

theory is used to explain the observed temperature dependence.

MM 42: Nanostructured Materials III

Time: Thursday 14:00–15:45

Location: IFW A

MM 42.1 Thu 14:00 IFW A

Virtual diffraction of MD-simulated nanocrystalline Pd under compression — ●JÜRGEN MARKMANN^{1,2}, DMITRIY BACHURIN³, PETER GUMBSCH³, and JÖRG WEISSMÜLLER^{1,2} — ¹Universität des Saarlandes, FR7.3 Technische Physik, 66123 Saarbrücken, Germany — ²Forschungszentrum Karlsruhe in der Helmholtz-Gemeinschaft, Institut für Nanotechnologie, 76344 Eggenstein-Leopoldshafen, Germany — ³Universität Karlsruhe, Institut für Zuverlässigkeit von Bauteilen und Systemen, 76131 Karlsruhe, Germany

Wide angle x-ray scattering (WAXS) is a widely used tool to characterize the microstructure of crystalline materials. During the last few years the means were developed to calculate x-ray spectra out of molecular dynamics simulation data and to use these spectra to analyse the MD structures. This approach has been validated to yield reliable results for well defined structures. Now, more realistic samples were investigated and analysed in-situ during a deformation by uniaxial compression. These samples consist of 100 randomly oriented palladium grains and were analysed for microstructural parameters like lattice constant, grain size, microstrain, and stacking fault density which can be evaluated out of the position and shape of the x-ray reflections. The most interesting finding is the increase of stacking fault density with the onset of plastic deformation. The stacking fault density reduces to zero when the structure is unloaded. This will be discussed together with the other microstructural parameters in terms of nucleation, dissolution, and movement of dislocations.

MM 42.2 Thu 14:15 IFW A

Segregation-induced near-surface lattice strain in CuAu nanoparticles — ●DARIUS POHL¹, ELIAS MOHN¹, JURI BARTHEL², KARSTEN ALBE³, LUDWIG SCHULTZ¹, and BERND RELLINGHAUS¹ — ¹IFW Dresden, D-01171 Dresden, Germany — ²FZ Jülich, D-52425 Jülich, Germany — ³TU Darmstadt, D-64287 Darmstadt, Germany

For binary nanoparticles, experimental findings indicate the segregation of one alloy constituent towards the particle surface which leads to a surface-near relaxation of the lattice. Due to a difference in the surface free energies between copper and gold, a gold segregation for nanoparticles is expected. In the case of CuAu nanoparticles, Au segregation towards the surface is suggested to lead to a lattice expansion. Through aberration-corrected HRTEM imaging with minimum delocalisation is achieved, and the position of individual atom columns can be determined with unrivalled precision.

Molecular dynamic (MD) simulations of CuAu nanoparticles are conducted under the constraint that the particle surface is terminated with either Cu or Au. A comparison between HRTEM contrast simulations of the simulated structures and experimental HRTEM images then allows to distinguish between the different possible scenarios. It is hereby proven that single crystal CuAu nanoparticles with a mean diameter of 3 nm exhibit a lattice relaxation due to a segregation of Gold atoms. This finding is in agreement with previous results obtained for multiply twinned FePt nanoparticles and thus corroborates that the surface-near expansion of the lattice is due to a segregation rather than to internal stress within the particle structure itself.

MM 42.3 Thu 14:30 IFW A

Size-dependence melting transformation of nanoparticles confined in an Al-rich glass — ●NANCY BOUCHARAT¹, HARALD RÖSNER², and GERHARD WILDE² — ¹Forschungszentrum Karlsruhe, Institut für Nanotechnologie, Karlsruhe — ²Universität Münster, Institut für Materialphysik, Münster

A common way to study the size dependence of the melting transformation is given by incorporating nanoparticles within a matrix. This configuration implies additional contributions related to the nature of the interfaces between the particles and the host matrix which strongly affect the melting process. Thus, the incorporation of nanoparticles within a glassy matrix opens a new angle to revise the size-dependence of melting by limiting the constraints imposed on the particles by the presence of a crystalline matrix with anisotropic lattice mismatch. Here, AlYFe glasses containing immiscible Pb or In additions have

been synthesized via rapid solidification. As expected for liquid-liquid phase separation, the Pb-containing samples consist of spherical Pb inclusions homogeneously dispersed within the matrix. In contrast, the In-containing samples show non-crystalline In-enriched regions. Applying a subsequent low-temperature treatment involves the crystallization of spherical particles with sizes smaller than 5 nm, which melt at extremely low temperatures compared to the bulk material. In this context, the melting behavior is discussed with respect to the size dependence and to the energetic contributions from the particle/matrix interfaces.

