). On the other hand, recent advances on materials characterization techniques have provided new platforms that enable the in-situ study of crystallographic, mechanical, electronic, and electrochemical properties under a wide range of environmental conditions, not only for fundamental research but also in order to recreate a more realistic environment in the case of applied research [8,11]. Here, we also present preliminary results on LSTC S-SOFC from our recently commissioned characterization device, capable to perform simultaneous XRD and Electrochemical Impedance Spectroscopy measurements.

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# Structural characterization of fragmented kraft lignin by biological processes by SAXS

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The enzymatic fragmentation of lignin was investigated by direct cultivation of basidiomycetes fungus in 10% (v/v) Kraft black liquor (KBL) containing 10 g L-1 dextrose. Four fungi isolates (BT-10, BT-21, BT-40, JAU-3) were grown in KBL, kept under stirring for 15 d at 30 °C. The lignin fractions (LF) were recovered after KBL acidification at pH 3.0 and drying at 60 °C. Prior to SAXS analysis, LF were purified on a gel filtration column Superdex 30 pepgrade (70 cm x 0.6 cm) in an AKTA@ system equipped with UV detector in order to separate the LF from proteins. The LF were eluted and diluted in NaOH solution (0.1 M/pH 11.0). 1 mL was injected into the column at the rate of 0.3 mL.min-1 and detected at 280 nm. The LF analysed by SAXS were selected based on the first and third absorption peaks detected at 280 nm and collected in microtubes by AKTA@ system. SAXS analysis indicated that the profile of the LF obtained after cultivation of BT-10 and BT-40 in KBL were similar, with identical radius gyration values (Rg) (1:58) and very close polydispersity. The profile of the LF biodegraded by JAU-3 and BT-10 (first and second peaks respec.) also were similar to Rg and  $\sigma$  values, however differing in the sizes of fractals (D). Based on the obtained data, SAXS analysis contributed to the structural characterization of LF fragmented by biological processes, regarding to the size, shape and polydispersity.

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### Iterative Reconstruction of Tomographic Images Using Accelerated Projection/Backprojection Techniques

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We will present techniques to speed up iterative reconstruction techniques for tomographic imaging. We will apply the techniques in several sub-optimal acquisition settings. Applications include sparse-view reconstruction and interior-exterior tomographic acquisition with mixed resolution.

## Small angle X-ray scattering applied to Glycoside hydrolases from families GH5 and GH6

Presenter: Neto M. de O.

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Glycoside hydrolases (GHs) are a group of enzymes of great importance for carbohydrate metabolism (HENRISSAT, 1991), and therefore are key enzymes for biotechnological processes, such as the production of biofuels from plant feedstocks (LI et al, 2009). GH5 and GH6 are formerly known as cellulase families A and B, respectively. The cellobiohydrolases of these families are widely believed to act processively from the nonreducing ends of cellulose chains to generate cellobiose. The encoding genes for the cellulases GH5 and GH6 were identified through functional screening of a metagenomics library derived from the soil. Both genes were cloned in a vector of expression pET28a, submitted to heterologous expression on Escherichia coli and purified by affinity chromatography and gel filtration chromatography. New theoretical and computational approaches were developed for the treatment and interpretation of the small angle X-ray scattering (SAXS) data, allowing a detailed analysis of the global structure of proteins in solution (SVERGUN et al, 1996), prediction of quaternary structures (SVERGUN et al, 1998) (OLIVEIRA NETO et al, 2008), among others. In this study we utilize SAXS to characterize two glycoside hydrolases from families GH5 and GH6 of metagenomics origin. Using SAXSMoW (FISCHER et al, 2008), we determine that both enzymes are monomers in solution. Pair distance distribution function p(r) and shape analysis were done using ATSAS package (KONAREV et al. 2006). The maximum diameter (Dmax) of the GH5 enzyme presented a significant decrease in the presence of calcium, suggesting that calcium ions provide a higher stability to the enzyme. GH5 in the presence of calcium ions had a radius of gyration (Rg) of approximately 27 Å and a Dmax of 90 Å and in the absence, respectively, 41.8 Å and 120 Å.

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### In situ study of austenite decomposition during thermal cycles and under application of stress in ferrous alloys

Presenter: Nishikawa A. S.

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In this work, the kinetics of austenite decomposition during the application of the novels thermomechanical treatments of Quenching and Partitioning (Q&P) and Hot Stamping + Quenching and Partitioning (HSQP) on a ductile cast iron and a TRIP steel, and also during the application of tensile stress in a metastable austenitic steel were studied by means of in situ synchrotron X-ray diffraction. Both experiments were performed at the XTMS experimental station facilities at LNLS/LNNano using a Gleeble thermomechanical simulator integrated to the XRD1 beamline. Real time information about kinetics of competitive reactions was obtained by following the lattice parameter and phase fractions changes. The results were validated by global and local measurements techniques, e.g., high resolution dilatometry and electron microscopy, and also confronted with information on literature. The results demonstrated to be in fairly agreement with the expected from previous experiments and also provided complimentary information about the bainite and martensite reactions occurring in the studied samples.

### Synchrotron small angle X-ray scattering Investigation of niobium oxyhydroxide nanostructured

Presenter: Pereira I. M.

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Our research group has developed an material based on niobium. They are: (i) hydroxide niobium (NbO2OH), (ii) niobium oxyhydroxide (NbO2OH/H2O2) and (iii) amphiphilic niobium oxyhydroxide (NbO2OH/amp). In this work SAXS was employed to explore the niobium nanoparticle form. The symmetric maxima observed at NbO2OH, NbO2OH/H2O2 curve suggests the existence of small spheroid like particles. Results suggstes that the O-H surface groups increase nanoparticle size without altering significantly the particle size. The asymmetric maximum presented on NbO2OH/amp curve and the second-order peak presented as a 'bump' at larger q-values suggests elongated lamellar structure. The niobium compound plots shows: a well defined bump around 2.9 nm-1, attributed to NbO2OH particles, an overlaid bump around 4.0 nm-1, attributed to O-H surface groups and, observed at NbO2OH/amp, a defined peak at 1.658 nm-1 attributed to the surfactant molecule. The Porod's Law negative deviation is not observed at NbO2OH/H2O2, apparently, the compact boundary is the result of the hydrogen bonds between the hydrogen atoms of the hydroxyl groups and oxygen atoms. On the other hand, the Porod's plot of NbO2OH/amp showed more complex information, suggesting that the hybrid system has significantly different surface composition. The strong negative deviation is due to the diffuse-boundary and due the electrostatic interaction between the surfactant organic molecular chains and Nb=O, Nb–O or Nb(O-O) groups which leads the interfaces to stimulate the negative deviation. The interphase thickness,  $\sigma$ , of NbO2OH is 0.06. On the other hand, NbO2OH/amp  $\sigma$  is 0.23.

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### Investigation of the Morphology Exhibited by Multilayered Films of Collagen and Cellulose Nanowhiskers by Small-angle X-ray Scattering

Presenter: Pereira I. M.

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Collagen (COL)/glycerol (GLI)/cellulose nanocrystal (CNC) nanocomposites were prepared by solution blend. The blends were obtained by integrating the COL solution with GLI/CNC solution and varying the CNC. Films were produced by casting the dispersions in a PVC mold and allowing them to dry at 400 C. Synchrotron small angle X-ray scattering (SAXS) was employed: (i) to investigate the macromolecular structure of neat COL/GLI and COL/GLI/CNC nanocomposites, (ii) to explore, in water solution, the CNC form and dispersion and (iii) to investigate the lyophilized cellulose whiskers surface. SAXS results indicated that CNC in concentrations up to 1 wt. % present an intermediate shape between a rod and a plane with 9.34 nm radius of gyration (Rg). Apparently, when dispersed in water, the CNC rod-like particle expands to a planar shape with a very smooth particle surface. SAXS patterns of lyophilized CNC present anisotropic feature typically observed for rod like particles. Lyophilized CNC scattering curve shows only one power-law regime, alpha = -3.7, which describe a surface fractal structure formed by CNC clusters with irregular surface. The COL/GLI/CNC nanocomposites present two different structural levels with two types of particles with very different Rg. At the intermediate power-law regime, it is observed a largescale mass fractal aggregate. In the high power-law regime, it is observed scattering from primary particles smaller the 1 nm. Besides, the primary particle of net COL/GLI is a rod where the shrunken CNC rods attach. As the CNC concentration increases the original particle distorts from a rod to a plate. As expected, the CNC concentration controls the particle radius because while the CNC is introduced into the system it attaches to the original primary particle which raises the particle radius. From the TEM data, it is not possible determine the dimensions of the isolated particles. However, the smaller cluster would have less than 3 nm in radius.

