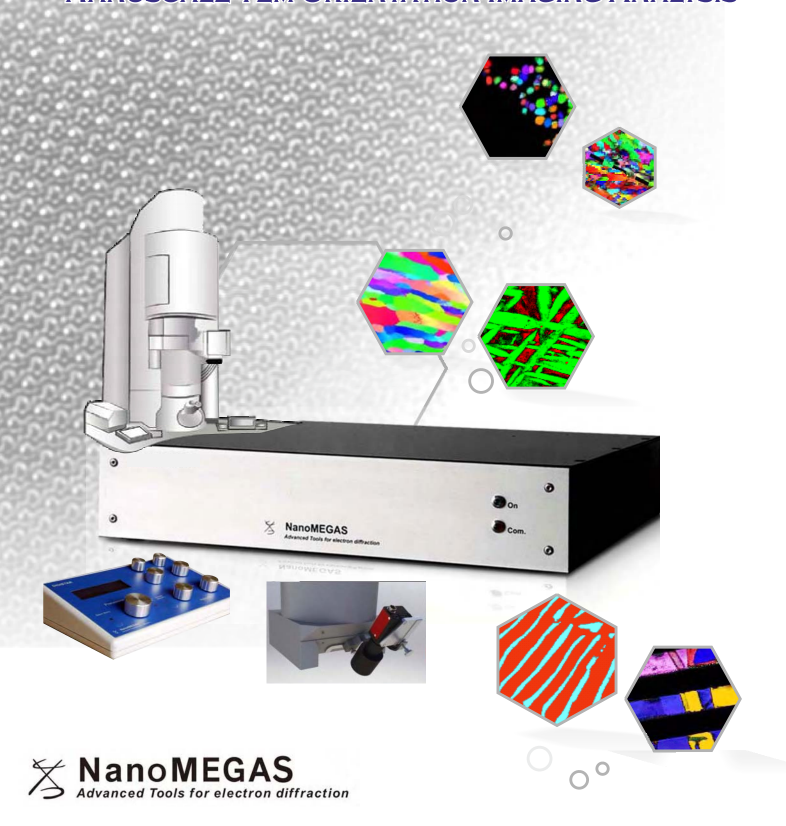
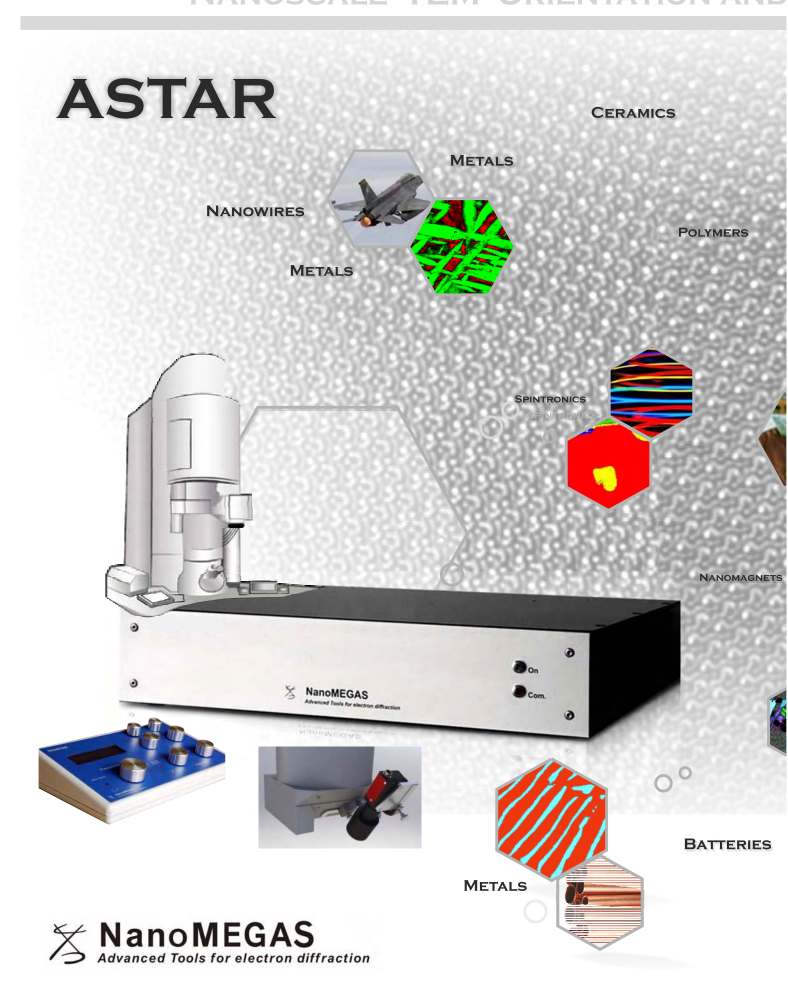
## **ASTAR**

# APPLICATION NOTES NANOSCALE TEM ORIENTATION IMAGING ANALYSIS

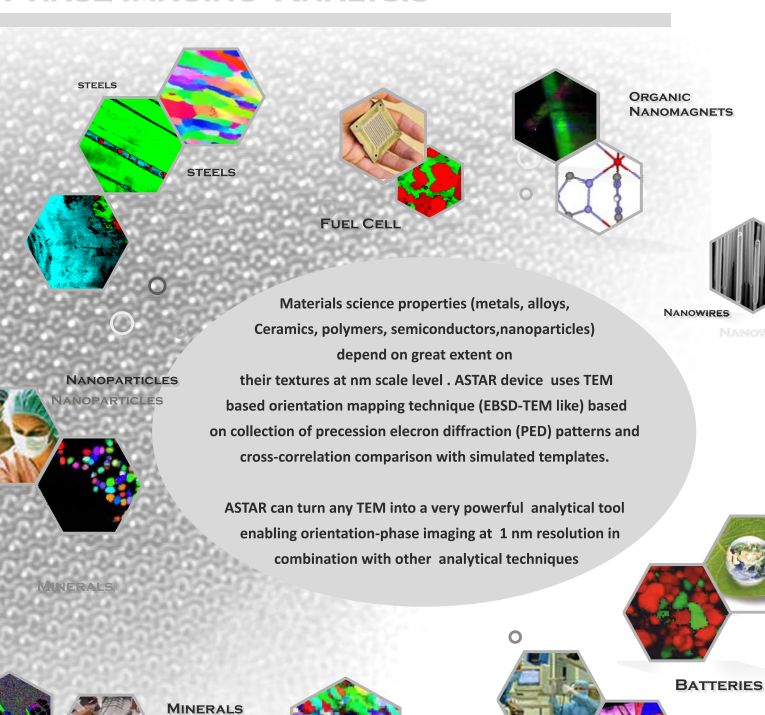


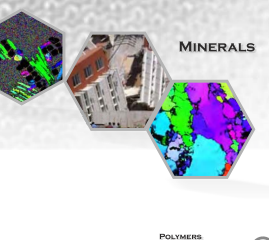
### NANOSCALE TEM ORIENTATION AND

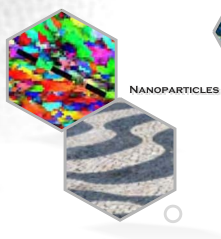


# ASTAR

### PHASE IMAGING ANALYSIS

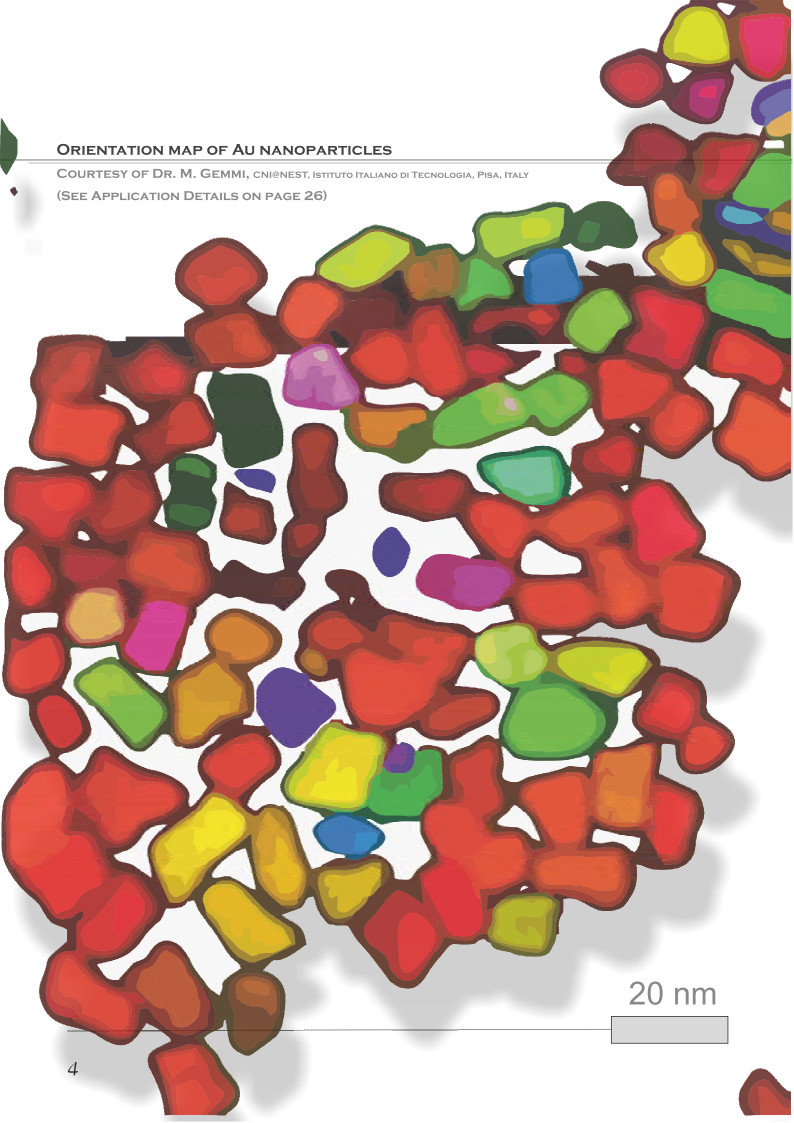








**METALS** 





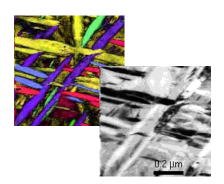
### **CHALLENGE**

### REVEAL TEXTURE AT NANOMETER SCALE



### METALS / ALLOYS

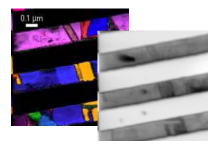
Texture of metals is linked to specific physical properties, so the need to characterize it at nanometer scale





### **SEMICONDUCTORS**

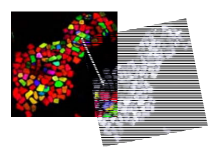
Faster chip performances in electronic devices push Copper Interconnects in close to nanometer scale





### **NANOPARTICLES**

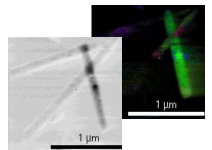
Nanoparticle size and texture are very important for drug delivery of poor water soluble drugs





### POLYMERS / ORGANICS

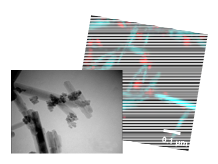
Crysralline polymers and other organics (pharmaceuticals) need new techniques for structure characterization





### **MINERALS**

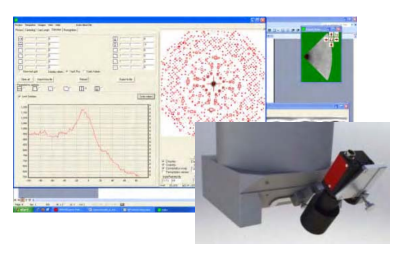
Textures of minerals at nm scale are intimately linked to their physical & chemical properties

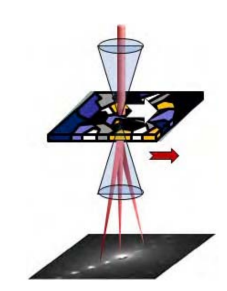


### **SOLUTION**

### **AUTOMATIC TEM ORIENTATION/PHASE MAPPING PRE**

### HOW IT WORKS





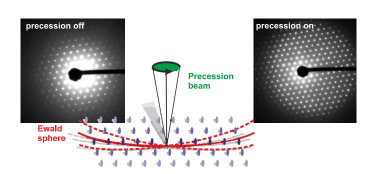
ASTAR is an automatic crystallographic indexing and orientation/phase mapping tool, developed for any TEM.