MM 42.4 Thu 14:45 IFW A

Gold nanoparticles under synchrotron X-rays — ●CHANG-HAI WANG¹, CHI-JEN LIU², TZU-EN HUA², CHIA-CHI CHIEN², WEI-HUA LENG², SHIN-TAI CHEN², CHENG-LIANG WANG², and YEU-KUANG HWU² — ¹Institute of Inorganic and Analytical Chemistry, Johannes Gutenberg - University, 55099 Mainz — ²Institute of Physics, Academia Sinica, Taipei, 11529 Taiwan

Gold particles with length scale less than 100 nm develop unique physical properties and biocompatible features that render extensive applications in nanotechnology. To materialize the functionalities, gold nanoparticles with controlled size, favorable surface properties and size distribution would be a pre-requisite. This work describes a new room-temperature synchrotron X-ray irradiation method to prepare reductant- and stabilizer-free colloidal gold solutions. Typical characterization tools include TEM, UV-VIS, FTIR, XRD and ICP-OES. The influence of processing parameters such as the pH value, exposure time, ionic strength and radical scavenger on the structure of gold nanoparticles was investigated. The mechanisms underlying the X-ray-triggered reduction of gold ions and the formation of gold clusters are discussed in detail. An interesting morphological evolution as a function of exposure time, from cross-linked network-like structure to individual particles, has been discovered. This approach could be easily extended to the preparation of polymer-modified colloidal gold by simply adding the polymer species to the precursor solutions. As an implication for nanotechnology, the interactions between gold nanoparticles and cells are also studied and reported.

MM 42.5 Thu 15:00 IFW A

Temperature dependent vibrational fingerprints of gold nano clusters: a DFT study — ●LUCA GHIRINGHELLI and MATTHIAS SCHEFFLER — Fritz-Haber-Institut der Max-Planck-Gesellschaft, Faradayweg 4-6, 14195 Berlin, Germany

The study of gold nanoclusters is a flourishing topic, due to its importance for catalysis and the presence of unexpected phenomena (e.g. chirality). Here, we apply the recently developed all-electron ab-initio code “FHI-aims” for the density functional (DFT) study of the relative energies and vibrational properties of known isomers of small Au_n (n ≤ 10) clusters. In particular, besides the calculations of the traditional harmonic frequencies, we focus on the less common evaluation of the (temperature dependent) vibrational spectrum, via Fourier transform of the velocity autocorrelation function, where the velocities come from DFT based Molecular Dynamics trajectories at given temperatures. We underline differences between harmonic and an-harmonic spectra at different temperatures and compare with available experimental data.

MM 42.6 Thu 15:15 IFW A

Delocalization mechanisms of excess free-volume in nanoglasses — ●DANIEL SÖPU¹, KARSTEN ALBE¹, YVONNE RITTER¹, and HERBERT GLEITER² — ¹Institut für Materialwissenschaft, TU Darmstadt, Petersenstr. 23, D-64287 Darmstadt, Germany — ²Institute of Nanotechnology, Karlsruhe Institute of Technology, P.O. Box 3640, D-76021 Karlsruhe, Germany

Molecular dynamics simulations are presented which provide a detailed picture of the structure of nanoglasses, a class of material which can be synthesized by consolidating nanometer-sized glassy particles. Our results provide evidence the existence of glass-glass interfaces between

the consolidated particles. These interfaces are characterized by an excess free volume. By comparing simulations for covalently bonded Ge nanoglass and metallic CuZr nanoglass, we find that the delocalization of this free volume is driven by homogeneous plastic flow and by thermally activated diffusion. These results suggest that the overall density, microstructure and atomic structure of nanoglasses can be adjusted by the initial particle size and chemical composition as well as by the annealing conditions.