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### Use of oyster Crassostrea rhizophorae as biomonitor in analysis of heavy metals pollution in the marine environment under influence of the ports of Santos and São Sebastião.

Presenter: Pezzatti R. R.

Pezzatti R. R.<sup>1</sup>, Yokoyama L. Q.<sup>1</sup>, Ignacio B. L.<sup>1</sup>, De Jesus E. F. O.<sup>2</sup>, Mársico E. T.<sup>3</sup>, Ribeiro R. de O. R.<sup>3</sup>, Barbosa R. de F.<sup>1</sup>

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All living organisms change the environment in which they are inserted. Hover, the human is changing the environment with higher intensity compared with other living beings. The coastal region is, due to the high population concentration and polluting industrial activities, the subject of considerable emission of heavy metals, suggesting potential effects on their populations, the marine environment as a whole and to the human population that consumes (whether or routinely) organisms in in this ecosystem. Thus, this study had the objective to examine the contractions of Ca, Cr, Cu, Fe, K, Mn e Zn in two different tissues of Crassostrea rhizophorae, the adductor muscle of the shells and the edges of the mantle. These organisms were collected from four areas between the ports of Santos and São Sebastião, Milionaries Beach, Boracéia Beach, Guaecá Beach and Araçá Bay. The metal concentrations were determined by the technique of X-ray fluorescence by Total Reflection (TXRF). Through the obtained concentrations, we observed elevated concentrations of metals compared with other studies conducted by the scientific community, so that such sites are admittedly influenced by polluting activities of these two ports. Moreover, compared to the maximum allowable limits for human ingestion, such sites had higher concentrations of metals, which can pose risks to human health, since several people have the habit of consuming oysters as food.

### Refinement of single crystal structures by X-ray multiple diffraction.

Presenter: Remédios C. M. R.

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The methods of X-ray, neutron and electron diffraction are of fundamental importance in crystallography where resolution in determining crystal structures relies primarily on refinement procedures. The collection of large data sets of diffracted intensities and adjustment of parameters in model structures to simulate the experimental intensities are the very basic steps common to all refinement procedures. Apart from these procedures, there are also validation tools necessary in structural biology to avoid serious errors when resolving macromolecular crystals from electron-density maps (Read et al., 2011). Structural resolution in crystallography in the 21st century is therefore limited to structural details producing an unambiguous set of diffracted intensities in atomic models presenting physical and chemical consistency. In this work A pair of enantiomer crystals is used to demonstrate how X-ray phase measurements provide reliable information for absolute identification and improvement of atomic model structures. Reliable phase measurements are possible thanks to the existence of intervals of phase values that are clearly distinguishable beyond instrumental effects. Because of the high susceptibility of phase values to structural details, accurate model structures were necessary for succeeding with this demonstration. It shows a route for exploiting physical phase measurements in the crystallography of more complex crystals.

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### Thermal transformations metakaolin – spinel type aluminosilicate: Al and Si k-xanes characterization

Presenter: Requejo F. G.

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Since the pioneering work of G.W. Brindley and M. Nakahira it is known that one of the open problems concerning the metakaolin phase, the manner in which it transforms into a spinel-type phase and mullite, and the relation of this spinel-type phase to mullite, because the series of reactions by which kaolinite transforms to mullite is perhaps the most important in the entire field of ceramic technology [1]. The thermal transformations of kaolinite (K) in metakaolin (MK) and the metakaolin in a spinel type aluminosilicate (SAS) were studied by Al and Si K-XANES to correlate the information obtained with this technique with that obtained by conventional X-ray Diffraction, and clarify and assign the structural changes. Simultaneously with XANES technique is possible to observe changes in the electronic structure of the system. For K à MK transformation, by Al K-XANES it was obtained results consistently with reported Aluminium coordination changes occurred in the kaolinite dehidroxylation process [2]. In general, Al K-XANES spectra of aluminum-containing oxides showed three distinguishable peaks: one at 1566.0 eV assignable to tetrahedral AlO4 and two assigned to AlO6 characteristic at 1568.0 and 1572.0 eV [3]. In the case of SAS there are several features, particularly there is a shoulder at 1566 eV and two AlO6-peaks one at 1567.8 and other at 1571.4 eV. No major changes in the Si K-XANES were observed, i.e. the environments of Si it is not primarily affected for thermal transformations, except for a slight variation in the 3p-holes density. This leads to the conclusion that the structural changes are attributable to changes in the Al environment.

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### Study of Eu3+! Eu2+ reduction in BaAl2O4: Eu prepared in different gas atmospheres

Presenter: Rezende M. V. dos S.

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The effect of different gas atmospheres such as H2(g), synthetic air, carbon monoxide (CO) and nitrogen (N2) on the Eu3+! Eu2+ reduction process during the synthesis of Eu-doped BaAl2O4 was studied using synchrotron radiation. The Eu3+! Eu2+ reduction was monitored analyzing XANES region when the sample are excited at the Eu LIII-edge. The results show that the hydrogen reducing agent are the most appropriate gas for Eu2+ stabilization in BaAl2O4 and that only a part of the Eu ions can be stabilized in the divalent state. A model of Eu reduction process, based on the incorporation of charge compensation defects, is proposed.

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# Ionization and fragmentation of the acetaldehyde (CH3CHO) molecule by 20-330 eV photons

Presenter: Ribeiro L. C.

Ribeiro L. C.; Santos M. de J.; Dos Santos A. M.; Arruda M. S.; Mendes L. A. V.; Dos Santos A. C. F.; Marinho R. dos R. T.; Prudente F. V.

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The aim of this proposal is to perform a study of the ionization and fragmentation of the acetaldehyde molecule (CH3CHO) in the gas phase after interaction with 20-330 eV photons. The acetaldehyde is a prebiotic molecule. From the physical and chemical point of view, the study of the origin of life is related to the formation of biomolecules, as the nucleic acids (DNA and RNA) and the proteins. A hypothesis to explain the origin of life suggests that the biomolecules were created in space by the recombination of simpler organic molecules subjected to radiation and collisions with the solar wind particles, such as protons, electrons and ions, and then brought to Earth by meteors and comets. Therefore, the study of simple prebiotic molecules is of fundamental importance to understand the formation of complex organic molecules. The experimental set up includes the time-of-flight mass spectrometer assembled at the rotating chamber. The chamber needs to be in ultra high vacuum and liquid nitrogen traps were used for the mechanical pumps. The molecule were inserted into the chamber through the gas inlet system. To take into account the influence of the higher harmonics orders we measured the photoionization for a known gas, the N2. We have determined the branching ratios for fragmentation products as a function of the energy of the projectiles. We have measured the total (TIY) and partial (PIY) ion yields, as well as the single (PEPICO) and double (PEPIPICO) mass spectra, for interactions with photons between 20 eV and 330 eV. Our group had previously measured some prebiotic molecules in the low (TGM beamline) and high (SGM beamline) energy range (for example, see reference [1]). Besides comparing the results of the interaction with different projectiles, with this work we could also provide information about the photofragmentation process with energies from opening channels until core excitation.