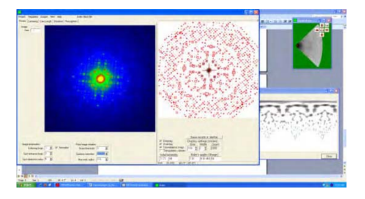
The electron beam is scanned in combination with beam precession through the sample area of interest (a); beam scanning is done by the NanoMEGAS DigiSTAR precession unit without using an inbuilt STEM mode.

A number of electron diffraction (ED) spot patterns from several sample locations (b) are acquired at high speed using a dedicated fast CCD camera (c) placed in front of the TEM screen; local crystal orientation(s) are obtained by comparing all individually obtained ED spot patterns via cross-correlation matching techniques with pre-calculated ED templates.

Detection and orientation/phase mapping of different (known) crystallographic phases and orientations in a crystal structure requires collection

of high quality ED patterns. Electron beam Precession diffraction is extremely useful for obtaining patterns with a large number of spots (d) almost twice as many compared to conventional (e)



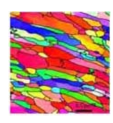


# ASTAR

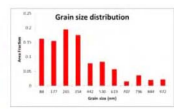
### CESSION UNIT

selected area electron diffraction (SAED) and without dynamical effects (e.g. Kikuchi lines). Therefore, phase and orientation identification based on pattern matching between experimental precession patterns with simulated templates is very reliable and precise. Required diffraction templates are generated every 1º through the

### HOW IT WORKS







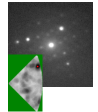
respective symmetry invariant section (f) of orientation space (stereographic triangle for cubic crystals). Resulting colour maps show with nm detail grain structure including boundaries (g), grain size distribution, and relative pole figures.

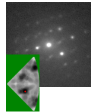
ASTAR resolution on a TEM is being determined by the electron probe size and can reach 1 nm on orientation maps with TEM-FEG microscopes.

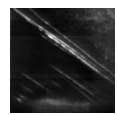
### VIRTUAL STEM (BF/DF)

**ASTAR** orientation-phase maps leads to **digital files** where information (e.g. about the corresponding ED pattern) can be retrieved from every pixel. It is possible then to create virtual bright / dark field limages by selecting specific reflections from the ED pattern. In the example below (**TWIP deformed steel**), a number of twins are present that can be revealed either using a **virtual aperture centered on twin reflections** (b-c)

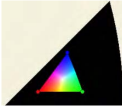
(a) ED pattern corresponding to matrix reflections, (b) ED pattern with twin reflections and virtual aperture centered on a particular reflection, (c) corresponding virtual dark field showing twins, (d) orientation map, (e) pole figure with color code corresponding to twin orientations within the stereographic triangle (f), Courtesy Prof. S. Godet, Univ. Brussels (ULB), Belgium.

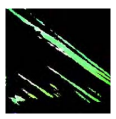




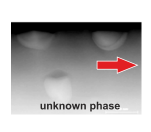


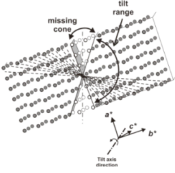




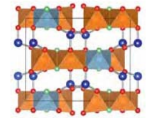


### SOLVING UNKNOWN STRUCTURES: 3D PRECESSION DIFFRACTION TOMOGRAPHY







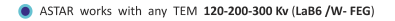


Courtesy Ute Kolb group, Mainz

TEM allows to study nm size crystals and by tilting (manually / automatically) around an arbitrary axis a single nanocrystal (tilt usually >  $120^{\circ}$  e.g 120 ED parterns with step  $1^{\circ}$ ) in combination with precession diffraction (usually  $1^{\circ}$ ), the reciprocal cell of any unknown phase can be recontructed and crystal cell can be evaluated automatically (error 2-5%). ED intensities can also be measured automatically to enable complete solution of crystal structure (optional software).

### **SPECIFICATIONS**

### TEM AUTOMATED ORIENTATION / PHASE IMAGING ANALYSIS

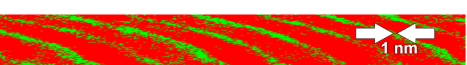


ASTAR orientation-phase map 1 nm resolution with TEM-FEG \*

 ASTAR may work with any type of diffracting material (inorganic / organic) using standard TEM specimen preparation techniques

> ASTAR can work in combination with beam precession (Patent Pending technique) for ultra-precise orientation /phase maps and ab-initio solution of unknown phases

- ASTAR can work / retrofit between multiple TEMs in same lab
- Galvanic Isolation system (GIS)
   via optical fiber
   for ASTAR-TEM connection



### **ASTAR INCLUDES**

- Digital Scan Generator for area beam scanning no need for STEM unit
- Scanning step from 0.1 nm to 100 nm in TEM / nanobeam /microprobe mode (TEM dependent)
- ASTAR dedicated highly flexible ultra-fast (> 100 frames/s; 256 gray levels) CCD Stingray camera to capture ED patterns to produce high speed orientation maps / diffraction pattern calibrations
- CCD Stingray can be placed in front of TEM fluorescent screen via special mechanical adaptor

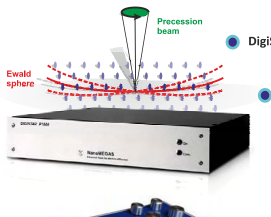


- ASTAR/ CCD Stingray camera can be adapted / retrofitted to any TEM in less than 5 minutes
- ASTAR CCD Stingray can be perfectly stay together with any other on axis / 35 mm port CCD cameras / GIF system









DigiSTAR digital precession unit compatible with any TEM

- Beam precession 0-4° & frequency 0.1-2 kHz (values TEM configuration dependent).

  Precession angle 0.5-1.0° recommended for ASTAR orientation imaging
- DigiSTAR –TEM alignment values are memory stored
- Manual user interface for precession angle visualization and precise alignment adjustments



### **ASTAR (SOFTWARE)**

- TEMDPA: Transmission Electron Microscope (TEM) Diffraction Pattern Acquisition Computer system workstation generates software that controls signal over different channels connected to TEM deflecting coils. Stingray CCD camera can be used for data (ED patterns) acquisition.
- DIFFGEN: Diffraction pattern generator

ED simulated templates are generated every 1° (or less) through the Ewald for every phase / crystal symmetry

- \* User friendly graphical interface (diffraction pattern, pole figure)
- \* Diffraction patterns generated for all crystal systems.
- INDEX : Orientation identification through Diffraction Pattern Matching

Every calculated-simulated pattern generated with DIFFGEN is compared to the experimental measured one through template matching

- \*Optimizing routines for orientation resolution <1°; automatic camera length calibration
- \*Correlation Index map calculation and display (for reliability checking)
- \*Pseudo-bright field or pseudo dark field image reconstruction
- MAP VIEWER: Orientation map viewing software

Crystal orientations extracted for every pattern related to the scanned area are stored in result files.

- Orientation map: pixel color is related to the sample x, y or z crystallographic direction.
- \* Phase map: pixel color is related to different existing crystal phases (eg cubic, hex, tetragonal) in crystals Crystallographic features: grain boundaries may be apparent on orientation maps, grain size and pole figures analysis.
- Export facilities compatible with the most available image / EBSD-SEM analysis software (TSL, HKL etc)
- \* 1 nm resolution tested on reference Al-9nm / 1 nm TiN nanolayer composite sample (see application note) and 0.4 nm thick twin structure detected at < 5 nm Pd nanoparticles

### APPLICATION NOTES

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## DESIGN MATERIALS WITH SPECIFIC APPLICATIONS

Texture of metals is linked to specific physical properties, so the need to characterize it at nanometer scale

Titanium alloys are metals which contain a mixture of titanium and other chemical elements. Such alloys have very high tensile strength and durability (even at extreme temperatures). They are light weight, have extraordinary corrosion resistance and the ability to withstand extreme temperatures. However, the high cost of both raw materials and processing have limited so far their use in military applications, aircraft, spacecraft, medical devices, connecting rods on expensive sports cars and some premium sports equipment and consumer electronics. Auto manufacturers

The challenge:

Solution:

Identify orientation of a variants with respect to the mother phase

ASTAR technique coupled with precession electron diffraction

Porsche and Ferrari use titanium alloys in engine components due to its durable

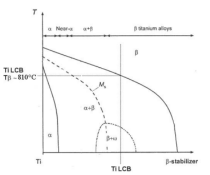
properties in these high stress engine environments.

Although "commercially pure" titanium has acceptable mechanical properties and has been used for orthopedic and dental implants, for most applications titanium is alloyed with small amounts of aluminum and vanadium, typically 6% and 4% respectively, by weight. This mixture has a solid solubility which varies dramatically with temperature, allowing it to undergo precipitation strengthening. This heat treatment process is carried out after the alloy has been worked into its final shape but before it is put to use, allowing much easier fabrication of a high-strength product.

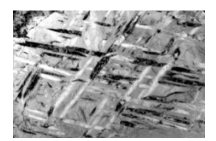
Nowadays, new titanium alloys are being designed, using cheaper elements as alloying elements. For example, Vanadium is replaced by Molybdenum or Niobium. This new alloy family is called Low Cost Beta (LCB) because of the stabilization at

fight aircraft

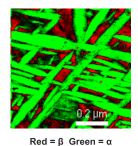


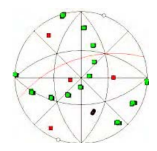


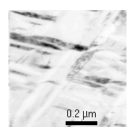
high temperature of the b bcc phase, giving rise to many variants of a hcp phase. These new compositions allow titanium alloys to be developed towards new applications such as in architecture (see the Guggenheim museum in Bilbao) where the exceptional properties of the alloy give rise to new shapes that defy gravity!

