D2AM, status and highlights
of the French anomalous CRG beamline at ESRF.
presented by J.-François Bérar
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This document is the collective work of all those participating in the beamline. They are to be thanked for their written contributions and for their help and comments in preparing this booklet.
Introduction

Built to respond to a broad range of problems in materials science, the D2AM (Diffraction et Diffusion Anomale Multilongueurs d’onde) beamline quickly focussed an important part of its activity on selected methods involving either anomalous scattering, high resolution or coherence. It remains however completely open to all materials science subjects, as can be seen from the contents of the section on scientific results, such as diffuse scattering, soft condensed matter, material engineering, ...

Its design as an anomalous beamline allows the wavelength to be changed easily and has encouraged some researchers to exploit not only the anomalous contrast but also the whole of the absorption fine structure in diffraction using DAFS. After the first experiments on orbital ordering in bulk materials they progressed to 2 dimensional structures such as multilayers, epilayers and then to quantum wires or quantum dots. Encapsulated dots in semiconductors are now being investigated.

At the same time, scientists engaged in SAXS experiments have taken advantage of the coherence of our bending magnet beam and defined the experimental protocol needed to obtain the requisite quality of the beam. This work is the basis of our high resolution SAXS configuration, which extends the range of the instrument to cover that normally investigated by visible light scattering. By this means, anisotropically scattering systems can be investigated by virtue of the 2 dimensional detection.

After a brief outline of the beamline, this report to the 2004 ESRF review panel goes on to describe a selection of experiments that are representative of the variety of results obtained on D2AM in recent years. This is followed by a small number of complete papers. The entire list of publications for the years 2001-2004 is given in an Appendix; other significant papers are referenced in the text.

1 Beamline presentation

A French CRG beamline

The CRG beamlines at the ESRF were founded by the CEA and CNRS in France in order to facilitate access of the French community to synchrotron radiation. The D2AM beamline was dedicated to anomalous scattering and studies of diffuse scattering at small or high angles. This project received strong support from laboratories in the Grenoble area.

The beamline was built to reconcile three experimental instruments, each sharing the beam time: a diffractometer dedicated to protein crystallography, a small angle camera and a materials science goniometer. The optics were designed to satisfy the common requirements of these groups: the project leaders, J.P. Simon (CNRS) and M. Roth (CEA) attached particular concern to the signal-to-noise ratio.

The beamline was among the first ESRF beamlines to open to users, in September 1994. Its scientific activity was reviewed in May 1997. As a result of the increased demand for protein crystallography, a new French CRG beamline was constructed (FIP). After February 1998 D2AM was no longer the priority of the protein crystallography community and it can be now considered as a materials science station with two instruments:

- a small angle camera (responsible J.P. Simon),
- a 7-circle goniometer (responsible J.F. Bérar).

At the present time, about 1/3 of the beam time is dedicated to the small angle scattering camera while the remaining 2/3 involve the goniometer.

The annual budget allocated by the CNRS and the CEA in 2002 amounted to 218 kE and in 2003 to 190kE. This budget does not include staff salaries but takes into account all charges incurred to the ESRF, running costs and investment, including our detector development program.

1.1 Staff and associated laboratories

As the D2AM beamline is a CRG, its organization differs slightly from that of ESRF beamlines, which enjoy the backing of the Experimental Division support groups. The beamline is managed by a small staff of 4 permanent CNRS employees (a scientist engineer having been appointed in 2001 to reinforce it).
In addition, the beamline receives support from Grenoble laboratories for development, support of external users and in-house research activities. About 10 scientists participate in these activities on a part-time basis, or about 2 - 3 full-time equivalent staff members. The support laboratories are:

- Laboratoire de Cristallographie (LDC)
- Laboratoire de Spectrométrie Physique (LSP)
- Laboratoire de Thermodynamique et Physico-chimie Métallurgiques (LTPCM)
- Service de Physique des Matériaux et Microstructures (SP2M).

The following figure shows for each of these four support laboratories those scientists who are most involved and their field of interaction with the beamline.

1.2 User access to the D2AM beamline.

From the point of view of users, access to the D2AM CRG beamline is similar to those of the ESRF. The ”user dedicated time” is allocated by two different review committees : the ESRF committees allocate 1/3 of the time, while the French CRG committees allocate the remaining 2/3. The scientific topics correspond to those of the ESRF review committees: HS, ME, SC (CH, IN). The ratio between requested and allocated shifts (or proposals) is about 2-2.5.

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Beamtime delivered to users by review committees

A few shifts reserved for teaching are distributed over about 5 days each year. Since 2001, one or two shifts each year are used by industrial companies under contract with the ESRF. Commissioning, maintenance and in-house research share the remaining time. In the period of reference, the allocated beamtime has been totally used by experiments. During these years very few experiments were interrupted by beamline failures (motors, cryostat) and this lost time has been replaced in the following months using beamtime reserved for maintenance.

1.3 Optics and instrument overview

For better stability with respect to energy, the D2AM optics is symmetrical, with a double monochromator situated between two mirrors. The beamline is located on the BM02 bending magnet and uses
the 0.85 Tesla source with a critical wavelength of 0.6 Å (20.6 keV). At the entry to the optical hutch, a primary collimator accepts up to 3 mrad of the horizontal divergence. Primary slits located upstream of all the optical components are used to define the effective vertical and horizontal divergence. The first mirror, a platinum-coated silicon single crystal, acts as a low pass filter and enables the beam to be focussed in the vertical plane. These first elements lie in a very clean vacuum ($10^{-9}$ without beam, $10^{-7}$ mbar with beam) that extends up to the last Beryllium window. The beam is then processed by a double crystal monochromator equipped with Si(111), or occasionally at high energies, with Si(311). This monochromator has a bandwidth of $10^{-4}$ and focusses the beam sagittally in the horizontal plane.

A second mirror, identical to the primary one, provides focussing in the vertical plane. Rear slits, located just after the second mirror and also near the instruments, reduce spurious signals from optical aberrations.

This whole symmetrical arrangement acts as fixed exit optics and should yield a 3:1 demagnification of the source. An exhaustive description has been published \(^1\). It delivers photons from 5 to 25 keV to the experimental hutch in a spot of a few hundred microns. Spots of $70 \times 100 \mu m^2$ can be attained for experiments requiring a very small beam size.

The experimental hutch is equipped with a "7-circle" goniometer, which consists of an Euler cradle for the sample, a detector arm that can move in the plane vertical to the polarization and up to 75° in the horizontal plane. All angles are driven by step motors with a resolution better than 0.001°. A two circle analyser can be fixed on the detector arm. Attention has been paid to ancillary equipment: evacuated sample holder cell, displex, furnaces, etc., are available.