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### Heavy metals measurement in sandy beaches: influence of the benthic fauna associated

Presenter: Ribeiro R. M. M.

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Heavy metals in trace amounts are naturally found in the environment some of them are essentials to life, however, they cause toxicity at high concentrations1. The absorption of heavy metals on intertidal sandy beaches sediment is related to the contamination of its inherent biota2. It may affect not only on the metal concentration on body tissues but also on the diversity and distribution of these organisms. The scientific literature did not present results that consider the influence of benthic fauna at concentrations of trace metals in the sediment. Based on the literature about heavy metals contamination at sediment from sandy beaches, this work aims to analyze the influence of the benthic fauna (macro- and meiofauna) in the quantification of trace metal contents of sandy beaches. Sediment from the intertidal regions of Itararé, located at the São Paulo State coast (southeast Brazil), will be sampled and analysed for heavy metal contents and granulometric composition. The collected sediment will get through different processes for comparison: the first will not be processed; the second one will be sieved to macrofauna's separation. Macrofauna will be sorted out by sieving the sediment through a 0.5 mm mesh. Heavy metals concentrations will be analyzed using TXRF technique. Among the metals found this work Ag, Ca, Cr, Cu, Fe, K, Mn P, S Si, Ti e Zn and some of these have a possible contributions in concentrations of metals in benthic macrofauna and coarse and very coarse sediment were found. The metals were analyzed to find out the main contribution of each element in each fraction of the sediment and organisms. Keywords: TXRF. Sediment. Macrofauna. Metals.

### Exploratory Methodology for Retrieving Oxidation State Information from X-ray Resonant Raman Scattering Spectrometry

Presenter: Robledo J. I.

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It has been observed recently that the resonant Raman scattering (RRS) peak of an X- ray spectrum contains information about the chemical environment of the irradiated matter [1]. This information is extracted with complex processing of the spectrum data. Principal component analysis (PCA) [2] is a statistical multivariate technique that allows exploring the variance–covariance structure of a set of data. This methodology can be applied to obtain information from any kind of spectra (In particular RRS spectra). To analyze its potentiality, several measurements of different oxides in surface nanolayers were measured in total reflection conditions using synchrotron radiation from the LNLS, in Campinas, Brazil. PCA was used to obtain the information encrypted in the RRS peak, and to establish a new methodology [3]. The results show that multivariate analysis techniques are suitable for the analysis of this kind of spectra, foreseeing its application in future research.

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Robledo, J. I.; Sánchez, H. J.; Leani, J. J.; Pérez, C. A., Anal. Chem. 2015, 87, 3639-3645.

## The effect of annealing on the electronic structure, morphology and charge transport in polymer: fullerene blends for photovoltaics

Presenter: Rocco M. L.

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In the last decade internal donor-acceptor copolymers emerged as the most promising polymers for organic solar cell applications. Photovoltaic devices with power conversion efficiency (PCE) higher than 10% have been achieved using these polymers as active layer. Several reports show good photovoltaic device characteristics especially after thermal annealing. Therefore, it is highly important to investigate the effect of thermal annealing on the electronic structure, morphology and charge transfer dynamics in these polymers and their blends. For that we applied Near-edge X-ray Absorption Fine Structure (NEXAFS), Xray Photoelectron Spectroscopy (XPS) and Resonant Auger Spectroscopy (RAS) in the context of the core-hole approach for the poly[2,7-(9,9-bis(2-ethylhexyl)-dibenzosilole)-alt-4,7-bis(thiophen-2-yl)benzo-2,1,3-thiadiazole] (PSiF-DBT) polymer and its blend with fullerene. The effect of the thermal annealing treatment at 100 and 200 °C on the electronic structure, charge transfer delocalization times and molecular orientation were probed. Edgeon and plane-on molecular orientations with respect to the substrate surface were measured for the thiophene and benzothiadiazole units, respectively, using angular dependent NEXAFS spectra at the S K-absorption edge. Molecular orientation of the silafluorene unit was also probed by NEXAFS at the Si K-edge. The improvement of the polymer ordering with annealing was evaluated by NEXAFS. Femtosecond charge transfer times were measured. Differences in charge transfer times at Si and S K-edges may be related to the localizeddelocalized character of the molecular orbitals involved in these excitation processes, which was corroborated by theoretical calculations, with explicit relaxation of molecular orbitals due to the core-hole. The authors would like to acknowledge CNPq, CAPES, LNLS and FAPERJ.

### Iron oxide nanoparticles coated with different Silica thicknesses: SAXS analysis of size, shape and agglomeration and its relationship with magnetic properties.

Presenter: Rojas P. C. R.

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Iron oxide nanoparticles were synthesized via high temperature decomposition of organic precursors, and the obtained nanoparticles were then coated with SiO2 via the reverse microemulsion method, attempting to regulate the coating thickness by varying the amount of tetraethyl orthosilicate (TEOS) and the reaction time. As a result five core-shell systems of single magnetic core (of fixed composition, structure, shape and diameter) with different silica shell thicknesses are presented here. TEM images corroborate the formation of such structures, with nearly-uniform spherical cores of mean diameter of 7.2nm and standard deviation of 0.2nm. The shell thicknesses increase with the amount of TEOS, reaching average values near 18nm for the highest TEOS amount. These values are in good agreement with those obtained by SAXS after analyzing the patterns with a core-shell model. The magnetic dipolar interaction strength dependence with the shell thickness was studied with static magnetic measurements, where the blocking temperature estimated from the temperature dependence of magnetization (FC/ZFC curves) decreases as the shell thickness increase, as expected for progressively less interacting systems. And the magnetization as a function of applied field curves show in all cases the behavior of a superparamagnetic system with interactions above the blocking temperature, according to the Interacting Superparamagnetic model formulated by Allia. The relationship between the size and morphological information provided by SAXS and the magnetic properties is exhaustively analyzed. Acknowledgements: authors would like to acknowledge CONICET (Argentina) for financial support and CNPEM (Brasil) for the use of synchrotron light (SAXS line, LNLS) and TEM facilities (LNNano).

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### Electronic Studies on Coordination Metal Complexes with Xanthates Ligands: S, Ni and Mn K-edge XANES

Presenter: Romano R. M.

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In this work, and as part of a general project aimed to the preparation and study of coordination complexes with potential pharmacological applications, we present the results of sulfur, nickel and manganese K-edge XANES study on a series of Ni and Mn coordination complexes with xanthate ligands and their potassium salts. Twenty three samples, with general formula (ROC(S)S-K+), (M(ROC(S)S)2) and (M(ROC(S)S)2(N-donor)1or2), with M = Ni, Mn; R = Me, Et, nPr, iPr, nBu, CF3CH2- and N-donor = py, bipy, phen, were prepared by our group in La Plata and characterized using vibrational (IR and Raman) and UV-visible spectroscopy, and also by single-crystal X-Ray diffraction analysis. The sulfur K-edge Total Ion Yield (TIY) spectra were investigated in the SXS beam-line under the proposal 16063 while the Ni and Mn K-edge XANES spectra were measured in the XAFS-1 beam-line under the proposal 17950. The XANES spectra of potassium xanthate salts (ROC(S)SK) present three resonances below the ionization energy of the 1s electrons of the sulfur atom, at 2468.3  $\pm 0.1$ , 2469.0  $\pm 0.1$  y 2470.6  $\pm 0.2$  eV, which were assigned to S1s $\rightarrow \pi^*C=S$  and S1s $\rightarrow \sigma^*C-S$ transitions. In the nickel complexes, besides transitions to antibonding orbitals of the ligands, a new band around 2469.6 eV, assigned to a S1s $\rightarrow \sigma^*$ S–Ni transition, was observed. The Ni K-edge spectra of the studied complexes present several resonances. The pre-edge peak around 8330 eV corresponds to the transition of the nickel 1s electron to its empty 3d orbitals. The low-intensity of this resonance is in agreement with its essentially forbidden character by the dipole selection rule. The most-intense peak, assigned to a Ni1s→Ni4p transition, appears in the region of 8340 to 8350 eV and shows the most significant difference in both its intensity and the chemical shift, in dependence on the ligand alkyl sustitutent. The Mn K-edge spectra present two peaks, the pre-edge resonance around 6540 eV and the main feature close to 6550 eV.