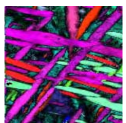


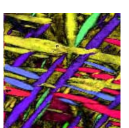
Ti α-phase:
Hexagonal P6<sub>3</sub>/mmc
a=2.97 Å, c=4.72 Å
Ti β-phase:
Cubic Im3m
a=3.28 Å

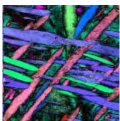












Experimental Data TEM type: Jeol 2100F Map resolution: 5 nm

(111) Pole figure

Spot size: 1.6nm Scanned area: 750x750nm

figure 1

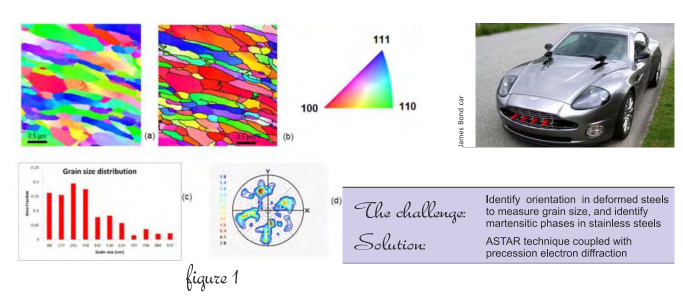
(A) Ti alloy bright field TEM image (B) ASTAR phase map (C) ASTAR orientation map at nm scale z direction, (d) x direction, (e) y direction

Orientation mapping in TRIP steels reveal their texture and help to understand their properties

### BETTER STEELS WITH IMPROVED **PROPERTIES**

Steel is an alloy made by combining principally iron and carbon, moreover other alloying elements are used to improve mechanical characteristics, such as manganese, or vanadium. The resulting steels are commonly know as High Strength Low alloy Steels (HSLA).

Carbon and other elements act as a hardening agent, preventing dislocations in the iron atom crystal lattice from sliding past one another. Varying the amount of alloying elements and the form of their presence in the steel (solute elements, precipitated phase) controls qualities such as the hardness, ductility, and tensile strength of the resulting steel. To harden a material, and especially a ferritic steel, it is also important to control the grain size, through cold rolling and recrystallisation processes.



Grain refinement analysis in severely deformed pure iron. Orientation map with the color related to (a) x axis, (b) z axis with grain boundaries highlighted, (c) grain size distribution and (d) {111} pole figure. The color code for the orientation maps is shown on the right

In metallurgy, stainless steel, also known as inox steel, is defined as a steel alloy with a minimum of 11% chromium content by mass. Stainless steel does not readily corrode, rust or stain with water as ordinary steel does. There are different grades and surface finishes of stainless steel to suit the environment the alloy must endure. Stainless steel is used

Experimental Data

Map resolution: 10 nm

Scanned area: 5 x 5 μm

TEM type: CM 200

Crystal Structure Ferrite: cubic, a = 2.86 Å Austenite: cubic, a = 3.59 Å **ε-martensite**: hcp Martensite: Cubic, a= 2.86 Å

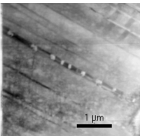


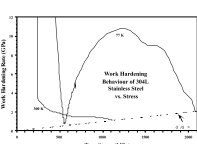
figure 2 TRIP 304 steel: Green = austenite fcc, Blue = e- martensite, red = martensite.

where both the properties of steel and resistance to corrosion are required. Some austenitic stainless steels have the spectacular property

to transform into martensite when

deformed, providing a unique combination of very high strength and ductility.

During deformation in tension, martensitic phases are formed in austenite, giving rise to a spectacular work hardening effect. Phases are clearly seen and characterized with ASTAR.



Courtesy Dr. A. Gholinia , Manchester University UK Spencer, M. Veron et al Materials Science and Engineering A 387-389 (2004)

## CHALLENGING MATERIALS FOR CHALLENGING APPLICATIONS

Texture of metals is linked to specific physical properties, so the need to characterize it at nanometer scale

Synthetic Sapphire is a single crystal form of corundum, Al<sub>2</sub>O<sub>3</sub>, also known as alpha-alumina, alumina, and single crystal Al<sub>2</sub>O<sub>3</sub>. Synthetic sapphire is the hardest of all known oxide crystals with a 9 on the Mohs scale. It is second in hardness only to diamond, and retains its strength at high temperatures. Synthetic sapphire crystals have good thermal properties, with excellent electrical and dielectric properties. In addition, it combines zero porosity with near total resistance to acids and alkaline substances. In its purest form, no porosity or grain boundaries are present, making it theoretically dense. The combination of favourable chemical, electrical mechanical ontical surface, thermal, and durability properties make san

electrical, mechanical, optical, surface, thermal, and durability properties make sapphire a preferred material for high performance systems and component designs. Due to these benefits sapphire has been used in various applications, such as sapphire wafers, lenses, substrates, watch glasses, and sapphire boule for many years. Synthetic sapphire is anisotropic hexagonal crystal. Its properties depend on crystallographic direction (relative to the optical C-axis). For various semiconductor applications, sapphire is the best choice in comparison to other synthetic single-crystals.





The challenge:

Being able to identify correctly orientations in Al<sub>2</sub>0<sub>3</sub> to study defects

Solution:

ASTAR technique coupled with precession electron diffraction

Sapphire is widely used for watch glasses. The world demand in the watch industry centres on scratchproof sapphire watch crystals. However, besides traditional flat shapes, crystals with spherical and cylindrical curves are also in great demand.

Did you know? Sapphire is the substrate for the growth of III-V and II-VI compounds such as GaN for LED's, used for LED.

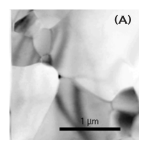


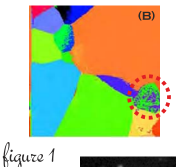
Another important application is the realization of hip prothesis, because of the biocompatibility and inert behaviour of sapphire in human bodies, combined with exceptional mechanical properties!

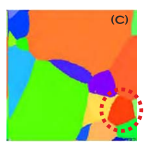
Thus, Analysis of polycrystalline sapphire, to understand the structure of defects and grain boundaries, was essential and was

performed in high resolution using the ASTAR in combination with preccesion electron diffraction technique.









Czystal Structuze  $Al_2O_3$ : Trigonal,  $R\bar{3}c$  a = 4.78 Å, c = 12.99 Å

(a) saphire grains (b) ASTAR orientation analysis without precession : orientation identification is poor, even wrong!

(c) ASTAR orientation map using 0.5° precession during acquisition allow to better capture Bragg reflection intensities and positions to analyze precisely grain orientations.





Experimental Data
TEM type: Tecnai F20
Map resolution: 1 nm
Scanned area: 2 x 2 µm

Steel degradation in nuclear plants leads to formation of new phases that is very important to monitor at nm level

## STEEL CORROSION IN NUCLEAR PLANTS

power plant

Generation IV reactors (Gen IV) are a set of theoretical nuclear reactor designs currently being researched. Most of these designs are generally not expected to be available for commercial construction before 2030. Current reactors in operation around the world are generally considered second- or third-generation systems, with most of the first-generation systems having been retired some time ago.

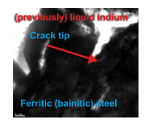
Relative to current nuclear power plant technology, the claimed benefits for 4th generation reactors include:

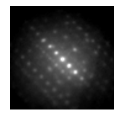
- Nuclear waste that remains radioactive for a few centuries instead of millennia
  - 100-300 times more energy yield from the same amount of nuclear fuel
  - The ability to utilize existing nuclear waste in the production of electricity
  - Improved operating safety.

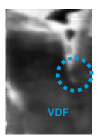
Fourth generation reactors are not yet fully designed and current research is exploring among other solutions the ability to use liquid metals as caloporters. Embrittlement of steels with liquid metals is a known phenomenon, and an issue of crucial importance for this development. Here, liquid indium was put in contact with a carbon steel, and the resulting damage was investigated in relation to the microstructure of the bainitic steel. A fine analysis

of all diffraction patterns allow to identify the presence of carbides at the crack tip as seen on the virtual bright field reconstructed with ASTAR.

While waiting for the 4th generation to be ready, it is important to study how Nuclear Centrals are ageing! Here is reported some observation of aged stainless steels aged 55000 h (6 years and 3 months) at 600°C. The specimen was prepared using FIB technique to select a specific area containing ferrite phase.





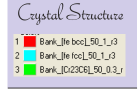


The challenge:

Identify orientation / phase in a thick specimen

Solution:

ASTAR technique coupled with precession electron diffraction





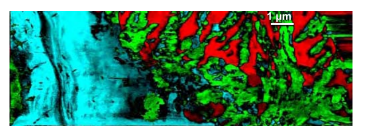


figure 1

Analysis of all the FIB section

(step of 20nm, size of the map:  $16x6 \mu m$  to record): ASTAR phase map combined with phase reliability. On the left, an austenitic grain is seen, on the right, a previously ferritic grain has decomposed into austenite and carbides.

blue = austenite fcc, red = ferrite bcc, green = carbides Experimental Data
TEM type: Jeol 3010
Map resolution: < 10 nm
Scanned area: 18 x 6 µm

## STRUCTURE OF EMBEDDED NANODOMAINS IN INTERMETALLICS

Ni-Te system is possible applied in thermoelectric generators converting heat into electrical power

Ni-Te is an intermetallic compound with possible applications in thermoelectric generators (TEGs) that convert heat into electrical power by exploiting a temperature gradient. Ni<sub>1,x</sub>Te<sub>1</sub> system is a two-solid phase mixture where nanodomains of modulated (layered) unknown phase (phase b) were found inside the NiTe matrix. Despite many attempts to solve this unknown Ni-Te phase b, its structure and stoichiometry was revealed only using 3D diffraction

The challenge:

Embedded nanodomains with size up to 70nm, two Ni-Te phases

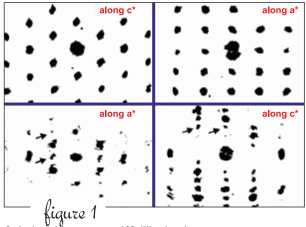
Solution:

Automated 3D Diffraction Tomography combined to Precession electron diffraction

tomography coupled to precession electron diffraction (PED).

3D diffraction tomography data were collected on the edge of a milled particle, revealed the presence of two phases, the matrix  $Ni_1Te_1$  (phase  $\alpha$ ) and the superstructure Ni5Te4 (phase  $\beta$ ). Structure determination of the modulated  $\beta$  phase was hampered by the small dimensions of the domains and the fact that most of them are embedded into the  $Ni_1Te_1$  matrix. Reflections coming from the superstructure were recognized and showed a doubling of both c and a parameters, leading to an monoclinic. It is found out that is locally similar to NiTe, but there is a shift of one Te layer by (1/3, 1/3, 0) every two c-periods referred to the initial hexagonal NiTe-type structure.