The small angle camera \(^2\) is situated at the rear of the same experimental hutch and receives the incoming beam through an evacuated pipe passing inside the goniometer. The camera itself, on a single granite bench, consists of antiscattering slits, sample holders, exchangeable vacuum pipes up to the beam stop and a CCD detector. The sample-detector distance ranges from 0.3 m to 2 m, with a normal $q$ range between $3.10^{-3} \AA^{-1}$ and $1 \AA^{-1}$ between 8 and 15 keV and a flux of $5.10^{10} - 10^{11} ph/s$. Ultra small scattering has reached $4.10^{-4} \AA^{-1}$, thus providing complete overlap with standard visible light scattering. By contrast, experiments at high pressure or high temperatures need hard X-rays (up to $2 \AA^{-1}$ at 20 keV). The flexibility of the camera has been put to advantage in several in-situ experiments in materials science and engineering as well as for Grazing Incidence SAXS.

Both instruments share detectors. These consist of scintillators/photomultipliers and cameras. The most commonly used is a CCD fiber optic coupled ($60 \times 50 mm^2$). Since 1999, gas detectors have no longer been in use.

### 1.4 Future perspectives and plans for upgrading

To define realistic perpectives, it is essential to situate the place of the CRG beamlines in the context of the planned beamlines at ESRF and SOLEIL. Our major specific expertise lies in skillful use of a low noise, anomalous beamline in the range 5 – 25 keV. Instead of moving on to new and different physics, therefore, it seems to be both more pertinent and of more immediate urgency to improve the instruments so as to investigate more exacting samples, for instance, samples prepared by emerging technologies.

The optical parameters of the beamline were defined long ago, on the basis of the techniques available in 1994. Since its design, the ESRF source has increased in brightness. The beamline optics should keep abreast of these improvements in order to ensure the achievable beam quality for the experiment. A rejuvenation plan was submitted to our founding authorities for this purpose.

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most sensitive element of our optics, the sagittal bender, was replaced in 1998, but we are now limited by the most onerous components of the beamline: the mirrors and the monochromators.

- **Mirrors** - Major improvements have been made in mirror technology since 1994. Our mirrors, made from a silicon single crystal coated by 50 nm platinum, were polished to slope errors lower than 5 µrad (rms) and a roughness of 5 Å (rms). Modern mirrors attain 1 µrad and 1 Å. The vertical structure of our beam appears complex, with a strong dependence on the resolution of the observation. It is probable that such structures are associated with intermediate frequencies coming from the slope errors. At present, to attain high resolution on D2AM the vertical aperture must be reduced to minimize this effect. Our mirrors need to be repolished and their bender optimized: this is the most important item in our rejuvenation plan. Simultaneous with this operation, the beryllium windows must be polished in order to remove the speckle pattern that they generate.

- **Monochromators** - The sagittal bender was replaced in 1998 by the "standard" ESRF design. This improved its reliability, eliminating the backlash associated with the friction between the pusher and the crystal holder. This gives us reproducible control of the curvature and permits dynamical focussing in the energy scan, as is required in DAFS experiments. However due to its design, the focal point of a crystal with ribs is far from perfect. At high resolution two focal lengths are revealed: one associated with the ribs, the other with the intervening material. The consequence is to increase the horizontal size of the spot from 100 µm to 250 µm. This is a heavy penalty for experiments that require very low noise, since one component cannot be removed from the focal plane. We therefore intend to improve the bending system of our monochromator to gain simultaneously in flux and purity.

As these considerations on the mirrors and the monochromator do not at present take into account the various "upgrade plans" that appear at the ESRF, some remarks are in order. We recall that the beamline is competitive in the range 5–25 keV. This range, which allows edges of nearly all the elements to be reached, should be maintained. Another major virtue of the D2AM beamline is its long-term optical stability (greater than 24 h even in coherent operation). This property must also be preserved; it is one of the reasons for which we have not at present decided to move to cryogenic cooling of the monochromator.

- **The 7 circle goniometer** still has good precision on its motors since they are all coded. However it has been operating for 10 years: the 2θ movement is becoming hard and should be repaired. The sphere of confusion of the Euler cradle, repaired in 2002, is about 50 µm. However, since the Eulerian geometry of the sample cradle is not optimised for grazing incidence experiments, we decided to add a new circle this year to assist such experiments. Nevertheless, if this kind of experiment is to become more frequent in the long term, we must consider the choice of adding a dedicated goniometer in the experimental hutch. This must be done while keeping in mind that the most important contribution that D2AM can offer will come from measurements of weak or anomalous signals, as in DAFS.

- **The SAXS camera** is a very flexible instrument with various sample holders and in-vacuo vessels. There are, however, limits to its use, but we must focus our choice on the characteristics of this beamline camera and to ensure that the planned improvement will really be of help to the community. Although the signal-to-noise ratio is very good, a very weak background remains even without a sample; although it can be corrected for, it handicaps measurements of weakly scattering samples. Tests are in progress to identify the principal cause: the optics, the beam monitoring or the detector. The other limitations arise from beamstop size. 2D high resolution experiments stand to benefit from an increase of the sample-detector distance to 3 (4) meters.

These considerations on the instruments imply modifying and enlarging the experimental hutch, an operation that seems reasonable as the control room and user room are sufficiently wide and could be shortened.

- **Detectors** - The beamline started in 1994 with NaI:Tl scintillators associated with photo multipliers and gas detectors. An effort was made earlier to improve the detection efficiency with avalanche photo diodes but the effort was quickly directed towards 2D detection. Even though CCD detectors are far from ideal detectors for our application, a first scientific CCD (1242 × 1152) was bought in 1997. This camera became outdated and developed defects. It has since been replaced by a new one (1340 × 1300) in 2002. Owing to progress in CCD technology, its dark current has been strongly reduced, thus allowing longer exposures.
Nevertheless, the detectors currently available on the market are still far from making full use of the beam quality of modern synchrotron sources, mainly owing to their small dynamic range and relatively long read-out time. To overcome these defects we initiated the development of a hybrid pixel detector. This project is detailed in section 2.7.

2 Scientific results

To represent the variety of topics investigated at D2am over the last few years, some results presented below are the summary of an activity, while others are examples that illustrate selected points. These are the strong points of the beamline and come both from external users and from in-house research.

2.1 DAFS, a probe of local environment in complex samples

In the 90s, DAFS appeared as an opportunity for accessing local environments. Attempts were made to develop energy dispersive DAFS while one group, led by H. Renevier (Grenoble) favoured a monochromatic approach well suited to the optics of D2AM [H-5].

Diffraction Anomalous Fine Structure (DAFS) spectroscopy uses resonant elastic x-ray scattering as an atomic-, shell- and site-selective probe that gives information on the electronic structure and the local atomic environment as well as on the long range ordered crystallographic structure. A DAFS experiment consists in measuring the elastic scattering intensity as a continuous function of the incoming x-ray beam energy in regions spanning absorption edges. Like X-ray Absorption Fine Structure (XAFS) spectroscopy, it provides information on the chemical state and the local environment of the resonant atom (also known as the anomalous atom), like X-ray Absorption Fine Structure (XAFS) spectroscopy. But in contrast to XAFS, it is a chemical-selective and site-selective spectroscopy. Like Multiple-wavelength Anomalous Diffraction (MAD), DAFS provides a means of recovering the phase of the structure factor, which is important for solving the long-range average crystallographic structure.