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### Atomic Pair Distribution Function at LNLS: A New Tool for Material Science

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The atomic pair distribution function (PDF) method is an attractive alternative to the X-ray absorption and diffraction methods for studying the local and medium-range atomic arrangements in material science. [1,2] In the PDF method, Bragg diffraction and diffuse scattering intensities are measured and the structural information is obtained by a Fourier transformation to real space. This technique can identify local atomic order much beyond the nearest neighbor atoms. With this powerful technique we can detect small local distortions that conventional of X-ray diffraction studies miss, since the sharp Bragg peaks only reflect the information about the average structure.[1,2] The XDS beamline of the LNLS, was designed to take advantage of the 4T superconducting multipole wiggler inserted in the storage ring. This multipurpose beam line is employed for X-ray diffraction and X-ray absorption spectroscopy in the energy range between 5 and 30 keV. The X-ray total scattering experiment can be perfomed in two different arrangements: a) Bragg-Brentano configuration, using a scintillation detector with an analyzer and b) Debye-Scherrer configuration, where the sample is mounted into capillaries and the diffraction pattern is acquired with an arrangement of 6 Mythen detectors or a scintillator. The sample can be measured at different atmospheres and temperatures. To study the viability of the beamline for this kind of experiment were acquired data of two standard materials at LNLS and then we validated our experiment measuring the same material at a dedicated PDF beamline of the Advanced Photon Source (APS). Our results are very promising. With our configuration we can study both the local order and the meso-scale (r up to 300Å). In this work we summarize our first results on the standards, ferroelectrics and superconductor materials

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#### Multivariate SAXS Profiles Analysis Applied to Synthesis of Heterogeneous Titania Photocatalysts by Sol-Gel Method

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The Persistent Organic Pollutants (POPs) that result from some industrial processes are a major source of water pollution, and they may create health problems for humans. In recent years, Advanced Oxidation Processes (AOPs) have been presented as an alternative to conventional methods. Among all of the AOPs, the degradation of organic pollutants by heterogeneous photocatalysis based on titania is one of the most successful applications that uses this technology for recalcitrant pollutant degradation in waste water at ambient conditions [1]. In parallel, silica-based catalyst supports can be produced by sol-gel technology and have been widely used for catalysis applications [2, 3]. Nevertheless, to our knowledge, silica nanoparticles have not been employed as supports to remove POPs in water. In the present investigation, microporous nanoparticles and mixed-structured silicas were employed as supports for photocatalysts. TiCl4 was used as the titanium precursor in the preparation of the nano- and mixed heterogeneous supported titania photocatalysts. For comparative reasons, photocatalytic tests were carried out with commercial titania (P25). The systems were evaluated in the photodegradation of Rhodamine B (RhB). The solids were characterized by nitrogen porosimetry, small-angle X-ray scattering (SAXS), zeta potential (ZP), dynamic light scattering (DLS) and diffuse reflectance spectroscopy in the ultraviolet (DRS-UV). All intensity profiles of X-ray scattering were treated by multivariate analysis tools [4], using the Hierarchical Cluster Analysis (HCA) and the Principal Components Analysis (PCA) in the modulus of scattering vector (q) region from 0.04 to 4.0 nm-1. The supported nanometric photocatalyst prepared with silica nanoparticles achieved the highest percentage of RhB degradation under the ultraviolet (68.0%) and visible (45.1%) radiation. The photocatalyst activity of nanometric catalyst was better than the commercial P25 (12.3% under visible radiation.

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### The importance of the active site molecular interactions to the oligomerization and reactivity of the typical 2-Cys Prx

Presenter: Dos Santos M. C.

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Typical 2-Cys Peroxiredoxins (Prx) are antioxidant proteins able to decompose hydroperoxides with second-order rate constants of 10<sup>6</sup>-10<sup>8</sup> /Ms. To decompose hydroperoxides, typical 2-Cys Prx use a Cys residue (peroxidatic cysteine - CP) and, during catalysis, it is oxidized to sulfenic acid (CP-SOH), which condenses with a second Cys residue (resolving cysteine - CR) to form a disulfide. These enzymes are obligate dimers which can associate in  $(\alpha 2)5$  decamers in response to several factors, as the redox state. In reduced form, these enzymes are decamers and oxididation to disulfide results in their dissociation in dimers. The high reactivity of Prx is related to the activation of CP by two neighboring residues: Thr/Ser and Arg, the so-called catalytic triad, that allow the establishment of a transition state. The analysis of Tsa1 crystallographic structure, a typical 2-Cys Prx from yeast, revealed that the active site lies in the dimers boundaries in the decamer. Thereby, a hydrogen bond (CH- $\pi$ ) between Tyr77 residue and Thr44 from catalytic triad of adjacent dimer, very conserved among the Prx, apparently positions the Thr favorably to perform the interactions involved in the transition state establishment. To shade a light in this hypothesis, we generated mutant enzymes carrying the replacement of Thr44 by Val (Tsa1T44V) or the Tyr77 by Ala (Tsa1Y77A). Analysis by size exclusion chromatography revealed that both substitutions resulted in decamer destabilization and the evaluation of the catalytic activity of by steady state kinetics and fluorometric approaches showed a high decrease of the catalytic efficiency to both mutants. Together, our results evince the importance of molecular interactions between Tyr77 and Thr44 in the 2-Cys Prx oligomerization and the full catalytic activity. To gain insights of the substitution effects over Tsal active site environment, crystallization trials were performed and the refinement of promising initial conditions are in progres.

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#### Microemulsions for application as corrosion inhibition: a SAXS approach

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Microemulsions (ME) are clear, stable and isotropic mixtures of oil, water and surfactant, frequently in combination with a cosurfactant. Microemulsions are a class of microheterogeneous systems having unique features of stability, solubilization capacity, structural morphology, physical property and applicability. They have found numerous applications in different fields from drug delivery vehicles up to corrosion inhibitors [1,2,3]. In the latter, microemulsions have generated considerable interest over the years. Advantages associated with their thermodynamic stability, optical clarity and ease of preparation enable corrosive agents to be solubilized, surfactants to be adsorbed on metal surface and hence the corrosion is reduced. Depending on the types of oil and surfactant and environmental conditions, microemulsioned systems of varied categories, consistence and internal structures may result. This study aims to obtain microemulsioned systems using vegetable inhibitors based on A. muricata L. (Am) and A. squamosa L. (As) extracts, (Annonaceae) as oil phases (FO). Tween80 and ethanol were used, respectively, as surfactant and cosurfactant and saline NaCl 3.5% (corrosive environment) and aqueous phase (AP). The formation of ME were analyzed and verified by SAXS and rheology allowing such systems are promising corrosion inhibitors. The results shows a broad peak typical of microemulsion systems with a maximum at approximately q = 1nm-1 for all samples. Quantitative structural information was obtained from the ellipsoidal "core-shell" model. The adjustment of the model provided information about the maximum radius (Rmax) and minimum (R min) and therefore the size of the ME.