The analysis of the experimental data confirms that  $Ni_1Te_1$  (phase  $\alpha$ ) has a hexagonal structure, of NiAs type, where the Te atoms form a BCBC... hexagonal sequence. The remaining Ni atoms are distributed over octahedral sites between the Te Layers



Projections of the reconstructed 3D diffraction of phase α along the c\* & the a\* axis; (top, hexagonal) phase β along the c\* & the a\* axis (down, Moniclinic) Superstructure reflections are marked with Black arrows.



(figure 2).

The new  $\mathrm{Ni_5Te_4}$  (phase  $\beta$ ) has a monoclinic structure where the Te atoms form a sublattice with BABCBA... hexagonal sequence, while the Ni atoms occupy deformed octahedral sites with a

...c,c,a,a,c,... sequence and tetrahedral sites with a ...c,b,c,... sequence. It is found out

Crystal Structu	ire
Ni <sub>1</sub> Te <sub>1</sub>	Ni <sub>5</sub> Te <sub>4</sub>
(α-phase)	(β-phase)
Hexagonal P6,/mmc	Monoclinic Pm
a= 3.85 Å b= 3.85Å	a= 6.95 Å b= 4.02 Å
a= 3.85 Å	a= 6.95 Å b= 4.02 Å c= 11.96 Å
a= 3.85 Å b= 3.85Å	a= 6.95 Å b= 4.02 Å c= 11.96 Å α= 90°
a= 3.85 Å b= 3.85Å c= 5.22 Å	a= 6.95 Å b= 4.02 Å c= 11.96 Å

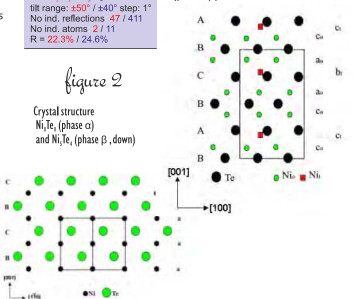
Experimental data

Ni<sub>1</sub>Te<sub>1</sub> / Ni<sub>5</sub>Te<sub>4</sub>

that is locally similar to NiTe, but there is a shift of one Te layer by (1/3, 1/3, 0) every two c-periods referred to the initial hexagonal NiTe-type structure.

ADT together with PED allowed the structure

determination of the nickelrich phase superstructure  $Ni_5Te_4$ (phase  $\beta$ ) for the first time.



High pressure torsion effects in Al-Cu metastable alloy at nm scale can be observed in orientation maps

### METALS UNDER HUGE STRESS

Silver and copper have a wide miscibility gap therefore special methodologies are needed to obtain a metastable alloy. The usual results of a Cu-Ag alloy synthesis is spinodal decomposition of Ag, Cu phases intergrown in bands some hundred nanometers wide. A way to push the spinodal decomposition to a smaller scale is given by the severe plastic deformation induced by a high pressure torsion (HPT).

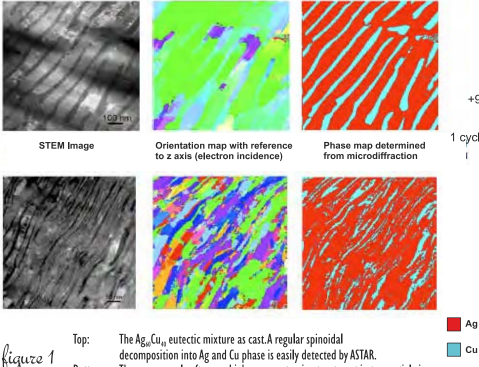
Several cycles of HPT mixed together the Ag and Cu bands in smaller grains of an Ag-Cu alloy. This process is extremely difficult to map in terms of phase separation, grain size and crystal orientation since it occurs on a nanometer scale.

The ASTAR technique can follow the processes in all phases. In the as cast starting mixture, an alternating sequence of Ag and Cu lamellae are detected. The orientation mapping reveals a strong orientation correlation between the lamellae. After one cycle of HPT the lamellar structure is hardly recognizable in planar view and becomes thinner and irregular in sections tangential to the deformation cylinder. The deformation induced by the torsion also changes the grain size and the texturing.

Identify orientation / phase of 2 different The challenge: phases particles with < 50 nm Solution: ASTAR technique

Within each lamella some small grains with irregular shape start to appear. The orientation correlation between the lamellae is strongly reduced.

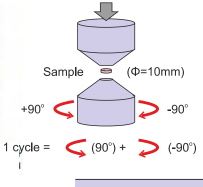
The process can be characterized with ASTAR at each step until a complete alloy is formed after 10 HPT cycles.



The same sample after one high pressure torsion treatment in tangential view.



**Schematic Illustration** of HPT process



Crystal Structure Silver: cubic, a = 4.086 Å Copper: cubic, a = 3.615 Å

Experimental Data TEM type: Tecnai F20 ST Map resolution: 4 nm Scanned area: 800 x 800 nm

The spinodal decomposition bands become thinner and less corrected.

Ag

Bottom:

### PAVEMENTS OF THE NANOWORLD

Texture patterns of electrodeposited Ni-based mesoscale structures resemble the patterns seen in cobblestone pavements

One effective method of producing nanocrystalline materials (grain size < 100 nm) is pulsed electrodeposition. It has been observed that electrodeposits form and grow in a nodular fashion each nodule representing a "colony" of nanosized grains (Cizek et al.). Orientation measurements performed in the vicinity of the boundaries separating two neighboring colonies suggest that the colonies in a nanocrystalline Ni-20% Fe electrodeposit might display a distinct "cobblestone" mesotexture characterized by a <001> fiber axis approximately perpendicular to the local curvature of the colony growth surface.



TEM has been used to investigate the local microtexture and misorientation characteristics of nanosized grains comprising mesoscale colonies in electrodeposited Ni, Ni-20 % Fe, and Ni-50 % Fe materials. Optical microscopy revealed the presence of mesostructure colonies in all three Ni-based electrodeposits studied. Consequent TEM analysis on specimen sections parallel either to the Deposition Plane (DP) or the Deposition Direction (DD) provided the size and shape distribution of the nanograins for all the materials studied.

isbon Pavement, Portugal

ASTAR analysis indicated that both the bulk texture and the local microtexture and mesotexture of the as-deposited specimens are dominated by the <001>//DD fiber, which has also been observed in many other experimental studies.

The challenge:

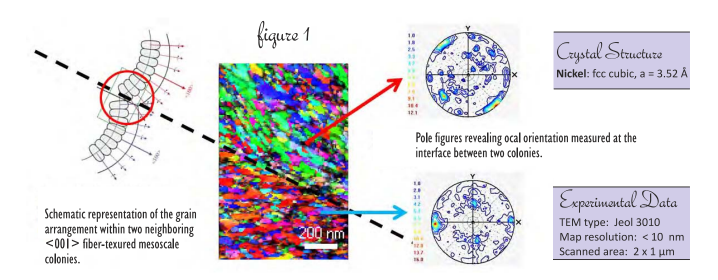
Identify the orientation of crystallites with size < 50 nm

Solution:

ASTAR technique coupled with precession electron diffraction

Researchers attribute this fact to the influence of hydrogen codeposition which is more active in {001} than {111} crystallographic planes. For all three alloys, the coarsened grains, obtained by annealing, within the mesoscale colonies showed with EBSD-SEM a fiber mesotexture characterized by a <111> axis approximately perpendicular to the colony hemispherical growth surface (parallel to the local DD). From such EBSD-SEM measurements it has been

surmised that the as-deposited texture patterns in Ni-based materials might resemble the patterns often seen in cobblestone pavements found in many European cities. This has indeed been confirmed by ASTAR orientation measurements made on a local nm scale in the as-received materials (see the figure below). Further work is needed to reveal the exact formation mechanisms of this cobblestone mesotexture as the aforementioned features will likely affect both the homogeneity and anisotropy of mechanical and physical properties of the nanocrystalline electrodeposits and many properties such as corrosion resistance, toughness or ductility, coefficient of friction and wear resistance could be affected by the characteristics of the mesoscale colonies.



Understanding the Li insertion / extraction mechanism at microscopic level is important to design next generation batteries

# UNDERSTANDING LI INSERTION MECHANISM FOR NEXT GENERATION BATTERIES

Lithium iron phosphate (LiFePO $_4$ ) is a good candidate for positive electrodes in lithium-ion batteries to be installed in new electric vehicles. Its cost is low, it has an excellent life cycle and it is safe. To obtain the best performances in terms of electrical conductivity it is synthesized in nanocomposite with grain size between 50 and 300nm.

The discharge-charge mechanism implies a delithiation of LiFePO<sub>4</sub> into FePO<sub>4</sub> and a successive insertion of Li into the iron phosphate. The mechanism of Li insertion / extraction at the microscopic level wasn't confirmed and at least four different models have been proposed.



The difference between the various models lies in the relative position, shape and dimension of the two phases, LiFePO<sub>4</sub>, FePO<sub>4</sub> during the process. Two of these models imply core shell structures (bulk or radial), another one a spinoidal decomposition at the nanoscale level of two Li and Fe rich end members (spinoidal model), and the last one where the nanoparticles are either fully lithiated or fully delithiated (Domino-cascade model).

A phase mapping technique able to identify the lithiated and delithiated phase at the nanoscale can obviously solve the dilemma. Unfortunately the problem is complicated by the very similar unit cell parameters of LiFePO<sub>4</sub> and FePO<sub>4</sub> which differ from each other by

The challenge:

Solution:

Identify Li doped particles between two different phases having <5 % difference in cell parameters and particle size <50 nm

ASTAR technique coupled with precession electron diffraction

less than 5%. ASTAR system can solve the problem by collecting the NanoBeam Electron Diffraction (NBED) patterns in precession mode.

Precession diffraction, on the one hand reduces the dynamical effects so that the patterns are closer to a kinematical simulation of the templates and on the other, it enables the collection of a higher number of

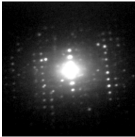
reflections from high order Laue zones reducing the ambiguity in indexing the patterns. In this way ASTAR system has been able to detect that the nanocrystalline grains of the nanoparticles are either fully lithiated or fully delithiated giving strong support to the domino-cascade model (fig. 2). The results have been confirmed also using HRTEM imaging and EFTEM / EELS analysis.

Czystal Structuze

LiFePO₄: Orthorhombic
a=10.33 Å, b=6.01 Å, c=4.69 Å

FePO₄: Orthorhombic
a=9.81 Å, b=5.79 Å, c=4.78 Å

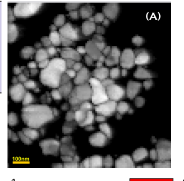
Experimental Data
TEM type: Jeol 2010 FEF
Spot size: 2.7 nm
Step: 5 nm
Scanned area 1 x 1 µm



00

figure 1

Experimental precessed patterns and its indexation with the LiFePO $_4$  (red) and FePO $_4$  (green) templates with the corresponding agreement factors.



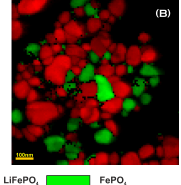


figure 2

Reconstructed bright field image (a), phase determination map (b). Crystal grains of the nanocomposite mixture of LiFePO<sub>4</sub> (red), FePO<sub>4</sub> (green) during different phases of the charge-dicharge cycle.

There is no mixing of the two phases inside the same crystal grain.