The requirements of the experiment are to measure the intensities as a function of the energy as fast as possible, with a monitor corrected signal-to-noise ratio as high as 1000 or more (comparable to a typical XAFS experiment), and without distortions of the spectra. Since the mechanical precision and stability of all motors is necessarily limited, we developed a feedback control of the sample rocking-angle position (based on a sample-holder that rocks the sample around the \( \omega \) axis) for measuring the maximum intensity. To cope with and take advantage of the high counting rate on the monitor, the sample fluorescence, or the diffraction peak, we developed a detector based on photodiodes operating in photovoltaic mode at room temperature, which provides high linearity and very large dynamic range. To speed up the Energy-Scanned experiment we implemented a quick-DAFS (q-DAFS) procedure that allows counting during motor movements. The whole settings is described in [3-28].

One of the most interesting DAFS studies \(^3\) performed at the beamline was the attempt to investigate charge ordering in magnetite Fe\(_3\)O\(_4\) by measuring the energy and azimuthal dependence (dependence upon the incoming x-ray beam polarization) of the \(d\)-glide plane (002) and (006) forbidden reflections, below and above the Verwey transition (125K), at the iron K-edge [E-9, T-14, 1-12, 1-13, 1-27]. We have observed a) virtual exitation-deexitation transition through dipolar-quadrupolar channels at the tetrahedral iron atom (A site), b) strong resonance at the 1s – 4p energy transition due to the anisotropy of the dipolar scattering factor of the trigonal \((\bar{3}m)\) octahedral iron atoms (B site) and c) for the first time, the extended part above the edge that has the same origin as the main resonance and exhibits oscillatory behaviour as a function of the energy. The analysis of the experimental data shows that the anomalous scattering factor is anisotropic but is identical for all the octahedral iron atoms at room temperature and remains unaltered across the Verwey transition. The conclusion was drawn that octahedral iron atoms in magnetite cannot be described as pure ionic Fe\(^{3+}\) or Fe\(^{2+}\) ions, either spatially or temporally.

**Nanostructured semiconductors**

For the last years DAFS spectroscopy has proven to be a very successful tool for studying the crystallographic structure of thin films, superlattices and interfaces [T-9, 3-15, 3-23, 1-17]. More recently DAFS in grazing incidence (GI-DAFS)

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has been developed and used to study uncovered InAs quantum wires (QWrs) grown on InP(001)\[1-16\]. The QWrs are aligned along the [110] direction with a typical length above 5 μm, a height between 0.6 and 2 nm, a period of 20 nm with an equivalent InAs coverage of about 2.5 monolayers (fig.1). We measured the DAFS spectra at the maximum of the QWrs satellite reflections, near the (420) and (440) InP substrate Bragg peaks at the As K-edge (11867 eV). The measurements were performed in grazing incidence geometry, with an incidence angle kept constant slightly above the critical angle of InP (about 0.2°). Grazing incidence is mandatory to enhance the QWr scattering contribution with respect to the substrate contribution.

Figure 1: AFM tridimensional view of InAs quantum wires on InP buffer

We extracted the Extended DAFS oscillations that appear after the edge (fig.2a) and analysed them using an EXAFS data processing scheme to get local parameters such as distances and atomic populations. Theoretical multiple-scattering EXAFS signals were calculated by the FEFF code simulating the fine structure signal from an As atom inside a cluster containing In atoms, As\(_x\) and P\(_{1-x}\) atoms. The polarisation of the incoming photons was perpendicular to the surface, so that the contribution of the As and P next-nearest-neighbour atoms to EDAFS is due only to the out-of-plane atoms. Refinement of the data was completed by the least-square fit procedure of the FEFFIT program. The relevant results are the P concentration, (1 – x) = 0.4, and the As-P distance, found to be 4.17 Å, close to the P-P distance in bulk InP (4.15 Å). Therefore, we could exclude the hypothesis of a fully relaxed InAsP epilayer. The P atoms contributing to EDAFS belong to the interface region, 0.5-2 monolayers, and the core of the quantum wire is essentially strained InAs. Two types of interface or a combination of both can explain the results: a) an abrupt InAs/InP interface with periodical strain strips generated in the InP buffer layer due to the interface mismatch strain; b) a corrugated InAs/InP interface with the same periodicity as the wire.

![Figure 2: (a) InAs quantum wire Grazing Incidence Extended DAFS oscillations, after background subtraction, with best fit (continuous line), (b) EXAFS of the quantum wires. The curves have been rescaled for clarity.](image)

For comparison we measured a glancing-angle EXAFS spectrum at the As K-edge (fig.2b) at beamline BM8 (GILDA, ESRF). The spectrum shows a clear As oxide shape with a strong low-frequency component that corresponds to a huge peak at 1.2 Å in its Fourier Transform (FT) (fig.3b). The oxide layer causes a significant loss of information, in particular for shells beyond the first, whereas, for a DAFS spectrum, it lowers the overall diffracted intensity and the jump at the edge, but does not perturb the fine structure signal of the interesting atoms.

For use in practical devices, the nanostructures are encapsulated or embedded in a superlattice. They must therefore be homogeneous in size, shape and composition, to provide well defined emission wavelengths. A knowledge of the strain field, chemical gradients, chemical mixing at the interface, is of great importance to understand the growth dynamics as well as the electronic and optical properties of the nanostructures.

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\(\text{--- Refer also to } [2-13], \text{ S. Grenier et al., Eur. Phys. Lett. (2002) in Appendix for details.}\)
We have recently used GI-DAFS to study the strain, size and composition of InAs Quantum Sticks, embedded in InP [4-2]. Compared with previous works, we used anomalous diffraction in grazing incidence at the As K-edge to study a very challenging system: nano-objects having a small size, which are encapsulated and the x-ray scattering contribution of which is mingled with that of bulk InP, whatever the momentum transfer is. We have shown that the structure factor (i.e. the Fourier Transform) of embedded InAs stick-like nanostructures can be directly extracted, allowing the average height and strain of the QSs to be determined, their composition to be defined and the As/P exchange to be checked. GI-DAFS oscillations, in the energy range above the edge, also give direct information on the local composition and on strain accommodation inside the sticks.

In conclusion, the above results clearly show that grazing incidence anomalous diffraction and GI-DAFS are powerful tools for studying nanostructures. It is of a great importance to be able to carry out 2D anomalous diffraction mappings and GI-DAFS measurements together on a dedicated beamline with a high beam stability, such as D2AM. However, an optics update (the beamline is 10 years old), i.e., polishing the mirrors and improving the sagittal focusing of the second monochromator crystal must be carried out if the development of GI-DAFS is to be pursued. The diffractometer must also be adapted for surface scattering.