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#### Phase Contrast X-ray Imaging of Human Peripheral Nerves

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Diagnostic imaging techniques play an important role in assessing the exact location, cause, and extent of a nerve lesion, thus allowing clinicians to diagnose and manage more effectively a variety of pathological conditions such as entrapment syndromes, traumatic injuries, and space-occupying lesions. Ultrasound and nuclear magnetic resonance imaging are becoming useful methods for this purpose, but they still lack spatial resolution. In this regard, recent phase contrast x-ray imaging experiments of peripheral nerve allowed the visualization of each nerve fiber surrounded by its myelin sheath as clearly as optical microscopy. In the present study, we attempted to produce high-resolution x-ray phase contrast images of a human sciatic nerve by using synchrotron radiation propagation-based imaging. The images showed high contrast and high spatial resolution, allowing clear identification of each fascicle structure and surrounding connective tissue. The outstanding result is the detection of such structures by phase contrast x-ray tomography of a thick human sciatic nerve section. This may further enable the identification of diverse pathological patterns, such as Wallerian degeneration, hypertrophic neuropathy, inflammatory infiltration, leprosy neuropathy and amyloid deposits. To the best of our knowledge, this is the first successful phase contrast x-ray imaging experiment of a human peripheral nerve sample. Our long-term goal is to develop peripheral nerve imaging methods that could supersede biopsy procedures.

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### Structural 3D Characterization of Silica, Zirconia and Titania Monoliths and Columns for Capillary Liquid Chromatography

Presenter: Da Silva C. G. A.

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The morphology and the pore structure of a monolithic chromatographic bed are very important features in the design of stationary phases since these aspects directly influence the hydrodynamic properties (e.g., flow properties), thermodynamic properties (e.g., loadability) and the mass transfer kinetics (e.g., efficiency) [1]. Imaging techniques like scanning (SEM) and transmission (TEM) electron microscopy rapidly provide direct information about monolith morphology from bulk or cross sectional capillary images. However, SEM images offer no reliable depth information and thus lack the required morphological details [2]. In this way, a 3D micro tomography images of silica (SiO<sub>2</sub>), zirconia (ZrO<sub>2</sub>) and titania (TiO<sub>2</sub>) monoliths prepared by the sol-gel process (SGP) using different experimental conditions and their columns were obtained. The 3D imaging information obtained by acquiring projection images of the sample along a number of different directions greatly improved our knowledge about the morphology of monoliths inside the capillaries and in the bulk and helped to understand and optimize the synthesis of the materials as well as improved the understanding the chromatographic performance of the columns.

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# Near edge structure at the lithium K-edge in LiH by inelastic X-ray scattering

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The excitation of core electrons of Li in LiH was studied by means of inelastic x-ray scattering spectroscopy (IXSS) at low and high momentum transfer. Measurements were carried out at the XDS beamline of LNLS. A Johann type spectrometer based on a spherical analyzer crystal [1] was used. The core electron contribution was extracted from the whole electronic excitation spectrum. At low momentum transfer the near edge fine structure of the spectrum is interpreted in terms of dipole allowed transitions to empty states. At higher momenta an additional structure was observed, which was assigned to the opening of excitation channels other than dipolar. A chemical shift of the Li K-edge of 2.2 eV to higher energies relative to Li metal was measured. Calculation of the density of states and simulations of the excitation spectrum were made using the FEFF code [2]. Core-hole effects were found to be appreciable and need to be taken into account in the simulations in order to reproduce the observed spectral features. The present experiment demonstrates the feasibility of investigating near edge structures of light elements by means of IXSS with high energy resolution at the XDS beamline.

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### Characterization of nanostructured A1-xSrxFe0.8Cu0.2O3-d perovskites (A=La, Ba) as IT-SOFC cathodes.

Presenter: Suescun L.

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Solid Oxide Fuel Cells operating in the 500-800 °C temperaure range (IT-SOFCs) require the development of new cathode materials with high mixed ionic-electronic conductivity and good catalytic activity of the oxygen reduction reaction (ORR).[1] Traditional La1xSrxMnO3-d perovskite material does not operate well at low temperatures due to the absence of oxygen conductivity and strong reduction of ORR activity. Cobaltites Sm0.5Sr0.5CoO3-d and related compounds show excelente conducting properties and are good ORR catalysts, but they show a large thermal expansion coefficient (TEC) and high reactivity towards comon electrolyte materials which prevents their efficient aplication as IT-SOFC cathodes.[2] In recent years our group has been exploring the preparation of cobaltfree, nanostructured cathode materials, with good ORR activity based on Fe-Cu perovskites. We have designed a novel gel-combustion technique that allowed the preparation of Ba0.5Sr0.5Fe0.8Cu0.2O3-d (BSFCu) and La0.6Sr0.4Fe0.8Cu0.2O3-d (LSFCu) materials, compatible with standard Ce0.9Gd0.1O1.95 (CGO) electrolyte and characterized its phase evolution with temperature, TEC, microstructure and electrochemical performance in the the 500 - 700 C temperature range of symmetrical cells with the configuration cathode/CGO/cathode prepared at different temperatures. Both LSFCu and and BSFCu materials show very promising ASR values of 0.035 and 0.15 ohm.cm<sup>2</sup> at 700 °C. Structural and microstructural characterization of the materials show that BSFCu shows a very isotropic and homogeneous arrangement of nanoparticles 18nm wide and a cubic Pm-3m structure at all studied temperatures while LSFCu shows larger particles with average size of 41 nm (as determined by TEM) with rhombohedral R-3c space group symmetry at RT, that transforms to cubic Pm-3m above 375 °C. This phase transition is accompanied by a significant change in the TEC that suggest BSFCu is a better candidate material for IT-SOFC cathodes [3,4].

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# Study of Ag@Fe3O4 nano-heterostructures by synchrotron radiation techniques

Presenter: Tancredi P.

Tancredi P.<sup>1</sup>; Moscoso-Londoño O.<sup>1</sup>; Muraca D.<sup>2</sup>; Pirota K.<sup>2</sup>; Knobel M.<sup>2</sup>; Wolff U.<sup>3</sup>; Damm C.<sup>4</sup>; Neu V.<sup>3</sup>; Rellinghaus B.4; Socolovsky L.M.<sup>1</sup>

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Nanostructured samples of Ag@Fe3O4 with controlled morphology were prepared by chemical synthesis. The bi-phase nanoparticles were obtained by decomposition and epitaxial growth of Fe3O4 over previously prepared Ag seeds. By this reaction pathway, the morphology of the samples can be tuned as a consequence of the electron-donor capacity of the employed solvent, either 1-octadecene, phenyl ether or a mixture of both. In this way, two kinds of nanoparticles were prepared and observed by microscopy techniques: dimer-like or flower-like. Further structural characterization was performed by synchrotron radiationrelated techniques in order to study the morphology of the nanoparticles (SAXS) and possible structural distortions at the interface caused by the distinct coupling of Fe3O4 at the Ag seeds in flowers and dimers (XANES and EXAFS). Also, Fe3O4 nanoparticles prepared without the Ag seeds were studied and characterized to compare the single phase material properties In previous works, we have studied how with the Ag@Fe3O4 coupled product properties. the nanostructure final shape influences the magnetic properties of the sample, and we found significant differences in the magnetic behavior between flower-like and dimer-like Ag@Fe3O4, and isolated Fe3O4 nanoparticles[1]. In this sense, complementary analysis derived from synchrotron radiation-related techniques provides helpful evidence to try to understand and relate structural features with the observed magnetic behavior. Acknowledgements: Argentinian agency CONICET and LNLS-CNPEM.

[1] O. Moscoso-Londoño, D. Muraca, P. Tancredi, C. Cosio-Castañeda , Kleber R. Pirota, L. M. Socolovsky; Physicochemical Studies of Complex Silver–Magnetite Nanoheterodimers with Controlled Morphology; J. Phys. Chem. C (2014), 118, 13168–13176

# Investigation of band structure of insulators using vacuum ultraviolet spectroscopy

Presenter: Teixeira V. de C.