This work was done by collaboration between the LETI, LITEN, INAC (CEA Grenoble - Minatec/PFNC -Nanocaracterisation Center) and UJF (Grenoble). This work has been supported by the French National Research Agency (ANR) through the AMOS Program

## NANOSCALE REACTION MODELS IN SOLID OXIDE FUEL CELLS

Reaction models on fuel cells can be established based on detailed orientation / phase map analysis

Solid oxide fuel cell (SOFC) technology is a promising energy conversion option at a time when efficient solutions are sought. During the electrochemical reaction of a fuel with an oxidant gas (usually air) at an operation temperature of  $\sim 800^{\circ}$ C, SOFCs cogenerate heat and electricity with high efficiency: Electric efficiencies close to 60% are obtained now with a SOFC alone. When coupled with a micro gas turbine (SOFC-GT), these can increase up to 70%. Advantages of SOFCs also include flexibility of fuel use (e.g. CO, H<sub>2</sub>, CH<sub>4</sub>), delocalization of energy conversion, and reduction of pollutants such as NO<sub>x</sub> and SO<sub>x</sub>.



A SOFC is composed of three main components: an anode, an electrolyte and a cathode. The standard design is based on the use of a porous ceramic-metal (cermet) composite anode made of yttria  $(Y_2O_3)$ -stabilized zirconia (YSZ) and Ni.

The anode acts as a fuel oxidation catalyst, hydrocarbon reforming catalyst, current collector and often as a mechanical supporting layer of the whole fuel cell. NiO particles are sintered with YSZ and then reduced by the fuel to metallic Ni during the first operation of the cell. During service life, minimum 5 years at ~ 800 °C, the anode runs on fuel and Ni is kept in its reduced state.

In this project, environmental transmission electron microscopy (ETEM) is used to understand the reaction mechanisms and the

The challenge:

Solution:

Monitoring crystallographic changes after reduction of fuel cell: orientation / phase of two different phases with details of 1 nm

ASTAR technique coupled with precession electron diffraction

Czystal Structuze

Y<sub>2</sub>O<sub>3</sub>: Cubic, Ia3, a = 10.58 Å

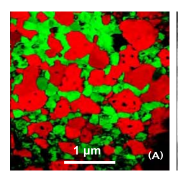
NiO: Cubic, Fm3m, a= 4.168 Å

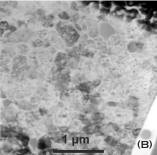
Ni: Cubic, Fm3m, a = 3.52 Å

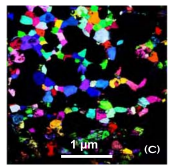
Experimental Data
TEM type: Jeol 2200 FS
Map resolution: 1 nm
Scanned area: 4 x 4 µm

structural evolution of the as-sintered NiO-YSZ anode during reduction. A thin window, which is prepared using a dual beam FIB/SEM, is reduced inside the ETEM in a total pressure of 1.3 mbar of H<sub>2</sub>. NiO reduction is observed to start at the grain boundaries with the YSZ phase. Pores form within the NiO grains to accommodate the volume shrinkage linked to the removal of oxygen. A post mortem analysis using ASTAR orientation imaging allows monitoring of the crystallographic changes induced by reduction at nm scale. It can be seen that the reduced Ni grains maintain the same orientation as the

initial NiO grains, while the YSZ phase is left unchanged by the reaction. Based on these observations, reaction models can be established.







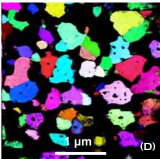


figure 1

- (A) ASTAR phase map where both NiO phase (red) and YSZ phase (green) can be seen
- (B) same area TEM bright field image
- (C) ASTAR orientation mapping for NiO phase and (D) ASTAR orientation map for YSZ phase.

Novel twin-modulated structures in high SFE metals using polar ceramic layers need to be investigated at atomic level

## ATOMIC SCALE OF NANOLAYER COMPOSITES

A stacking fault is a one or two layer interruption in the stacking sequence of the crystal structure. These interruptions carry a certain stacking fault energy (SFE).

High SFE materials such as Al do not form twins easily. However, researchers have shown that it is possible to form Al layers in twin orientation to each other across polar TiN layers, if these are grown so that both the Al and TiN layers have a  $\{111\}$  surface as their growth front. Al-TiN multilayers were deposited at room temperature on Si substrates using dc magnetron sputtering in the two following different layer thicknesses: Al-9nm/TiN-1nm .

HREM-STEM image I collected by a HAADF detector, showed two Al layers separated by a TiN layer. It was seen that the Al layer is always the one that has

a twin orientation with respect to the TiN and Al layers below it. The situation is schematically illustrated in the figure below. Theoretical ab initio calculations based on density functional theory (DFT) revealed that nitrogen termination in the {111} growth plane of the TiN layers greatly favors the growth of twin oriented Al layers on them, thus, being in excellent agreement with experimental observations.

In conclusion, it was shown that in magnetron sputtered Al-TiN multilayers, consecutive Al layers grow in twin orientation with respect to each other when the intervening TiN layers are 1–2 nm thick.

The above HREM experimental data were explored further by means of the ASTAR system that was able to produce orientation /

The challenge: Solution: Identify orientation / phase of 1 nm TiN layer in a Al-9nm / TiN-1nm nanolayer composite while TiN and Al differ <5% in cell parameter

ASTAR technique coupled with precession electron diffraction

(A) (B) (B) (D) (D) (D)

phase maps of the Al-TiN nanolayers revealing clearly the TiN layer of 1 nm thickness. ASTAR orientation maps show clearly the twin orientation relationship between succesive Al layers.

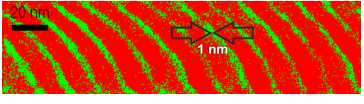
Moreover, these findings suggest a method of introducing nanoscale twins in high SFE materials in general. Since the deposition of Al and TiN layers is used in the formation of diffusion barriers, it is important to investigate and understand these structures at the nanometer length scale and hence to be able to control them.

Crystal Structure

Al: fcc cubic a = 4.05 Å

TiN: fcc cubic a = 4.24 Å

Experimental Data
TEM type: Jeol 2100 F
Map resolution: 1 nm
Scanned area: 200 x 200 nm





(a) HAADF image of Al-9nm/TiN-1nm multilayer system (b) atomic model of the Al-TiN-Al interface (c) ASTAR orientation map where Al layers are shown in red, TiN 1nm layer shown in green (d) line profile of ASTAR orientation map showing twin-related Al layers (E) ASTAR orientation map with 1nm TiN layers shown as black strips

## NEXT GENERATION CIRCUITS FOR SEMICONDUCTOR INDUSTRY

The scaling required to reach faster chip performances in electronic devices has pushed the dimensions of copper interconnect (CI) lines to the nanometer domain. This constant downscaling of CIs implies a change in their microstructure. A change in the grain boundary type distribution and local texture will strongly influence the resistivity and the mechanical reliability of downscaled CIs. A different texture can imply different mechanical properties and a different local distribution of stresses. It is, therefore, necessary to map the texture evolution with the size of CIs.

Unfortunately, CIs that reach lateral sizes of less than 100 nm, with Cu grains smaller than 50 nm, cannot be characterized by conventional techniques like EBSD and XRD, because they do not have enough spatial resolution.

For reliable texture quantification, it is important to acquire extensive and reliable data sets to have statistically meaningful results and at the same time have a high spatial resolution. ASTAR with a parallel nanosized beam, coupled with a small precession angle, has the appropriate spatial resolution (1 nm) and the reliable pattern indexing which reduces the 180° ambiguities by sampling reflections from higher order zones

(HOLZ). In addition, ASTAR is able to acquire data and index patterns rapidly, which makes it possible to acquire statistically relevant information.

The challenge:

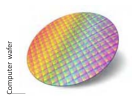
Characterize Copper Interconnect lines (CI) grain boundaries and texture at 1nm scale

Solution:

ASTAR technique coupled with precession electron diffraction

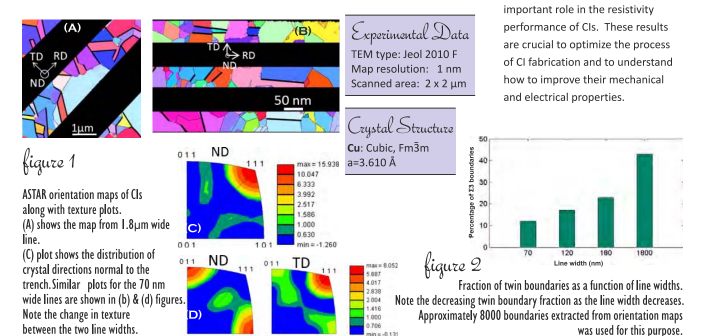






ASTAR orientation maps reveal the evolution of texture as the CI are scaled from CIs from 1.8  $\mu$ m to 70 nm. In large CIs the Cu grains have a strong <111> fiber texture normal to the trenches (Fig. 1). In the smallest CIs, a <110> texture is observed, normal to the trench

while {111} planes are now parallel to the trench sidewalls (Fig. 1). The microstructure also changes from a bamboo-like structure in large CIs into a polygranular structure in small CIs, where a polygranular microstructure is a microstructure in which there are continuous grain boundary paths along the length of the interconnect. In the 70 nm CIs grain size is not uniform and clusters of small grains are formed at the bottom of the trenches. Such clusters of small grains adversely affect the reliability of CIs. In addition, a decrease in the fraction of coherent twin boundaries was observed with decreasing line widths (Fig. 2). Twin boundaries play an



Spintronic device properties strongly depend on a nanometer scale structure

Solution:

### STRUCTURE OF MAGNETIC **SEMICONDUCTORS**

Magnetic semiconductor materials have attracted widespread attention in recent years due to their potential applications for the transport of information by exploiting both the intrinsic spin of the electron and its associated magnetic moment, in addition to its fundamental electronic charge. The new technology which emerged from discoveries in the 1980s concerning spin-dependent electron transport phenomena in solid-state devices was called Spintronics ('Spin Transport Electronics') also known as magnetoelectronics.

The material system (Mn,Ga)As on GaAs is of great interest for the development of such spintronic devices because the unstrained-bulk-MnAs is ferromagnetic at room temperature (α-MnAs, P6<sub>3</sub>/mmc). It undergoes a

phase transition to paramagnetic phases at 40°C (β-MnAs, Pnma) and above 125°C (γ-MnAs, P6<sub>3</sub>/mmc)). "Surface contactable quasiembedded" MnAs crystallite precipitates are formed in a [001] - oriented GaAs matrix during the cooling down stage of a MnAs layer that was deposited by metal-organic chemical vapor deposition (MOCVD) above 600 °C.

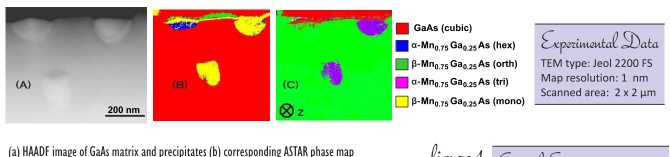
In order to obtain a thorough understanding of the (Mn,Ga)As crystallite formation process, the crystallographic phase and orientation of the precipitates with respect to the matrix have to be determined. The chemical composition of the precipitates was determined by quantitative EDXS analysis, their chemical composition was defined as Mn<sub>0.75</sub>Ga<sub>0.25</sub>As.