2.2 Applications of anomalous scattering at wide angles

This topic has been the subject of a general review, "Resonant Diffraction" by J.L. Hodeau (Grenoble) [1-18]. The notoriety of this research team has attracted many experiments on this topic, working in close collaboration on the beamline. The tunability of the beamline has also been used to characterize liquid systems and amorphous materials. Anomalous scattering of these systems seems to be the way to characterize the atomic distribution at the level of accuracy needed to validate models. S. Hosokawa (Marburg) began his experiments on the beamline on Ge-Se liquids [3-19, 3-20, 2-16, 2-17, 1-19]. He observed a modification in the local ordering which has been associated with a transition. He is now investigating glassy ternary alloys with As-Se.

Following some real time investigations of the recrystallisation of refractive oxides, a careful investigation of molten Y$_2$O$_3$ at high temperature was performed by L. Hennet and coll. (Orléans) [E-1, 3-16, 3-17]. To preserve the oxide from contamination, it was levitated in a gas flow while being heated by a laser. The local structure observed is close to those of the high temperature H-phase with 7 Y-O and 11 Y-Y bounds.

Single crystals

In a single crystal sample, V. Favre-Nicolin (Grenoble) uses the anomalous x-ray diffraction technique at the Ta L$_3$ edge to investigate the nature of the tantalum displacement pattern in the modulated phase of the charge-density-wave compound (TaSe$_4$)$_2$I [E-5, 1-11]. In addition to the known acoustic-like modulation, they find the first direct evidence for the condensa-

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5 refer to [3-16], L. Hennet et al., Apl. Phys. Lett (2003) in Appendix for details
tion of optical-like Ta displacements along the metallic chains corresponding to an LLSS pattern of long and short in-chain Ta-Ta distances (Ta-tetramerization modes). The anomalous signal allows the selection of this optical-like signal which is one order of magnitude lower than the acoustic-like one. This result justifies the model in which the interaction of the electronically coupled optical modes with long-wavelength acoustic shear modes leads to the condensation of a modulation of mixed character.

Also, high-resolution diffraction at 33 keV (fig.4) gives insight into the domain structure, proving that each domain corresponds to a single modulation wavevector.

**Multilayers.**

To improve the ferroelectric properties of the BaTiO$_3$ perovskite, artificial superlattices such as (BaTiO$_3$/SrTiO$_3$)$_n$ were studied. Multilayers with artificially induced strain and chemical modulation were prepared using a CVD method and analyzed on the D2AM beamline in the thesis of M. Nemoz (Grenoble). The major feature of the analysis of 00L diffraction profiles is that a unique set of parameters enables one to describe simultaneously several diffraction orders (L from 1 to 8) over 3 to 4 orders of magnitude. The fitting parameters enable us to evaluate the bilayer periodicity, the layer thickness and the thickness fluctuations, the macro-strain and the atomic intermixing at the BaTiO$_3$/SrTiO$_3$ interface as well as the coherence length of the stacking (Fig.5). We observe the presence of large gradients of the lattice parameter (macro-strain) and of the composition (atomic mixing) along $<001>$ across the interface, which dominate the diffraction curves. This significant migration of the cations throughout the bilayer is permitted by the high miscibility of Ba and Sr in all proportions in the Ba$_x$Sr$_{1-x}$TiO$_3$ solid solution [1-4].

**Figure 4:** High-resolution image of satellite diffraction peaks, with harmonics up to 3rd order. Note the splitting of all peaks, different for all families of satellites, revealing the mono-modulation nature of each of the 4 modulated domains. An insert shows typical intensity dependence with the energy near the Ta L$_3$ edge of satellite peaks, which was used to characterize the optical-like modulation along the Ta chains.

**Figure 5:** The HL map around the (303) peak of the same sample shows no orthorhombic distortion: BaTiO$_3$ and SrTiO$_3$ sub-layers exhibit the same a (= b) = 3.955 Å parameter, compared to a = 3.905 Å for the SrTiO$_3$ substrate. This common large in plane width of all reflections could indicate that a mirror twinned domain phenomenon occurs due to the quadratic strain induced by the SrTiO$_3$ substrate. Top left: measured (blue) and calculated (red) diffraction (004) curves of a CVD-grown multilayer composed of 15 bilayers (80 Å STO/ 80 Å BTO) at $\lambda$ = 0.78 Å. Top right: HL map around the (303) peak The bottom left graph shows the variation of the lattice parameter d(001) across one bilayer. The bottom right graph shows the variation of the Ba and Sr concentration per unit cell in one bilayer.

**Powder diffraction.**

As a specialized powder beamline exists at the ESRF, our effort in powder applications focussed on the use of the anomalous difference to solve problems related to cation occupancy: in numerous materials cation substitution strongly modifies the properties, but conventional refinements do not allow the structural modification to be specified. This method was used in the thesis of J. Lorimier (Dijon) to characterize the valence distribution occurring in nanometric magnetite and complex ferrite, Fe$_{3-x}$Ti$_x$O$_4$, by in-situ studies [T-11, 3-22, 3-24, 2-14, 1-20].

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In bicationic zeolite, due to the partial occupancy of the active cation site, only anomalous refinement can be used to locate cations such as Sr\(^{++}\) in one site and Br\(^+\) in the other. Moreover, these system are very sensitive to the reaction conditions and to approximate the catalytic process experiments must be carried in-situ.

A dedicated experimental cell was developed and tested with success during the thesis of H. Pallanchner (Grenoble). The quality of the data measured with this cell (fig.6) permitted the cations in Sr(Rb,Na)X zeolites to be localised on the site revealed on the Fourier map. Working near the absorption edge allows the use of the anomalous contrasts. Depending of the energy some lines increase while others decrease. Using this variation, it now possible to detail the Sr\(^{2+}\) and Rb\(^+\) cation distributions on the insertion sites.

Figure 6: Part of the powder diffraction spectrum by Sr\(_{25}\)Rb\(_{16}\)X. Insert : Anomalous dependence of Sr\(_{25}\)Rb\(_{16}\)X diffraction lines at Sr edge and fourier map.

2.3 Diffuse scattering and weak signals

The good signal-to-noise ratio achieved on the beamline together with its harmonic purity were soon put to use by users. M. Amara (Grenoble) employed this set-up to detect a new quadrupolar ordering in NdMg. Superstructure peaks associated with the 4f orbital ordering were measured below the 35 K antiferromagnetic transition \[3-1, 2-24\]. Several days were needed to do the measurement because of the low counting rate. This success was due to the beamline stability. This set-up was also used by S. Ravy (Orsay) in the study of vanadium-doped blue bronze, which are quasi-one-dimensional conductors. They discovered\(^7\) Friedel oscillations, which were predicted long ago but have remained unobserved : the pinning of the charge density wave by the defect induces an oscillation of the electron density which leads to unusual diffraction profiles for satellites \[E-6, 2-31, 1-26\].