Teixeira V. de C.1; Valerio M. E. G.2

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Vacuum ultraviolet spectroscopy is a powerful tool to investigate the band structure of semiconductors and insulators. At this work some studies were performed to investigate the optical band gap of calcium aluminosilicates (CAS) used as matrix to prepare luminescent materials. For this, it was used a special geometry for optical measurements the TGM (Toroidal Grating Monochromator) beamline at the LNLS (Synchrotron Light National Laboratory). The main aim of this work is to better understand the band structure of CAS and propose luminescent mechanism for them when they are exposed to the ionizing radiation. It was recorded both the photoluminescence yield and the emission spectra for all samples. The samples studied were from the family of Ca2Al2SiO7 (CAS) doped with transition metals as rare earths. They were prepared using a hybrid methodology that combines sol-gel and solid state reaction and are potential to be used as scintillators for ultraviolet and soft x-rays [1]. Results carried out in photoluminescence yield indicated a strong absorption around 6 eV in all samples suggesting CAS optical band gap is around this value. The emission spectra indicated typical transitions from dopants and large band emissions from near ultraviolet to visible range from pure samples were also observed suggesting this emission are associated to the exciton. With this set of results it was possible to observe CAS optical band gap is around 6 eV, which is good agreement with previous result [1] and the behavior of emission when the samples are excited immediately above the optical band gap. It was also possible describing the excitation from fundamental levels, that is an important key to know the whole luminescence process when the material is exposed to the ionizing radiation.

TEIXEIRA, V.C., MONTES, P.J.R., VALERIO, M.E.G., "Structural and optical characterizations of Ca2Al2SiO7: Ce3+, Mn2+ nanoparticles produced via a hybrid route", Opt Mater, 36, pp. 1580-1590 (2014)

### XANES study of the oxidation state of Cu, Ni and Ce cations in the Cu-Ni/ce{Ce\_{0.9}Zr\_{0.1}O2} cermet in reducing atmospheres

Presenter: Toscani. M.

Toscani L. M.; Zimicz M. G.; Martins T.; Lamas D. G.; Larrondo S. A.

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Solid Oxide Fuel Cells stand as a promising technology for efficient and sustainable energy production. In this context, \ce{Ce {0.9}Zr {0.1}O2} (Ni/ZDC) cermets are an excellent alternative to Ni/YSZ commercial anodes. The aim of this work is to study the effect of partial substitution of Ni by Cu in that cermet and to evaluate its performance in redox and catalytic conditions. In situ XANES experiments using synchrotron radiation were performed at the Ce-LIII, Cu-K and Ni-K edges. Temperature programmed reduction (TPR) experiments were carried out in 5 mol.\% \ce{H2}/He atmospheres. Ce-LIII edge results showed Ni/ZDC and CuNi/ZDC almost identical profiles up to 670°C, while the bimetallic sample presented higher reduction percentages up to 800°C. Both samples concluded the experiment with a total 40\% of Ce reduction. In the Ni-K edge, the bimetallic sample presented a lower reduction onset temperature and a higher proportion of Ni surface species exhibited in the larger area under first peak of TPR profile, with 100\% Ni reduction in both cases. Cu-K edge results showed a two step Cu reduction for the bimetallic sample. When the cermets are exposed to  $ce{CH4}:ce{O2}=2.6$  ratio atmosphere, they exhibited low catalyst reduction percentages: 8% Ni reduction for both cermets, 4% and 2% Ce reduction for cermets containing Ni and CuNi, respectively. The mass spectrometer signals of the exhaust gases during the experiments showed the production of both \ce{H2} and CO for Acknowledgements: To LNLS (proposal XAFS1-15329), temperatures above 650°C. PIDDEF N°011/11 and ANPCyT (PICT 2013-1587).

### Photofragmentation of a perfluorocarboxylic acid using synchrotron radiation: Study of CF3CF2C(O)OH

Presenter: Védova. O. D.

Védova C. O. D.; Martínez Y. B.; Bava Y. B.; Filho R. C.; Romano R. M.

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Perfluorinated compounds have been extensively used in industrial applications during the last years. Recently, some perfluorinated species were detected in tissues of animals, in environmental waters, and in the atmosphere.[1,2] Different processes may be responsible for the presence of these compounds in the environment. For example, the thermolysis of fluoropolymers and the degradation of fluorotelomer alcohols result in the emission of perfluorocarboxylic species to the atmosphere. In this work, and as part of a general project aimed to the elucidation of the photofragmentation mechanisms of compounds relevant for atmospheric chemistry, we present the study of perfluoropropionic acid using synchrotron radiation with energies between 7.3 and 300eV. The photoexcitation and photofragmentation of this species was studied in the TGM beamline at LNLS. Above 11.7 eV different ions can be observed in the PEPICO spectra. At exactly 11.7 eV, COH+, C2F4+, and M+ were detected. This result shows that the cleavage of two covalent bonds is carried out at energies barely higher than the first ionization potential. The peak corresponding to C(O)OH+ was detected at 11.8 eV, and CF3CF2+ appeared in the spectra from 15 eV. The parent ion was only detected below 16 eV. From that energy, the heaviest ion observed in the PEPICO spectra was the M-OH+ fragment. Unlike the fragments observed in the previously studied analogous molecule CF3CF2CF2C(O)Cl,[3] CF+, CF2+, and CF3+ ions were not the most abundant peaks in the spectra.

<sup>[1]</sup> Hurley, M. D.; Ball, J. C.; Wallington, T. J.; Anderson, M. P. S.; Ellis, D.A.; Martin, J.W.; Mabury, S. A. J. Phys. Chem. A, 2004, 108, 5635-5642. [2] Shine, K.P.; Gohar, L.K.;Hurley, M. D.; Marston, G.; Martin, D.; Simmonds, P. G.; Wallington, T. J.; Watkins, M. Atmos. Environ., 2005, 39, 1759-1763. [3] BerruetaMartínez, Y.;Bava, Y. B.; Erben, M. F.;Cavasso Filho, R. L.; Romano, R. M.; Della Védova, C. O.J. Phys. Chem. A, 2015, 119, 1894–1905. Acknowledgements: This work has been largely supported by LNLS under Proposal TGM-17872. We thank Arnaldo Naves de Brito and TGM beamline staff for their assistance throughout the experiments, and Facultad de Ciencias Exactas, Universidad Nacional de La Plata, CONICET and ANPCyT for financial support.

### Phosphorus speciation during the production of phosphate fertilizers using a metallurgical acid residue

Presenter: Vergütz L.

Vergütz L.; Santos W. O.; Filho L. F. S. S.; Hesterberg D. R. L.; Mattiello E. M.

Departamento de Solos - Universidade Federal de Viçosa; Departamento de Solos - Universidade Federal de Viçosa; Departamento de Solos - Universidade Federal de Viçosa; Department of Soil Science - North Carolina State University; Departamento de Solos - Universidade Federal de Viçosa Universidade Federal de Viçosa leonardusvergutz@gmail.com

Soluble phosphate fertilizers are produced in the fertilizer industry by the acidification of rock phosphates (RP) using pure acids. Therefore, the use of acid residues (AR) to solubilize RP is a potential way of reusing any sort of AR. This way, the objective of this work was to evaluate the use of a metallurgical AR to produce phosphate fertilizer from three contrasting RP commonly found in Brazil. This AR comes from the leaching of pegmatite by a mixture of three acids (H2SO4/HF/HCl) in order to extract Ta and Nb. The work was carried out in a 3x5 factorial design: three RP (Araxá, Patos, and Bayóvar) and five RA concentrations (0,0; 12,5; 25,0; 50,0; and 75,0 % v/v in water). The changes occurring in the original RP and the P solubility of the products following the addition of AR were evaluated by X-ray diffraction (XRD), P K-edge XANES, and water soluble P (Pwater) and neutral ammonium citrate soluble P (PNAC). In terms of P solubility, the reactivity of the RP increased with increasing AR concentration. It means that the concentrations of Pwater and PNAC increased the higher the AR concentration. This increase in RP reactivity was also accompanied by the disappearing of apatite (P-Ca) peaks in the XRD, which is the main P species in the RP and not readily bioavailable. Phosphorus K-edge XANES spectrum of P-Ca species is characterized by the absence of pre-edge structures and the presence of strong post-edge shoulders. As AR concentration increased, these features disappeared in the products, agreeing with the XRD. What XRD could not show was the formation of non-crystalline P-Ca and P-Fe species. As AR concentration increased, an important feature arose, which was the pre-edge structure, characteristic of amorphous P-Fe species. At the end, linear combination fitting showed that as the AR concentration increased, P-Ca species present in the RP changed to more soluble species of P-Ca, with the formation of great amounts of amorphous P-Fe.