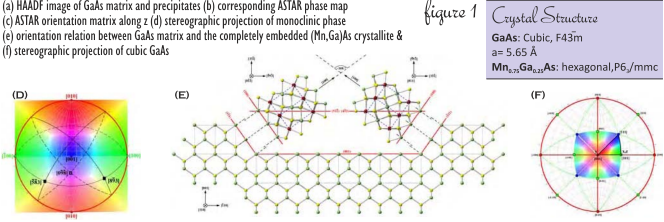
Identify unknown phase of MnGaAs precipitates The challenge: among 4 different crystal phases & establish orientation relation between GaAs matrix and precipitates ASTAR technique coupled

with precession electron diffraction

A careful inspection of the PED diagrams on the precipitates showed that there existed super structure reflections not consistent with MnAs bulk structure. ASTAR template matching analysis with all possible templates of 4 possible phases compatible with Mn<sub>0.75</sub>Ga<sub>0.25</sub> As

(total 38561 templates) shows a coexistence of precipitates with the monoclinic β-phase (space group P2,/m) and of precipitates with the hexagonal α-phase (P6<sub>3</sub>/mmc) at room tempereture. An example of ASTAR phase and orientation analysis of the embedded (Mn, Ga)As-crystallite precipitates in GaAs is illustrated below.





## HYBRID INORGANIC / ORGANIC SYSTEMS FOR OPTOELECTRONICS IN DAILY LIFE

Optical properties of hybrid systems depend on their nm level structural properties

Nanocrystalline ZnO is provided in an inorganic/organic system of ZnO deposited on sexiphenyl (6P). Initially 6P was deposited on a-plane ZnO substrate at room temperature forming mesoscopic 1-dimensional islands of 6P. In a second step, the islands were capped with ZnO via molecular beam epitaxy at 373 K.

Information on the nanostructure of ZnO grown atop 6P was provided by high-resolution TEM. Imaging was performed along the [00.1] direction of the ZnO substrate which showed that ZnO exhibits a perfect crystal structure when directly grown on the substrate. This was confirmed by Fourier transformation (FT) which also confirmed that ZnO grown atop 6P undergoes an imperfect growth.

Identification of the orientation of the nanocrystals and indexing of the lattice planes only based on FT is rather ambiguous and must be handled with care. For this reason, ASTAR was used to uncover any specifics of the overgrowth regime of ZnO on 6P at nm scale.



With ASTAR scanning the electron beam in nanobeam diffraction mode (probe diameter 1 nm) across an area of 100 nm in width and 80 nm in height provides the orientation maps depicted below. The maps were derived by comparing the diffraction pattern recorded at each individual position with a set of about 1000 calculated diffraction patterns of hexagonal ZnO. Focussing on ZnO atop 6P, columnar grains are identified that extend even up to the surface as signified by green and red areas.

The challenge:

Identify precise orientation texture between ZnO substrate - sexipephenyl (6P) - ZnO deposited layer at nm scale

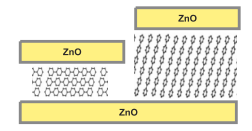
Solution:

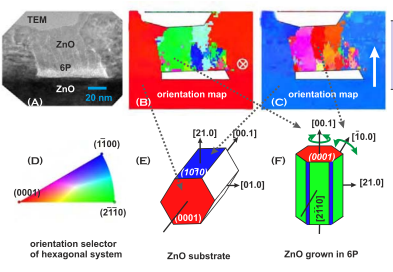
ASTAR technique coupled with precession electron diffraction

prismatic facets contribute to the diffraction pattern.

Research on ZnO is of immediate practical importance as this semiconductor has several favorable properties, including good transparency, high electron mobility, wide bandgap, and strong room-temperature luminescence. These properties are used in emerging applications for transparent electrodes in liquid crystal displays, in energy-saving or heat-protecting windows, and in electronics as thin-film transistors and light—emitting diodes (LEDs).

From the ASTAR orientation map it is evident that the [00.1] wurzite axis is well-aligned along the vertical growth direction indicating a strong texture. Electrostatic forces originating from the 6P molecules have to be considered as the origin of this texture. In addition, a significant lateral texture within the (0001) plane cannot be excluded, as it cannot be distinguished which of the six





Crystal Structure

ZnO: Hexagonal, P6₃mc
a = 3.22 Å, c= 5.20 Å

Experimental Data
TEM type: Jeol 2200 FS
Map resolution: 1 nm
Scanned area: 100 x 80 nm

Crystallographic orientation mapping of ZnO

as revealed by scanning nanobeam
diffraction (SNBD):

a) TEM image of the region analyzed by SNBD.
b) and c) crystallographic orientation maps of ZnO.
The white arrow markers represent the lattice planes color coded by the orientation triangle of d).
e) alignment of a representative ZnO unit of both, ZnO substrate and low temperature grown homoepitaxial ZnO, f) alignment of a representative ZnO unit for the textured growth atop sexiphenyl.

Nanoparticle size and texture are very important for drug delivery of poor water soluble drugs

## DRUG DELIVERY & STRUCTURE OF NANOPARTICLES

Iron oxide nanoparticles (usually below 20nm) are being investigated for biomedical applications because of their biocompatibility and unique physical properties. This biocompatibility results from their capacity for metabolism, transportation and storage in human tissues. Magnetic iron oxide particles with diameters above 20nm are ferromagnetic, i.e. they have a reminiscent magnetic moment that generates unstable particulate suspensions, as the particles aggregate together into biologically toxic, micron-sized clusters.

Small Fe<sub>2</sub>O<sub>3</sub> below about 20 nm are superparamagnetic, which means thermal energy is sufficient to cause large fluctuations of their magnetic

moment, so that in the absence of an magnetic field, the particles have no effective moment. This reduces inter-particle attraction and makes stable, biocompatible suspensions possible. These particles maintain a strong response to an external magnetic field, which make them ideal candidates for biomedical applications, e.g. improve effectively the poor water solubility of hydrophobic drugs.

Colloidal systems, including nanoparticles and liposomes, have been extensively studied as potential drug carriers for targeted or controlled release. Monitoring of the nano-assembly will facilitate an indirect determination of the site at which the therapy is administered.

The texture of nanoparticles is therefore the utmost importance to control undesired agglomeration effects for drug delivery applications. ASTAR has been used in order to reveal successfully 1 nm scale details of very small nanoparticles (Pt of

Solution:

The challenge:

Identify orientation textufe of very small <5nm nanoparticles & distinguish between particles of same cells but different space groups

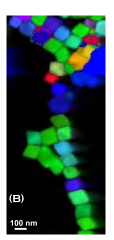
ASTAR technique coupled with precession electron diffraction

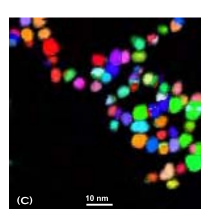
5-10nm in size and magnetite  ${\rm Fe_3O_4}$  parttcles of 100nm in size (figure 1).

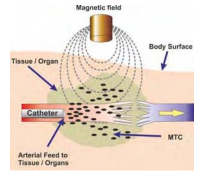
On the other hand, ASTAR has been used successfully to distinguish between a mixture of structurally closely related nanoparticle structures (e.g magnetite and maghemiite having essentially the same unit cell dimensions but different space groups, see table).HR-

TEM study is possible to use to distinguish between both phases, but is too time-consuming. Automatic ASTAR orientation / phase map analysis by comparing experimental PED patterns with both maghemite /magnetite templates , reveraled higher reliability for magnetite crystals (results confirmed independently by HREM techniques). Collection of quasi-kinematical PED was essential to distinguish between patterns that belong to different Space Groups and extinctions may become visibe with ZOLZ – HOLZ reflections.









Czystal Stzuctuze

Magnetite Fe<sub>3</sub>O<sub>4</sub>: Cubic,
Fd3m, a=8.32 Å

Maghemite y-Fe<sub>2</sub>O<sub>3</sub>: Cubic
P4,32, a=8.33 Å

Experimental Data

TEM type: Tecnai 20 & Jeol 2010F

Map resolution: 1 nm Scanned area: 2 x 2 μm

figure 1

(a) ASTAR orientation mapping of magnetite nanoparticles along z direction

(b) along x direction (c) Pt nanoparticles along z direction

## INSIDE THE NANOWIRE FOREST

Understanding texture at nm level is crucial to foresee and design the proper electrical and transport properties of nanowire heterostructure.

III-V semiconductor nanowires are at the forefront of the solid state physics and technology (SEM image on the right). Nanowires are quasi one dimensional crystals that can be heterostructured in different sectors having different chemical composition and crystal structure. They are grown by chemical beam epitaxy (CBE) from a III-V wafer with the Au-assisted growth method. In this method the wafer is coated with a thin layer film that by a dewetting thermal treatment in the CBE chamber transforms into small Au nanoparticles.

The Au nanoparticles act as a catalyst and drive the growth of the nanowire out from the wafer. At the final stage of the growth the wafer is a forest of nanowires each with its Au nanoparticle on top

The challenge:

Distinguish structurally similar phases & perform orientation / phase of two different phases in a nanowire

Solution:

ASTAR technique coupled with precession electron diffraction

Experimental Data
TEM type: Zeiss Libra 120
Map resolution: 10 nm
Scanned area: 3 x 3 µm

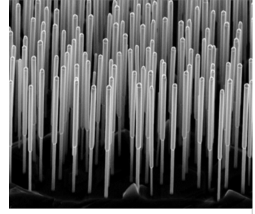
Czystal Structure

InSb: Cubic,  $F\overline{4}$ 3m

a = 0.64 Å

InAs: Hexagonal,  $P6_3$ mc

a= 4.27 Å c= 7.02 Å



SEM image of a forest of InAs / InAsSb heterostructured nanowires

(SEM image at the right).

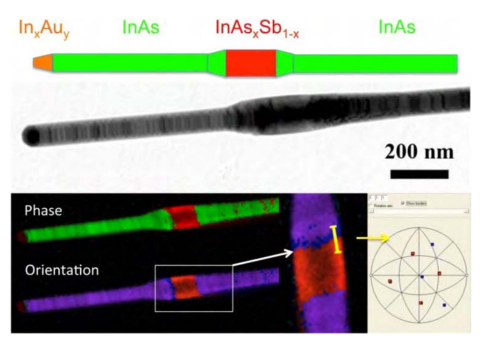
The growth of heterostructured nanowires is a challenge, especially if the two crystal structures, which must be piled up on top of each other, have a large lattice mismatch. This is the case of the InAs/InAs1-xSbx/InAs heterostructure, which can only be grown in nanowires, where the strain can be relaxed at the free surface, but not in a two dimensional growth. ASTAR is able to identify where the cubic Sb sector is located between the bottom and top hexagonal InAs stems. It determines the relative orientation of the two

crystal structures and is able to recognize twinned layers inside the cubic Sb sector (fig. 1).