The neutral-ionic transition in mixed-stack charge-transfer organic compounds serve as a model system to study the physics of non-linear excitations. E. Collet (Rennes) showed that the temperature dependence of diffuse scattering is very anisotropic. He established that the evolution of certain correlation lengths agree with a one-dimensional relationship \[E-3, E-10, H-2, 2-6\]. There is a crossover from a 3D to a pronounced 1D regime for the fluctuations. The strong cooperativity inside the stack is reflected in the size of the strings of length about 100 Åat T\(_{1D}\). This provides direct evidence for 1D Lattice-Relaxed Charge-Transfer fluctuations.

The signal-to-noise ratio efficiency of the beamline was also of great advantage for the study of diffuse scattering of quasi crystals by M. de Boissieu (Grenoble) for the thesis of A. Letoublon and S. Francoual.

Quasicrystals are a new paradigm for long range order: their diffraction pattern displays sharp Bragg peaks but with a symmetry that is incompatible with translational symmetry. Their atomic structure is now better understood but the details of the structure remain to be determined. Such details are important for the understanding of physical properties of quasicrystals, but also to get some clues on their formation and stabilising mechanisms, a question which is still open. In particular, some models predict that phason fluctuations play a key role. Phason modes are collective diffusive excitations, characteristic of the long range quasiperiodic order. They lead to diffuse scattering in a way that is similar to thermal diffuse scattering. Experimentally it has been shown that the diffraction pattern of the i-AlPdMn quasicrystal displays a fair amount of diffuse scattering which can be reproduced using the generalised elasticity of quasicrystals and long-wavelength phason fluctuations. Absolute scale measurements of the diffuse scattering carried out on the D2AM beam line allowed the determination of the corresponding 2 phason elastic constants. Similar measurements have been carried out on other quasicrystals, in particular the recently discovered binary CdYb icosahedral phase. A systematic exploration of reciprocal space reveals diffuse scattering around the Bragg reflections, with characteristic anisotropies. The diffuse scattering is also due to phason fluctuations in this quasicrystal, although the phason elastic constants have not been yet determined. A comparison on an absolute scale showed that the amount of diffuse scattering is of the same order of magnitude in both the i-CdYb, the i-AlPdMn and i-AlPdRe phases (fig.8). This may be related to entropy stabilisation of these quasicrystals.

2.4 High resolution and Coherent scattering at BM2

The source in an ESRF bending magnet is of high quality: its size is about $30 \times 70 \, \mu m^2$ ($v \times h$, rms), and its movement is limited to a few micrometers. F. Livet et al., (Grenoble), after a careful reexamination of the the characteristics of the beamline, was able to enhance the quality of the focussing and the stability of the optics of BM2. This improvement makes possible high resolution and even coherent scattering experiments. The corresponding loss of intensity was partly compensated by the use of a Direct Illumination (DI) CCD, which provides pixels of size $20 \mu m$ that can be used as photon counters.
Large scale fluctuations in a critical system

The dynamics of the size increase of antiphase domains below $T_c$ in AuAgZn$_2$ has been observed. The high resolution of the experiment and the use of our DI CCD enabled domain sizes from 0.1 to 0.5 $\mu$m to be observed. The classical $t^{1/2}$ dependence for the domain size was verified.

Fluctuations close to the second-order ordering transition of the AuAgZn$_2$ alloy were carefully studied for a wide range of correlation lengths, from 100Å to 1000Å. Fig.9a shows the domain of q-vectors and of intensities that could be measured with our DI CCD, when the temperature is varied from $T_c+14$ to $T_c+0.08$ K. In the corresponding q-range, the critical slowing-down could be observed from the increase in critical fluctuation times after quenching the sample to a few tenths of K above $T_c$ (fig.9b). This is one of the first direct observations of critical slowing down.

Coherent scattering

Coherent scattering is observed when the resolution approaches the diffraction limit: $\epsilon$ being the rms beam divergence, and $\sigma$ the rms size at the sample: $\epsilon \times \sigma \simeq \lambda$. This is easily observed with a collimated beam. Fig.10a shows diffraction of 1$\mu$m asymmetrical slits, with carefully polished edges. This development [3-21] was introduced to allow SAXS experiments to be extended to very low q-values (to $q = 4 \times 10^{-4}$Å$^{-1}$).

The SAXS setup was modified in order to do X-ray Photon Correlation Scattering (XPCS) in filled polymer systems [E-8] 11 These systems were filled with fumed silica particles or carbon black, and the diffusion process studied was very slow. Although our coherent beam intensity was low, of the order of $10^6$ ph/s, the long time fluctuations could be observed. As a CCD was used, we obtained the characteristic times over a wide q-range: from $10^{-2}$Å$^{-1}$ to $5 \times 10^{-4}$Å$^{-1}$. Results from the lower q-values were compared with dynamic light scattering. The long term stability of the SAXS setup was essential for these results (see next Section for details of this experiment).

We are now able to obtain X-ray beams having a high degree of coherence, and are aiming to take full advantage of the focussing optics so as to concentrate a large coherent X-ray intensity on small crystals. Fig.10b shows the diffraction of a sub-micrometric Au crystal. This diffraction is obtained in the neighbourhood of a 111 Bragg peak, and the oscillating streaks observed correspond to facets of the crystal.

2.5 Soft condensed matter

This heading includes not only polymers but also other disordered materials, for whose investigation the resources of D2AM proved invaluable. The accompanying articles published in Langmuir show an application of SAXS to partially ordered nanostructured materials (Ehrburger-Dolle et al.). László et al. illustrates the extended q range (more than 3 orders of magnitude) covered by D2AM and provides an example of the use of a contrast modifying medium in SAXS. The D2AM camera was also used with an efficiency of 50% at 7.8 KeV 9.

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Distribution of divalent ions around a polyelectrolyte chain

Swollen polyelectrolyte gels, when surrounded by aqueous solutions of a monovalent salt, undergo a volume transition when divalent ions above a certain threshold concentration are added to the solution. The distribution of divalent ions in this case is not known either theoretically or experimentally. The aim of this anomalous small angle X-ray scattering study was to determine whether the divalent ions form a cloud around the polymer chain, or whether they condense on it. The gel consisted of sodium polyacrylate in close to its volume transition, with strontium as the divalent ion. At the five different energies of the incident beam used to vary the contrast in the vicinity of 16 keV, the scattering curves have similar shapes, being separated only by constant multiplying factors (Fig. 11). This result, in conjunction with the SANS results from the same sample, indicates that the divalent ion (strontium) does not form an extended cloud but instead espouses the polymer backbone.

\[ \Gamma = Dq^2[1 + a \times \exp(-D_{SR}q^2t_0)] \]

where \( t_0 \) is the time delay after the mechanical disturbance of mixing the sample into the holder. Both relaxation processes are diffusive (i.e., follow a \( q^2 \) behaviour), with rate coefficients \( D = 5 \times 10^{-14} \text{ cm}^2\text{s}^{-1} \) and \( D_{SR} = 2 \times 10^{-15} \text{ cm}^2\text{s}^{-1} \) respectively. A second sample, in which the silica surface was hydrophobic, was thixotropic. In spite of its solid-like properties on a macroscopic scale, the same double diffusion mechanism was also observed, but with much slower rate constants, \( 2 \times 10^{-15}\text{ cm}^2\text{s}^{-1} \) and \( 9 \times 10^{-16}\text{ cm}^2\text{s}^{-1} \) respectively. The differences between the samples are a consequence of the different structures of the polymer that develop around the hydrophilic or hydrophobic silica surfaces. Measurement by XPCS reduces the characteristic time constants of the relaxation processes from months on a macroscopic scale to minutes, owing to the high values of \( q \) attainable with SAXS.