### Identification of the substrate binding sites in Actinobacteria sulfotransferase Cpz8

Presenter: Vieira B. D.

Vieira B. D.1; Fernandes A. Z. N.1; Kaysser L.2; Gust B.2; Trivella D. B. B.1\*

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The understanding of natural product biosynthetic pathways has received increased interest of the modern scientific community. Recently, our collaborators at the University of Tubingen (Germany) identified steps in the biosynthesis of the antibiotic caprazamicin (CPZ) in Streptomyces sp.. The CPZ sulfation pathway is catalyzed by a PAPS dependent sulfotransferase, Cpz8. Cpz8 uses PAPS as sulfate donor and a presuficidins as sulfate acceptors. Interestingly, Cpz8 doesn't display a typical sequence for PAPS binding domain. Low similarity is displayed by Cpz8 when compared to known sulfotransferases (19 and 29% similarity for human and S. mansoni respectively). Therefore, we attempted to determine the crystal structure of Cpz8 to elucidate its mechanism of catalysis and substrate binding. For this, the enzyme was heterologously expressed, purified and subjected to crystallization at RoboLab (LNBio-CNPEM). Diffraction data sets were collected at the MX-2 beam line (LNLS-CNPEM) with resolution better than 2 Å. It wasn't possible to determine the crystal structure of Cpz8 by molecular replacement. Therefore, SAD experiments were carried out also at the MX-2, using quick cryo-soaking with Nal, CsCl and GdCl3. After data processing, the phases were recovered using SAD data sets with iodine. The calculated electron density map allowed for the determination of the crystal structure of Cpz8. The atomic model was refined and analyzed. It was found that - despite the poor similarity with known PAPS binding motives in the primary sequence - the Cpz8 3D-structure presents a large cavity compatible with the PAPS binding site. The important chemical groups to interact with PAPS and sulfate transfer are preserved in this putative binding site. A second cavity is also found and predicted to bind the sulfate acceptors. The crystallization of Cpz8 in complex with PAPS and the sulfate acceptor (presuficidin A-D) are in progress.

#### Surface nanomodification of polypropylene

Presenter: Waldman W. R. Waldman W. R.<sup>1</sup>, Galante D.<sup>2</sup>, Fitaroni L. B.<sup>3</sup>, Cruz S. A.<sup>1</sup>

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Surface modification of polymeric materials by exposure to light occurs by absorption of luminous radiation and the subsequent excitation and reaction of specific chemical groups with each other or with the atmosphere. The modification advantage is the production of a highly reactive surface. The aim of this work is to study the polypropylene surface modification as a function of exposure time to extreme UV under vacuum, in order to verify the kinetics and the nature of the modification. Polypropylene was exposed to extreme UV, generated in Toroidal Grating Monochromator beamline (TGM) during different times, from 10 to 60 minutes. The obtained results showed that the modified region presented an evolution as a function of time of the surface modification pattern, as followed by scanning electron microscopy (SEM). The observed modification was a corrugated morphology, with dimensions ranging from 200 to 300 nm without any evidence of cracking or crazing, which is the opposite of the expected in photodegradation under vacuum or in presence of atmosphere.

Daniel E. Weibel, Polymer Surface Functionalization Using Plasma, Ultraviolet and Synchrotron Radiation. Composite Interfaces 17 (2010) 127-136

Masaki Ono and Eizi Morikawa. Ultraviolet Photoelectron Spectroscopy Study of Synchrotron Radiation-Degraded Polyethylene Ultrathin Films. J. Phys. Chem. B (2004), 1894-1897

#### Extreme UV and Selective Inner-Shell Fragmentation Studies of Novel Polymeric Resist Materials

Presenter: Weibel D. E.

Chagas G.1; Satyanarayana V. S. V.2; Kessler F.3; Belmonte G. K.1; Gonsalves K. E. 2; Weibel D. E.1

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Two key concepts are important in extreme ultraviolet lithography (EUVL) to be the candidate for mass production of future integrated circuits: the polymer formulation and the photo-fragmentation process. We present here a detailed study of EUV and carbon inner-shell excitation of homopolymers and copolymers that have in their structures sulfonium and triflate groups as EUV absorption-enhancing units. The goal is to achieve sub-20 nm line patterns having low line edge roughness (LER) of < 2.0 nm and high sensitivity. The present work demonstrates the lithographic performance of several photoresists, MAPDST homopolymer and MAPDST-MMA copolymer, and the correlation with the photodynamic fragmentation processes. The dependence of the photodynamic processes on the excitation energy was studied by scanning the synchrotron energy. Particular excitations were selected which finally led to selective fragmentation of the polymers. The unique properties of the synchrotron radiation, highly monochromatic and intense, allowed to obtain detailed information about the photodegradation process in the new resists. The SGM (spherical grating monochromator) and the Planar Grating Monochromator (PGM) beamlines for VUV and soft X-ray spectroscopy were use in the present study. NEXAFS, XPS and QMS techniques were employed as analytical tools. The polymers were found to be sensitive to EUV irradiation at 103.5 eV showing that the triflate and the ester group were the weakest part of the MAPDST homopolymer with important desorption of  $SO_2^+$ ,  $SO^+$  and  $CF_3^+$  fragments during irradiation. The presence of a metacrilate group (MMA) led to a lower gas desorption rate and higher resistant to irradiation than the homopolymer. When a new MANTMS homopolymer and co-polymer (having a biphenyl group) were irradiated at selected carbon inner-shell excitation energies it was found a highly selective process of bond breaking mainly under the  $\pi^*_{C=C}$  excitation of the phenyl functional groups. The defluorination and loss of sulfonated groups with the increase in the irradiation time for the MANTMS homopolymer was evident when the excitation energy was tuned to excite the  $\pi^*_{C=C}$  transition. On the contrary, C1s  $\rightarrow \pi^*_{C=O}$  and C1s  $\rightarrow \sigma^*_{C-F}$  excitations did not produce important changes in the polymer surface region. This methodical investigation will provide guidance in designing new resist materials with improved efficiency for EUVL through polymer microstructure engineering.

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Singh, V.; Satyanarayana, V. S. V.; Batina, N.; Reyes, I. M.; Sharma, S. K.; Kessler, F.; Scheffer, F. R.; Weibel, D. E.; Ghosh, S.; Gonsalves, K. E., Performance evaluation of nonchemically amplified negative tone photoresists for e-beam and EUV lithography. J. Micro. Nanolith. Mem. 2014, 13 (4), 043002-043002.
Satyanarayana, V. S. V.; Kessler, F.; Singh, V.; Scheffer, F. R.; Weibel, D. E.; Ghosh, S.; Gonsalves, K. E., Radiation-Sensitive Novel Polymeric Resist Materials: Iterative Synthesis and Their EUV Fragmentation Studies. Appl. Mater. Interfaces 2014, 6, 4223–4232.
Seletive Fragmentation of Radiation-Sensitive Novel Polymeric Resist Materials by Inner-Shell Monochromatic Irradiation Gabriela Chagas, Vardhineedi Sri Venkata Satyanarayana, Felipe Kessler, Guilherme Kretzmann Belmonte, Kenneth E Gonsalves and Eduardo Weibel.

### Purification, Crystallization and Preliminary Analysis of the X-Ray Diffraction Data from an Epoxide Hydrolase identified in Streptomyces sp.

Presenter: Wilson C.