These structural details are crucial to foresee and design the proper electrical and transport properties of the heterostructure.

### figure 1

(Top) Model and bright field image of a InAs / InAs<sub>1-x</sub>Sb<sub>x</sub>/ InAs heterostructured nanowire. (Bottom) Phase and orientation mapping of the same wire. In the phase map the cubic zinc blend structure of the Sb part is identified (red) while the orientation map shows that is separated from the hexagonal wurzite. InAs stem by a twinned sectorof 30 nm (blue). At the bottom left a pole figure calculated trough the twin boundary is displayed



Revealing nanoparticles texture at nm level helps to understand and predict their optical properties

## TEXTURE SECRETS OF NANOPARTICLES

Nanoparticles are now a hot topic in several different fields for their astonishing and unusual properties compared to bulk materials. New sciences like nanochemistry, nanomedicine and nanotoxicology are born and an accurate knowledge of the nanoparticle structure is required to model properly their useful properties. Among these, the grain size, the defects and the particle shape are of major importance, since they easily correlate especially with optical properties.

The knowledge of the grain size of nanoparticles can be easily obtained with conventional TEM techniques, but the coherent domains, those inside where the structure is defect free (no twinning or grain boundaries), are much more difficult to recognize and measure.

The nanoparticle shape is another important factor to be known, since it drives the self-assembly. All these structural details call for a characterization technique that is able to furnish crystallographic information with nanometric resolution. ASTAR technique is able to supply this information.

Two examples of orientation mapping obtained by ASTAR on nanoparticles are here displayed. In fig. 1 is showen an orientation map obtained on NaYF4 nanoparticles. These nanoparticles are known as upconverting, since they are able to emit visible and near-infrared light when excited with near-infrared radiation. They are very promising as fluorescent targets for biological imaging. NaYF4 crystallizes in two forms one cubic and one hexagonal. ASTAR identifies all

the nanoparticles as hexagonal. Those that have a hexagonal shape are oriented with the c axis (the 6 fold axis) normal to the plane, while those that are rectangular are oriented with the c axis parallel to the plane. This indicates that the nanoparticles are hexagonal prisms with a height comparable

to the base sides. The second example shows an orientation map of nanogold (fig. 2).

ASTAR identifies most of the nanoparticles as oriented with the [100] direction normal to the plane, indicating that they have well developed flat {100} faces on which they fall. Some of them exhibits {111} twinning as shown by the pole figure plot at the right (fig. 2c).

orientation map of NaYF, nanoparticles (Top) Index map (Bottom) Orientation map.

The hexagonal (A) nanoparticles are 'red', corresponding to an orientation with the c axis normal to the plane.

The rectangular (B) are 'green — blue', corresponding to orientations with the c axis on the plane.

The challenge:

Identify orientation & phase of nanoparticles with < 40 nm that crystallize in two different phases

Solution:

ASTAR technique coupled with precession electron diffraction

Experimental Data
TEM type: Zeiss Libra 120
Map resolution: 10 nm
Scanned area: 2 x 2 µm

Czystal Structuze

NaYF4: Hexagonal P6
a= 5.91 Å c= 3.50 Å

figure 2

Orientation map of Au nanoparticles.
a) Index map b) Orientation map

c) Pole figures of [111]. Equivalent directions of the two twins displayed

(A) (B) (C)

in the box at the center. Most of the nanoparticles have a "red" orientation corresponding to the [100] direction parallel to the plane normal. Therefore most of them have {100} flat faces on which they fall.

## CRYSTAL ORIENTATION IMAGING OF ORGANIC NANOMAGNETS

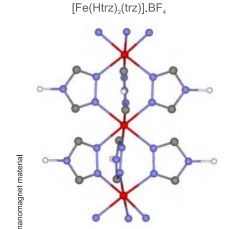
Organic nanomagnets are lighter than conventional magnets and need new techniques for structure characterization

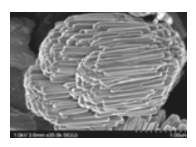
The development and characterization of nanometer-scaled functional molecular-based materials, which may exhibit fascinating magnetic, optical or electrical properties, have recently attracted much attention. Molecular nano-objects offer the possibility of tuning or combining these properties by a rational synthetic chemical approach. These materials present promising technological applications in molecular electronics, or magneto-optic memory devices providing that we can design single domain particles with hysteresis behaviour in the future.

Nanorods of the spin crossover material [Fe(Htrz)<sub>2</sub>(trz)].BF<sub>4</sub> are prepared by arrested growth within liquid crystal phases. A complete understanding of the crystal growth requires the identification of the orientation of the crystal axes with respect to the morphology of the nanoparticles.



The intrinsic physical properties (magnetic and optical hysteresis) rely on the domain structure of the nanoparticle. Molecular nanoparticles of [Fe(Htrz)<sub>2</sub>(trz)].BF<sub>4</sub> are extremely sensitive to the e-beam so that ASTAR at high frame rate is the only possible technique which allows the recording of the diffraction pattern before the destruction of the probed nanoparticle occurs.



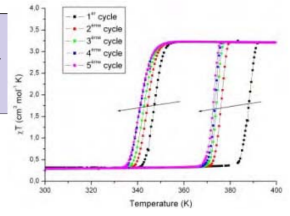


The challenge:

Identify if nanocrystals diffract and whether they are single crystal particles / mapping of the unit cell disorientation along the nanoparticle.

Solution:

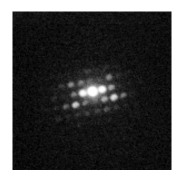
ASTAR technique at high frame rate >100 fps, no cryo-holder is used



1 μm

1 μm

Experimental Data
TEM type: Jeol 3010
Map resolution: 10 nm
Scanned area: 2 x 2 µm



Czystal Stzuctuze
Orthorhombic, Cmcm a = 9.37 Å, b = 17.05 Å c = 7.34 Å

A complete mapping of the diffraction pattern on a single nanoparticle shows that the particle is almost single domain, and that the disorientation of the

crystal lattice is limited to only 5%. ASTAR also shows that the disorientation is not progressive, but rather restricted to

Nanoparticles structural identification reveal the manufacturing process of ancient ceramics

### INSIDE A ROMAN CERAMIC FACTORY IN ANCIENT GAUL

Terra sigillata is certainly the most famous fine ware of the Roman period with characteristic decorations obtained by means of specially designed stamps (sigilla). This pottery was extensively produced in standardized shapes by a few specialized workshops and was widely distributed across the Roman Empire. Below is shown the diffusion map of one of these large workshops.

The success of this pottery was mainly because of the brightness and red color of the slip. Also in order to better understand the technical specificities of Terra sigillata, TEM and other characterization techniques have been used to investigate the microstructure and microcomposition of Roman period Terra Sigillata slips from various workshops.

A high proportion of spinel (≥25%) is characteristic of Italic productions (1st c. BC) while a high content of Fe-rich corundum(>30%) is the brand of south Gallic workshops (1st c. AC). This systematic difference is due to an evolution in the manufacturing process which permitted them to obtain a slip more resistant to the abrasion. ASTAR phase analysis indicates strongly that the ceramic sample provenance is from a factory at Graufesenque (South Gaule).

The challenge:

Determine the nature, size and orientation of the different nanocrystallized phases of the slip

Solution:

ASTAR technique coupled with precession electron diffraction



figure 1 Fragment of a decorated cup in Terra Sigillata



Experimental Data TEM type: CM 20 FEG Map resolution: 2 nm Scanned area: 2 x 2 µm

0.2 μm

figure 2

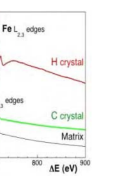
Crystal Structure Spinel, MgAl<sub>2</sub>O<sub>4</sub>: Cubic, Fd3m a= 8.09 Å Corindon, Al<sub>2</sub>O<sub>3</sub>: Hexagonal, R3c a=4.76 Å, c=13.0 Å Hematite, Fe,O<sub>3</sub>: Hexagonal, R3c a=5.03 Å, c=13.75 Å

TOP: ASTAR high resolution phase (b) and orientation (c) maps for the crystals present in the slip of a Gallic Terra Sigillata from La Graufesenque workshop.

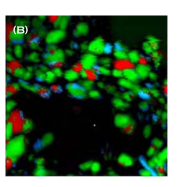
The slip coating consists of hematite (in red), corundum (in green) and spinel (in blue) crystals in a glass matrix (b). In contrast with the coatings of other types of Greek and Roman potteries, there is no particular orientation of crystals.

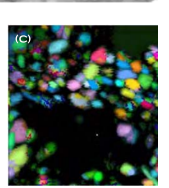
Courtesy S. Joulié, C. Roucau & Ph. Sciau, CEMES-CNRS Toulouse, France

LEFT: EELS analysis of the matrix, hematite and corundum crystals.



Fe L edges





Ph. Sciau et al J. Am Cer Soc 89 (2006) 1053, Y. Leon et al JRS 41 (2010) 1550

## SHED LIGHT TO EARTH SEISMICITY

Texture of olivine studied under high pressures help to understand Earth's mantle tectonic properties

The mineral olivine dominates the composition of the Earth's upper mantle and hence controls its mechanical behaviour and seismic anisotropy. Experiments at high temperature and moderate pressure, and extensive data on naturally deformed mantle rocks, have led to the conclusion that olivine at upper-mantle conditions deforms essentially by dislocation creep with dominant [100] slip. The resulting crystal preferred orientation has been used extensively to explain the strong seismic anisotropy observed down to 250 km depth.It has been shown that this high-pressure dislocation creep produces crystal preferred orientations resulting in extremely low



Earthquake, Santiago, Chile, 27 October 2010

seismic anisotropy, consistent with seismological observations below 250 km depth.

These results raise new questions about the mechanical state of the lower part of the upper mantle and its coupling with layers both above and below. In order to study the structural modifications /textures of olivine at pressures that exist at depths of 200-300 km below the surface the DIA deformation equipment coupled at Synchrotron X-Ray source was used .

As olivine is stable at ambient conditions, a post-mortem (after application of strong e.g 40 % deformation at 1373 K and 3 GPa ) analysis with ASTAR in TEM has been performed in order to reveal the texture at nanoscale size.

The sample studied was a forsterite; in fact, olivine is actually a name for a series

between two end members, fayalite and forsterite. Fayalite is the iron rich member with a pure formula of Fe<sub>2</sub>SiO<sub>4</sub>. Forsterite is the magnesium rich member with a pure formula of Mg<sub>2</sub>SiO<sub>4</sub>. The two minerals form a series where the iron and magnesium are substituted

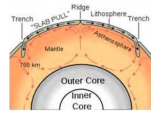
for each other without much effect on the crystal structure. Besides the high number of defects formed during DIA deformation cycles, it has been perfectly possibly to study in detail with ASTAR the texture of the deformed sample at nm level scale.