Dynamics of flocculation in filled polymer melts

In the manufacture of rubbers, e.g. for automobile tyres, finely divided fillers (carbon black, silica) are blended with uncross-linked polymer. Owing to their poor solubility in this medium, the filler particles flocculate during the period before Vulcanisation, an aging process that alters the elastic properties of the final product. This phenomenon was investigated by small angle coherent X-ray scattering on D2AM, in conjunction with small angle dynamic light scattering, by studying the motion of fumed silica aggregates suspended in a polymer melt, poly(dimethyl siloxane) (PDMS). (This topic is touched on briefly in Section 2.4 above.) The sample, in which the uncoated surface of the silica particles was hydrophilic, took the form of a high viscosity liquid (500 Pa s). The measured relaxation rates varied with the transfer wavevector \( q \) more weakly than \( q^2 \). The observed motion was found to be a combination of liquid-like diffusion in addition to a structural relaxation in which the diffusing silica aggregates recombine into larger agglomerates at long times. It obeys an equation of the form

\[ \Gamma = Dq^2[1 + a \times \exp(-D_{SR}q^2t_0)] \]
Figure 12: Relaxation rates $\Gamma$ of fumed silica suspension in PDMS melt. High $q$: XPCS, $t_0 = 1$ h. Low $q$: small angle dynamic light scattering $\times 10$ days, $t_0 = 14$ days. Line D is relaxation curve for undisturbed sample. Lines B and C are calculated from the equation for the given values of $t_0$. 
Filler networks in elastomers

Elastomers are soft materials that can be reinforced by dispersing into them solid particles or particle aggregates such as silica or carbon black. However, the mechanism of reinforcement is still not fully understood. Our work consists in investigating by ultra small-angle X-ray scattering (USAXS) the structure of the aggregate network permeating the matrix in the initial sample and its modification during and after elongation. The goal is to relate the macroscopic mechanical behaviour with the structure of the aggregate network.

For an uncross-linked EPR matrix (fig.13), the scattering pattern is isotropic both for the initial unstrained sample and for small strains. For $\epsilon > 0.20$, a characteristic "butterfly" shape appears (fig.13a). In this case, the SAXS intensity curves measured along the direction parallel ($I_\parallel$) and perpendicular ($I_\perp$) to the stretch axis (fig.14b) are no longer identical, unlike undeformed or weakly elongated samples (fig.14a). The power law domain observed in a log-log plot of $I(q)$ comes from the structure factor of CB fractal aggregates ($D_f \approx 1.8$). The value of the slope is close to -1.8 for dilute dispersions. The slightly lower absolute value (1.5) measured for the initial uncross-linked sample (fig.13a) indicates that CB aggregates form a slightly interpenetrating connected networks. For $\epsilon = 0.53$, the aggregates interpenetrate strongly in the perpendicular direction (compaction).

A similar feature is also observed for concentrations below the percolation threshold, in uncross-linked EPR. Fig.16b) shows that cross-linking of the matrix induces strong interpenetration of the CB aggregates, thus forming large agglomerates. The mechanical behaviour of the two series of composites (Fig.16b) is significantly different. The shape of the curve obtained for the uncross-linked system is characteristic of a hard material in which the stiffness results from a network of slightly interpenetrating CB aggregates. The appearance of the butterfly pattern that characterises the start of the disruption of the aggregate network coincides with the maximum of the stress-strain curve. In the same strain range, cross-linked systems display no maximum. Both SAXS and mechanical measurements on cross-linked EPR thus suggest that CB aggregates may be arranged into large agglomerates separated by a layer of polymer. This hypothesis will be checked by electrical conductivity measurements.

Authors: F. Ehrburger-Dolle, F. Bley, E. Geissler, F. Livet, I. Morfin, C. Rochas (Grenoble)
Related publication: [E-7, 5-11, 8-1, 9]
2.6 In situ materials science and engineering

Important contributions to engineering studies of polymers have come from experiments conducted at D2AM, as described in the book for graduate students by L. David, "Introduction à la physique des polymères" [2-10]. Metallurgical engineering has also made significant contributions using in-situ investigations.

Small-Angle Scattering for the study of precipitation in structural materials: in-situ measurements and microstructure mapping.

Structural materials, particularly those used in the transport industry, are constantly confronted with the challenge of improving their specific mechanical properties in order to achieve weight reduction. In the aerospace and automotive industry, aluminium-based alloys are widely used, and frequently gain their mechanical properties through a fine dispersion of precipitates of a second phase. The mechanical strength of these materials depends critically on the details of this dispersion, in terms of size (usually a few nm) and volume fraction (1-5 vol%), which in turn depend on the details of the temperature history that the sample has been subjected to. Small Angle Scattering proves to be the only tool that can access systematically these two parameters of the microstructure, both with good statistics and high precision.

Specific to synchrotron radiation source in terms of SAXS measurements is the reduced time of measurement (as low as 1s), which enables high quality in-situ measurements to be performed, as a function either of temperature or of deformation. This reduced time for measurement, associated with a small beam size (<200 µm) also gives access to mapping of precipitate size and fraction in materials that show a microstructure gradient. This is the case notably in welding, during which the heat generated by the process affects the precipitates located in the so-called heat-affected zone. Such mapping, related to local temperature cycles, are crucial steps in optimising the welding process and in designing new alloys specifically for their welding capacity.

Figure 1 shows the dissolution behaviour of an Al-Zn-Mg alloy for automotive applications (AA7108.50). This material initially contains precipitates of radius 40Å and 2.6% in volume fraction. When subjected to a temperature rise, many of these precipitates become unstable and dissolve. However, upon dissolution (i.e., increasing solid solution), the stability of the surviving particles is increased. After a well defined minimum in volume fraction, the surviving particles start to coarsen, and the volume fraction reaches its equilibrium value at the heat treatment temperature. This behaviour can be well described by a precipitation model (dashed lines), which is a powerful tool for describing the precipitate microstructures after complicated temperature cycles such as are met in welds. Fig. 17d shows such temperature cycles in the case of MIG (Metal Inert Gas) welding, as a function of the distance from the weld line. Fig.17e finally shows the comparison of a precipitate microstructural profile performed by SAXS and the model, calibrated on the reversion experiments, and applied to these temperature cycles.