Wilson, C.<sup>1</sup>, <sup>2</sup>; Santos, J. C.<sup>2</sup>; González, G. D. T.<sup>3</sup>; Oliveira, L. G.<sup>3</sup>; Dias, M. V. B.<sup>1</sup>, <sup>2</sup>

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Epoxide hydrolases (EHs) are co-factors independent enzymes which catalyse the conversion of epoxides to transdihidrodiols. They exhibits different biological roles such as metabolism of mutagenic and/or carcinogenic epoxides. In microorganisms, EHs are also responsible for hydrolysing specific carbon sources such as tartaric acid, limonene and epichlorohydrin. Epoxide hydrolases from Streptomyces sp. in addition to metabolic functions, exhibit biocatalytic activity proprieties using a limited range of substrates. They have biotechnological interest to discovery new process for enzymatic separation of enantiomeric mixtures of epoxides and region specific hydrolysis. The objective of this work was to perform the purification, crystallization and obtain data of X-ray diffraction for B1EPH2, an EH identified in Streptomyces sp. Initially, B1EPH2 gene was cloned into pET28b (+) and transformed into E. coli BL21(DE3). The enzyme was expressed in 2YT medium with induction by IPTG. Purification was carried out using affinity and size exclusion chromatographic. Crystals grown using the vapour diffusion methods. The diffraction screening and data collection of crystals was performed at the MX2 line of LNLS, Campinas, Brazil. The X-ray diffraction data were processed using the CCP4 program package. B1EPH2 was purified by affinity chromatography using a buffer containing 50 mM Tris-HCl pH 7.0; 100mM NaCl and 10% (v / v) glycerol and a gradient of imidazole from 0-500 mM. The pure protein was concentrated up to 12.5 mg/ml and co-crystallized with 10 mM of valpromide, an inhibitor of epoxide hydrolases, at 18°C. Crystallization occurred in different conditions. The crystals were diffracted up to 2.5 Å and belong to space group P412121. The protein B1EPH2 was crystalized in different conditions and the crystals were diffracted up to 2.5Å. The next step of this work is to perform the resolution of the structure and compare it to other epoxide hydrolases from different organisms.

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### Physical Simulation and Advanced Characterization of Structural Materials

Presenter: Wu L.

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The X-ray Scattering and Thermo-Mechanical Simulation (XTMS) experimental station has been co-developed by the Materials Characterization and Processing group of the Brazilian Nanotechnology National Laboratory (LNNano) and the Brazilian Synchrotron Light Laboratory (LNLS) Engineering and Technical staff. Such installation, which is located at LNLS XRD1 beamline, is operated by LNNano with LNLS support. It consists of a diffraction beamline built around an advanced thermo-mechanical simulator, the Gleeble® Synchrotron system, which allows the material of interest to be submitted to a wide range of thermo-mechanical conditions with high accuracy and reproducibility. Linear or area X-ray detectors are mounted in a high-resolution goniometer for fast data acquisition, which allows time resolved measurements.

# Micelle and mesoporous silica formation with different cethylammonium surfactants

Presenter: Zapelini I. W.

Zapelini I. W., Campos A. F. P., Modesto P. P., Ferreira A. R. O., Alkimim I. P., Araujo J. A., Silva L. L., Cardoso D.

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MCM-41 hexagonal mesoporous silica is of high interest by the scientific community because its easy preparation, pore diameter adjustment (between 1.5 and 10 mm), high superficial area, porosity and short range of pore diameter distribution. These characteristics allow this material to be used as catalysts, supports, adsorbents and membrane in many processes. This material is formed by the combination between polyanionic silica species and the micelles of the cationic surfactant. The way that this combination performs has high influence on the mesoporous phase obtained. Therefore, the objective of this study is evaluate the influence of surfactants with different hydrophilic head sizes on micelles formation and then on the MCM-41 hybrid silica. The studies were performed by small angle x-ray scattering curves of aqueous solutions of cethylammonium surfactants. The results showed a decrease of the micellar organization and of the bromide anion layer thickness around the micelles when the size of surfactant head increases (C16TMABr < C16TEABr < C16TPABr). It is believed that the high size of the surfactant head provides a decrease on the condensation of bromide anions around the micelles [1] and larger steric hindrance between heads, interfering on the mesoporous silica formation. According to the SAXS results, the xray patterns of the mesoporous silicas prepared by C16TEABr and C16TPABr showed the formation of structures with less organization and with only the (100) diffraction peak, compared to the MCM-41 prepared by C16TMABr surfactant which has, additionally, the (110), (200) and (210) peaks.

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### Fragment based drug discovery targeting proteins associated with virulence and resistance to antibiotics in Mycobacterium tuberculosis

Presenter: Zuniga G. A. L.

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The emergence of multidrug resistant Mycobacterium tuberculosis has led to the development of new anti tuberculosis drugs. L, D-transpeptidases (LdtMt) catalyzes 3-3 transpeptide linkages and usually are not inhibited by  $\beta$ -lactams. Its association with an intrinsic *β*-lactamase prevents the use of these antibiotics to treat tuberculosis. LdtMt1 and LdtMt2 are essential for Mt and their disruption results in severe morphological and functional alterations. Another M. tuberculosis protein named Eis, Enhance the survive of M. tuberculosis within of macrophages phagosomes by negatively modulating multiple signal that conduce to bacterial death, and also induces intrinsic resistance to aminoglycosides by acetylatilation and prevent its binding to the bacterial ribosome. Both, L.D transpeptidases and Eis proteins are interesting targets for Drug discovery. This work aim to identify simple molecules that maybe chemically modified for the generation of potent inhibitors against L.D. transpeptidases and Eis protein from Mycobacterium tuberculosis The genes for LdtMt1 LdtMt2 and Eis proteins were cloned from M. tuberculosis genomic DNA (Strain H37Rv) in pET28a, superexpressed in E. coli BL21(DE3), and the products were purified by immobilized-metal affinity (IMAC) and size-exclusion chromatography (SEC). We are performing a screening using thermal shift against a library of about 400 compounds to identify hits which have low affinity to these targets. The proteins have already been crystallized in different condition using hanging drop and the crystals are being used for in soaking experiments with compounds identified by the first trial. We will also further apply a variety of biophysical and crystallographic techniques to validate the candidate molecules. In collaboration with research groups of organic chemistry, molecules with higher affinity to the targets will be synthesised and promising inhibitors might be tested against M. tuberculosis cultures.

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### An investigation of the Morphology exhibited by HDPE composites after being subjected at very high rates of loading

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Due to the low weight allied with high mechanical resistance, high-density polyethylene (HDPE) composites are the current state-of-the-art body armor system. In ballistic systems, HDPE is the matrix of the ballistic panel. Because material properties might be enhanced controlling process conditions and introducing fillers, we aim to be effective in improving the ballistic properties of the ballistic panel matrix by the dispersion and orientation of a second phase. In the present study, the filler, vermiculite particles modified by carbon nanofibers, V900 and VTRITFe800, was synthesized by chemical vapor deposition (CVD) method. The V900 particle were synthesized at 900oC and the VTRITFe800 were impregnated with iron chlorideand synthesized at 800oC. Due to their amphiphilic character, hydrophobicity from the nanofibers and hydrophilicity from Si and Si-Al oxide surface, the particle were used in the production of polymeric composites using HPDE as matrix. In order to study the material behavior and it's mechanical properties, the specimen were subjected to high strain rates, 104 s-1, using a split Hopkinson pressure bar. At this article, using SAXS technique, we investigated the composite nanomorphology before and after deformation. Before deformation, the presence of the non-polar chains on the vermiculite surface favored the dispersion in the polyolefin. HDPE lamellar structure originated a peak at 0.246 nm-1. The peak position of HDPE composites slightly shifts to lower 0.234 nm-1 values, suggesting an increase of the spacing of the periodical lamellar structure and smaller degree of order. The lamellae thickness was controlled by the filler. After deformation, results showed that composites containing particles presented the optimal results for energy absorption. The after deformation morphology varies due the particle composition and concentration. HDPE lamellar peak did not change after deformation, indicatingits lamellar structure is less susceptible.

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