The challenge:

Solution:

Identify texture and structural defects at nm scale of olivine mineral applied at very high pressures (3 Gpa)

ASTAR technique coupled with precession electron diffraction





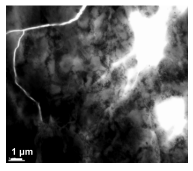
Experimental Data
TEM type: Jeol 3010
Map resolution: <10 nm
Scanned area: 5 x 5 µm

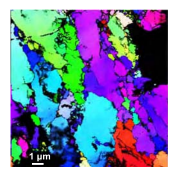
Czystal Structuze

Olivine (Forsterite) Mg<sub>2</sub>SiO<sub>4</sub>:

Orthorhombic, Pbnm

a= 4.76 Å, b=10.27 Å, c= 5.98 Å







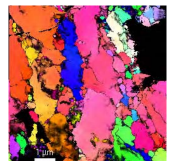


figure 1

ASTAR orientation olivine map with high number of defects (a) BF image (b) along x direction (c) Multi-Anvil Cell X-ray (X17 MAC) facility at the National Synchrotron Light Source Brookhaven USA (d) ASTAR orientation map along z direction

Textures of minerals at nm are intimately linked to their physical & chemical properties

### THE WORLD OF MINERAL POLYMORPHS & TEXTURES

Inorganic nanoparticles are widely used in catalysis. Research has led to control their morphology, crystallographic orientation and assembly. Metal oxide composites can be formed by heteroaggregation using electrostatic forces induced by the surface charges of the nanoparticles. It is important to understand the mechanisms of heteroaggregation and in particular the role of crystallographic facets.

ASTAR and TEM electron tomography has been carried out in order to analyse the

The challenge:

Identify orientation / phase of different mineral phases with particle size <50 nm

Solution:

ASTAR technique coupled with precession

electron diffraction

heteroaggregates homogeneity and their 3D structure in a case study of rods of goethite (α-FeO(OH)) with particles of brookite (TiO<sub>2</sub>) mixture.

ASTAR orientation / phase

map at nm scale revealed that the rods of goethite are assembled one to each other along their long edges. Particles of brookite are agglomerated and are linked to the tips of the goethite rods (fig.1). It is very important that exactly the same results have been revealed using TEM electron tomography. It is interesting to observe dramatic resolution in ASTAR phase maps generated with Cs corrected TEM in comparison with std Lab6 300 kV TEM (Jeol 3010) Fig.1a-b

Emphasis on the synthesis of nanoparticles is currently directed at designing their crystal structure and morphology and therewith their self-assembling behavior. We employed nonionic water-in-oil microemulsions as reaction medium to explore nanoscopic crystallization pathways for barium carbonate (witherite mineral).

Contrary to what was originally expected is the genesis of stable orthorhombic witherite in these media is not a straightforward process enabling thus the investigation of birth, deterioration and transformation of so far unprecedented metastable amorphous and crystalline nanoparticles. ASTAR gives the opportunity to identify the different polymorph phases in a crystalline nanoparticle ensemble (orientation/phase maps) and to determine major growth directions of individual nanoparticles or the adjacent faces of selfassembled nanoparticles (orientation maps) thus gaining deeper insight into the underlying processes. Fig. 2 shows monoclinic (P12,/m1 (11)) nanocubes together with orthorhombic (Pnma (62)) nanorods that have grown out in a direction.

zystal Structure

a= 9.17 Å, b= 5.44 Å, c= 5.13 Å (α-FeO(OH))Goethite: Orthorombic, Pbnm a= 4.59 Å, b= 9.95 Å, c= 3.02 Å

TiO, Brookite: Orthorhombic, Pbca

Whiterite BaCO<sub>3</sub>: Orthorhombic, Pnma a=5.31 Å, b= 8.9 Å, c= 6.43 Å

BaCO<sub>3</sub>: Monoclinic, P2<sub>1</sub>/m

a = 6,9 Å, b= 5,29 Å, c = 4,51 Å;  $\beta$ = 107,8°

Experimental Data TEM type: Libra 200 FE Cs(Fig.1c)

(A)

(B)

(C)

Bank\_[Brookite]\_100\_0.2

Bank\_['Goethite' Pnma]\_

(A) Brookite particles & Goethite

nanorods TEM bright field image

(C) ASTAR phase map with <nm resolution taken with aberration

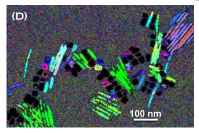
corrected Cs-TEM

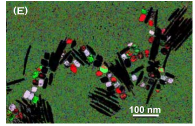
(B) ASTAR phase map of the same area

100 nm

Map resolution: <1 nm Scanned area: 2 x 2 µm

TEM type: Tecnai 20 F (Fig.2) Map resolution: 1 nm Scanned area: 800 x 600 nm





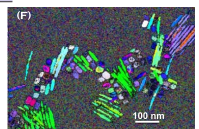


figure 2

ASTAR phase map of different whiterite polymorphs (blue-monoclinic, red ortho BaCO.) (b) colour code for orthorhombic structure (a) and for monoclinic (c) ASTAR orientation map for BaCO3 orthorhombic phase (y axis), (d) same for BaCO3 monoclinic phase (z axis) (e) ASTAR orientation map for both BaCO3 polymorphs (y axis)





### RESEARCH GROUPS & TECHNICAL DETAILS

#### **DESIGN MATERIALS WITH SPECIFIC APPLICATIONS**

Technical details: ASTAR - Jeol 2100F; precession angle 0.7°

Research Group: Dr. Dong Qui, Univ. of Queensland, Australia; Dr. P. Cizek, Dr. A. Sullivan, Deakin Univ., Australia

#### BETTER STEEL WITH IMPROVED PROPERTIES

Technical details: CM 200 - ASTAR; precession angle 0.5°

Research Group: Dr. A. Gholinia, Dr. S. Height, Univ. of Manchester, UK

#### CHALLENGING MATERIALS FOR CHALLENGING APPLICATIONS

Technical details: Tecnai F20 - ASTAR; precession angle 0.5°

Research Group: Dr. S. Lartigue, CNRS, Paris, France; Dr. M. Veron, INP Grenoble, France

#### STEEL CORROSION IN NUCLEAR PLANTS

**Technical details:** Tecnai F20 - ASTAR; precession angle 0.6° **Research Group:** Dr. F. Delabrouille, EDF Paris, France

### STRUCTURE OF EMBEDDED NANODOMAINS IN INTERMETALLICS

Technical details: Rotational tomographic holder; beam precession (1.2°) by DigitStar, NanoMEGAS.

Research Group: Prof. U.Kolb, Dr. E.Mugnaoli, Johannes Gutenberg-Universität, Mainz, Germany; Prof. N.Frangis, Dr. E.Sarakinou, Univ of Thessaloniki, Greece

#### METALS UNDER HUGE STRESS

Technical details: Tecnai F20 - ASTAR

Research Group: Dr. C. Kuebel, D. Wang, KIT Karlsruhe, Germany

#### PAVEMENTS OF THE NANOWORLD

Technical details: ASTAR - Jeol 3010; precession angle 0.8°

Research Group: Dr. P. Cizek, Dr. A. Shelivan, Deakin Univ., Australia,

#### **UNDERSTANDING LI INSERTION FOR NEXT GENERATION BATTERIES**

Technical details: Jeol 2200 FF - ASTAR; precession angle 1.2°

Research Group: Dr. G. Brunetti, Dr. J. L. Rouviere, CEA-LETI Grenoble, France

#### NANOSCALE REACTION MODELS IN SOLID OXIDE FUEL CELLS

Technical details: Jeol 2200 FS - ASTAR; precession angle 0.7°

Research Group: Dr. M. Cantoni, Dr. A. Duncan, EPFL Lausanne, Switzerland, Dr. M. Veron, INP Grenoble, France

### ATOMIC SCALE OF NANOLAYER COMPOSITES

Technical details: Jeol 2100F - ASTAR; precession angle 0.6°

Research Group: Dr. P. Cizek, Deakin Univ., Australia; Dr. Bhattacharyya, ANSTO, Australia; Dr. M. Veron, INP Grenoble, France

#### **NEXT GENERATION CIRCUITS FOR SEMICONDUCTOR INDUSTRY**

Technical details: Jeol 2100F - ASTAR; precession angle 0.7°

Research Group: Prof. P. Ferreira , Dr. A. Darbal , Dr. J. Ganesh , Dr. J. Kameswaran , Dr. S. Rajashekara, Univ. of Texas at Austin, Prof. K. Barmak , Dr. G. S. Rohrer, Carnegie Mellon University

#### STRUCTURE OF MAGNETIC SEMICONDUCTORS

Technical details: Jeol 2200 FS - ASTAR; precession angle 0.5°

Research Group: Dr. H. Kirmse, Dr. I Hausler, Humboldt Univ., Berlin, Germany

#### HYBRID INORGANIC / ORGANIC SYSTEMS FOR OPTOELECTRONICS IN DAILY LIFE

Technical details: Jeol 2200 FS - ASTAR; precession angle 0.5°

Research Group: Dr. H. Kirmse. Dr. I Hausler. Humboldt Univ.. Berlin. Germany

### **DRUG DELIVERY & STRUCTURE OF NANOPARTICLES**

Technical details: Tecnai 20 FS - ASTAR; Tecnai F20 - ASTAR; Jeol 2100F - ASTAR; precession angle 0.7°

Research Group: Dr. Hui Hu, Univ. of Singapore; Dr. S. Rouvimov, Dr. P. Moeck, Portland State Univ. USA, Dr. P. Ferreira, Texas Univ. at Austin, USA

#### INSIDE THE NANOWIRE FOREST

Technical details: Zeiss Libra 120 – ASTAR

Research group: Dr. M. Gemmi, Italian Institute of Technology Pisa, Dr. I. Sorba CNR Nano, Pisa, Dr. D. Ercolani Scuola Normale Superiore Pisa, Italy

#### **TEXTURE SECRETS OF NANOPARTICLES**

Technical details: Zeiss Libra 120 – ASTAR /

Research Group:Dr. M. Gemmi (CNI@NEST, Istituto Italiano di Tecnologia, Pisa, Italy), V. Voliani (NEST, Scuola Normale Superiore, Pisa, Italy)

### CRYSTAL ORIENTATION IMAGING OF ORGANIC NANOMAGNETS

Technical details: Jeol 3010-ASTAR

Research group: Dr. S.Pillet Univ of Nancy, Dr. E.Rauch, M.Veron INP Grenoble, France

### INSIDE A ROMAN CERAMIC FACTORY IN ANCIENT GAUL

Technical details: CM 20 FEG-ASTAR

Research group: Dr. P.Sciau CEMES Toulouse, France

### SHED LIGHT TO EARTH SEISMICITY

Technical details: Jeol 3010- ASTAR Precession angle  $0.8^{\circ}$ 

Research group: Dr. Bollinger, Prof. P. Cordier, Univ of Lille, Dr. E. Rauch, M. Veron, INP Grenoble, France

#### THE WORLD OF MINERAL POLYMORPHS & TEXTURES

**Technical details:** Libra 200 FE Cs – ASTAR & Techni 20F –ASTAR Precession angle  $1^{\circ}$ 

Research group: Dr. E. Rauch, INP Grenoble, France, Dr. T. Weirich, Dr. Wiebke, Sager RWTH Univ of Aachen, Germany & Prof. M. Veron INP Grenoble, France



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