Figure 17: (a) initial state (b) evolution of volume fraction as a function of time and temperature (symbols : experiments, lines : model) (c) id., precipitate size (d) temperature cycles in a weld experienced in various points of the HAZ (e) comparison between the precipitate microstructure mapped by SAXS across the HAZ and that predicted by the precipitation model.
Structure and mechanical properties of Nylon fibers reinforced with nanofillers.

It is well known that inorganic nanofillers provide efficient reinforcing effects in bulk thermoplastic polymers, mainly due to their high specific surface area, which ensures strong polymer-filler interactions. Multifunctional globular polymeric particles are also nanofillers with a three-dimensional structure and numerous terminal functions. Both kinds of nanofillers are added to nylon PA6 to improve the mechanical properties of spun-drawn fibers. Structural characterisation of unfilled fibers PA6, PA6 filled with 5% globular molecules (GM) and PA6 filled with 1% of surface treated montmorillonite particles was obtained by SAXS and WAXS as a function of draw ratio.

The WAXS patterns of three kinds of spun fibers (Figure 19) display differences in crystalline structure and orientation. The majority phase of the unfilled PA6 fiber is $\gamma$ (a). The PA6-clay fiber (c), also in majority in $\gamma$ form, displays stronger crystal orientation parallel to the draw axis. The PA6-GM fiber (b) exhibits the presence of the crystalline $\alpha$ form in addition to the $\gamma$ form:

Orientation is stronger for the filled PA6 fibers than for the neat PA6 fibers. The filler-matrix interactions in the melt therefore change both the rheological properties of the molten material and the crystallisation kinetics. For PA6-GM, these interactions are probably responsible for the occurrence of a significant amount of $\alpha$ phase, with an extended-chain conformation. The appearance of the sole $\gamma$ phase in PA6-clay fiber suggests that the inorganic filler has a $\gamma$ nucleating effect, as reported for bulk crystallised PA6. The presence of fillers also affects the fibrillar structure as shown in the following SAXS patterns (Figure 20), which display strong anisotropy with lobes:

From the projection along the fiber axis, the long period, $L_p$, lamellar crystal thickness, $L_c$, and amorphous layer thickness, were estimated. A transverse analysis of the SAXS lobes gives access to the fibril radius.
The intensity of the meridian lobes characteristic of the periodic amorphous-crystal arrangement of the fibrillar structure is strongly increased in the case of the as-spun filled fibers. The fibril radius increases with increasing filler content. The strong equatorial scattering of the clay-PA6 fiber (c) is due to the alignment of the clay platelets parallel to the fiber (draw) axis. Stretching the fibrils during the initial stage of drawing first involves an increase of the long period at secondly a decrease of the fibril diameter. Then, the slippage of the fibrils past each other involves peeling off the fibrils by chain unfolding from the lateral surfaces.

The SAXS pictures obtained for PA6 during elongation at points A, B and C were very different: As strain is increased (from A to C), the meridional lobes move toward the beamstop, indicating that the long period increases.

The SAXS pictures obtained for PA6 during elongation at points A, B and C were very different:

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2.7 Detector development

Currently available 2D detectors do not make full use of the high flux and high brilliance of third generation synchrotron sources. For this reason numerous experiments are still performed using slits and photomultipliers that allow only point detection. At the present time, the detectors in most common use are CCD cameras with indirect photon detection.

Modern CCD detectors exceed $10^6$ pixels and their reading-time is close to a second. They are suited for structure collection as integrating devices. The dark current and the converter noise amount to a few counts on a maximum scale of 16 bits. Their complex optical system degrades the properties of the incoming photons: low counts no longer obey a Poisson distribution and the spatial transfer function produces tails that preclude measurements of weak signals near a peak that may spill over its neighbouring pixels. Moreover, they require a cooling system to reduce the dark current as well as the use of a mechanical shutter.

Active pixel detectors have been developed for high energy physics, they present a new opportunity to improve the quality of our measurements.

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The XPAD project

The XPAD project is headed by CPPM-IN2P3 (Marseille), LdC (Grenoble) and the CRG-D2AM beamline.

The XPAD photon counting detector, suitable for materials science and small angle scattering experiments similar to those performed on D2AM, should have the following characteristics: wide dynamic range (> $10^9$ photons/pixel), saturation rate up to $10^7$ ph/s/pixel and a noise lower than 0.1 ph/s/pixel. In our detector, incoming photons are converted to an electron cloud in the sensor in which a polarization ensures migration of the charge through the bump to the electronic chip. In each pixel, the electron bunches are then treated by dedicated electronics. By virtue of this wide parallelism, very short reading times can be achieved (less than 1ms). On-board memories allow frames for real time kinetics experiments to be stored \[E^{-4}, 3^{-5}, 2^{-4}, 2^{-5}, 2^{-8}\] .

The prototype

The XPAD prototype, with pixel size $330 \times 330 \mu m^2$, was manufactured by AMS with $0.8 \mu m$ CMOS technology. The most sensitive part of the design was the analog block, owing to the conflicting requirements of high counting rate and high sensitivity. At 20keV, a photon produces 5500 electrons but only 1400 at 5 keV. These charges must be discriminated from the noise. In each pixel, the discriminator has to be adjusted independently because of the fluctuations in component values generated by the different processing steps.

Each pixel response can be measured with a charge injector to allow pre-configuration of the detector. At the discriminator exit, a logical gate enables counting for the desired acquisition time and, if necessary, can switch off a damaged pixel. This block is followed by a 16 bit counter located in each pixel. A common logical unit located at the bottom of the chip ensures the interface with the outside world: dialogue, timing, ... The chips are then mounted on an epoxy card together with a few elements and an ALTERA programmable chip to handle dialogue and control the memory for the external counters. All this is then monitored by a PC for the test acquisitions.

Results

Electronic tests in the lab show that the pulse width at the amplifier output was about 0.1µs, which is close to the expected value needed to reach the design counting rate. The configuration of the chips was carried out using a radioactive source and an X-ray tube. These preliminary tests exhibited a broad dispersion in the characteristics of the threshold level of the pixels. It revealed the difficulties for finding a configuration in which more than 90% can be simultaneously tuned.

Tests were then performed with the detector mounted on the D2AM SAXS station so as to define a very small and clean incident beam. The energy response of detector was tested between 10 and 24 keV. The energy resolution is estimated to be about 1 keV. A charge spread of $60 \mu m$ was measured by scanning the detector with a very narrow beam $10 \times 50 \mu m^2$. It can be reduced by setting the threshold level at exactly half of the incoming energy: in this case, whatever is the division, the photon is detected by one only of the pixels. The dynamic range was also checked: the xpad pixel detector measures counts ranging from $0.1 \text{ ph/s/pixel}$ to $2.5 \times 10^6 \text{ ph/s/pixel}$. At high count rates when the pixel saturated, no influence could be detected on adjacent pixels.

Figure 23: 3 modules of 8 chips connected to the acquisition card.

Figure 24: SAXS on Silver Behenate: up to 7 orders of diffraction have been recorded as well as intensities close to the direct beam.
In conclusion, pixel detectors can attain the performances needed for high dynamic studies. This is illustrated by the SAXS pattern of silver behenate (a standard for SAXS experiments, d=58.38 Å) and CdYb icosahedral quasicrystal Bragg peaks and diffuse scattering shown below. These images were measured at 20 keV, on a continuous diode bumped to 2 rows of 5 chips. White pixels are those for which the tuning is out of range. The number of truly dead pixel is fewer than 2 %.

Future developments

Our goal is to develop a real detector with more than 10⁶ pixels. Such a detector will consist of several modules to be tiled together. This step has recently been validated by assembling 8 modules of 8 chips each.

To achieve the number of pixels required in numerous experiments, we have already begun the design of a new chip XPAD3. It will use radiation-hard submicronic technology (0.25 microns), which will allow the pixel size to be reduced to 100 – 150μm with similar or enhanced performance.
Grazing-incidence diffraction anomalous fine structure of InAs/InP(001) self-assembled quantum wires

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PACS. 61.10.-x – X-ray diffraction and scattering.
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PACS. 68.65.-k – Low-dimensional, mesoscopic, and nanoscale systems: structure and non-electronic properties.

Abstract. – We have studied nanostructured samples of InAs/InP(001) by means of Grazing Incidence Diffraction Anomalous Fine Structure. The samples, grown by molecular beam epitaxy, show a periodic corrugation on the surface giving rise to an array of self-assembled quantum wires after deposition of 2.5 monolayers of InAs. We measured the (440) and (420) GIDAFS spectra, at the As K-edge, at incidence and outgoing angles close to the substrate’s critical angle. We analysed the anomalous diffraction line shapes vs. the energy, as well as the oscillatory part of the signal in the extended region above the edge and obtained, for the first time, information about composition and strain inside the quantum wires and close to the interface. Our results suggest possible interfaces.

Melting behavior of levitated Y₂O₃

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The yttrium environment in liquid Y₂O₃ at 2770 K has been measured with anomalous x-ray scattering, aerodynamic levitation, and laser heating. The Y–O coordination of 6–7 and the Y–Y coordination of around 12 imply that the close packing of the high-temperature (H-type) solid phase is preserved on melting, in contrast to the large structural changes exhibited by Al₂O₃. The unusually sharp main diffraction peak implies a high degree of chemical order and mirrors the diffraction pattern of the H-type phase. © 2003 American Institute of Physics.

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Phason elastic constants of the icosahedral Al–Pd–Mn phase derived from diffuse scattering measurements

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Abstract

The diffuse scattering in the diffraction pattern of the icosahedral Al–Pd–Mn quasicrystalline phase has been measured on an absolute scale by X-ray and neutron scattering on single-grain samples. Most of the diffuse scattering can be interpreted in the framework of the elasticity theory of icosahedral quasicrystals considering only phason fluctuations. At room temperature the absolute values of the $K_1/k_BT$ and $K_2/k_BT$ phason elastic constants are of the order of 0.06 and 0.031 atom$^{-1}$. The amount of diffuse scattering intensity is insensitive to the sample annealing treatment.
Small-Angle X-ray Scattering and Electron Microscopy Investigation of Silica and Carbon Replicas with Ordered Porosity

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Ordered nanoporous carbons can be prepared by a replica technique starting from an organized silica template. The silica template used, SBA-15, displays hexagonal arrangements of mesopores interconnected by micropores. Two routes are possible for introducing carbon into the pores of the silica: liquid impregnation by a solution of sucrose followed by carbonization or chemical vapor infiltration (CVI). After dissolution of the silica template by hydrofluoric acid treatment, a carbon material is obtained. Small-angle X-ray scattering (SAXS) measurements were performed over a broad range of wave vectors in order to investigate the multiscale structure of the carbon replica as a function of the method of infiltration (liquid or gas route) and the amount of infiltrated carbon. Because of the close match in electron density between silica and carbon, it is possible to investigate the silica mesopore filling. It appears that at least 50% of the pore volume must be filled in order to obtain an organized carbon replica after dissolution of the silica template. It is also shown that the gas route (CVI) prevents the spatially proportional (i.e., affine) shrinkage observed for replicas prepared by liquid impregnation. TEM confirms that the organized carbon is a nearly perfect negative replica of the silica porous structure.
Morphological Investigation of Chemically Treated Poly(ethylene terephthalate)-Based Activated Carbons

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Complementary techniques, including low-temperature nitrogen adsorption and small-angle X-ray scattering (SAXS), are applied to detect the effects of surface functionalization on the morphology of activated carbon derived from polyethylene terephthalate (PET). Scanning electron microscopy (SEM) is also employed as an auxiliary method to visualize the surface below the micron scale. The SEM images reveal a micron-sized ridgelike texture. Room temperature acid treatment makes the ridges become more pronounced, while treatment with boiling acid uncovers fiberlike structures of roughly 1 µm diameter. All samples display an apparent surface fractal dimension of Ds = 2.4 in the wave vector range 0.001–0.02 Å⁻¹. Nitric acid at room temperature increases the surface oxygen content by 2 at. %, while all the adsorption properties and structural parameters reported in this paper are virtually unaffected. Significant differences in the morphology at submicron scales appear only after boiling acid treatment. The resulting carbon remains highly microporous, but the loss of Brunauer–Emmett–Teller (BET) surface area from about 1150 to 304 m²/g is approximately 75%. In addition to the principal peak at about 8 Å, fresh peaks appear in the polydispersive Horváth–Kawazumi (HK) pore-size distribution owing to the burnoff of intervening walls. The average width of the slitlike pores calculated from the Dubinin–Radushkevich (DR) plot increases from 8.4 to 11 Å. The minimum slit width where the applied probe molecules, that is, nitrogen and hexane, can enter increases from about 5 to about 5.4 Å. The separation distance between the high structural units is practically unchanged. When, however, this carbon is in contact with hexane, this distance expands from about 19 to 27 Å. The swelling is consistent with the deformable nature of this sample also illustrated by the low-pressure hysteresis and the reduced helium density. Particular attention was paid to the surface areas derived from low-temperature nitrogen adsorption and X-ray measurements. Owing to the wide spatial range of the structures in these samples, estimates of the specific surface area of activated carbons can be substantially in error unless both upper and lower q ranges of the SAXS spectra are taken into account. Surface areas derived from the adsorption data either by the BET or the DR approaches were always below the values obtained by standard SAXS. As an example, the carbon sample functionalized at room temperature gave surface area values of 1134, 1393, and 1970 m²/g, respectively. The possibility that this difference is caused by inaccessible pores was excluded by contrast variation measurements with hexane.

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