MM 42.7 Thu 15:30 IFW A

Mechanical alloying of Fe-Cu powders: elaborating the microstructure at various scales — ●CATHARINA WILLE¹, TALÁAT AL-KASSAB^{1,2}, PYUCK-PA CHOI³, and REINER KIRCHHEIM¹ — ¹Institut für Materialphysik, Universität Göttingen, Germany — ²Material Science and Engineering, King Abdullah University of Science and Technology (KAUST), effective 15 April 2009 — ³Korea Institute of Science and Technology, Nano-Materials Research Center

In this contribution, the process of mechanical alloying has been stud-

ied for both sides of the binary Fe-Cu system. Ranging from light-optical microscopy on the mm-scale to atom probe tomography (APT) on the Ångström-scale, the morphology could be observed on several orders of magnitude. Since a ductile element (Cu, fcc) and a brittle one (Fe, bcc) were combined, striking differences in morphology were expected and found on all length-scales, depending on the mixing ratio. Results on powders with low concentrations of the respective minority component will be presented and discussed.

Regarding the widespread application and accessibility, Fe-Cu acts as an ideal binary model alloy to elaborate the enforced non-equilibrium enhanced solubility being immiscible and characterised by a large positive heat of mixing. Chemical identification on the Ångström-scale was granted by APT. Thus, not only the atomic mixing of Fe and Cu could be evaluated, but also the distribution of impurities, mostly stemming from the fabrication procedure.

Financial support from the DFG under contract KI-230/33-1 is gratefully acknowledged.

MM 43: Liquid and Amorphous Metals II

Time: Thursday 16:00–17:45

Location: IFW A

MM 43.1 Thu 16:00 IFW A

serrated flow behavior in bulk metallic glasses — ●GANG WANG, NORBERT MATTERN, and JÜRGEN ECKERT — Institute of complex materials, IFW-Dresden, 01069 Dresden, Germany

Plastic deformation of bulk metallic glasses (BMGs) is a complex inhomogeneous process characterized by avalanches in the motion of shear bands. In the present study, based on the CuZr-based BMG, we investigate the serrated flow behavior in the plastic deformation stage. We statistically analyze the serrated stress-strain behavior of BMGs with different plastic deformation ability. The cumulative possibility distribution of the elastic energy density follows the Weibull distribution. Accompanying with the compression tests, we measured the atomic-scale strain during the serrated flow stage by x-ray synchrotron radiation at room temperature. High resolution strain scanning reveals the relationship between the macroscopic serrated flow and the atomic-scale elasto-plastic deformation. Based on the potential energy landscape (PEL) theory of ductile glasses, we attempt to construct a clear physic image of the origination of the plastic deformation in glassy phase, i.e. the evolution from the disorder atomic cluster, to shear transforming zone (STZ) formation, to shear bands origination and then to the shear slip occurring.

MM 43.2 Thu 16:15 IFW A

Dynamics of shear localization and stress relaxation in amorphous Cu₅₀Ti₅₀ — ●MAX NEUDECKER and S. G. MAYR — I. Physikalisches Institut, Georg-August-Universität Göttingen, Friedrich-Hund-Platz 1, 37077 Göttingen

Dynamic heterogeneities of atomic mobility in metallic glasses are investigated for the model glass Cu₅₀Ti₅₀ with the help of classical molecular dynamics computer simulations. By rapid quenching from melt at various cooling rates (comprising 5 orders of magnitude), differently relaxed amorphous cells are prepared. During subsequent shearing, we observe a series of highly localized shear events, termed shear transformation zones (STZs). Detailed analysis focuses on geometrical shape and size of STZs, mechanical stress dynamics and correlations of mobility with local properties. We identify a local stress bias as physical origin of STZ formation during low temperature deformation. Further investigations concern the characterization and analysis of heterogeneous mobility and stress dynamics during shearing at higher temperatures.

[1] M. Neudecker and S. G. Mayr, *Acta Materialia*, in press (2008) *Financially supported by the DFG (PAK 36)*

MM 43.3 Thu 16:30 IFW A

Temperature and kinetics of shear band propagation in amorphous metals — ●FLORIAN H. DALLA TORRE, DAVID KLAUMÜNZER, ALBAN DUBACH, and JÖRG F. LÖFFLER — Laboratory of Metal Physics and Technology, Department of Materials, ETH Zurich, Wolfgang-Pauli-Str. 10, 8093 Zurich, Switzerland

Deformation in bulk metallic glasses (BMGs) far below their glass transition temperature is known to be spatially inhomogeneous and

restricted to narrow nanometer-sized shear bands. In this study the shear band characteristics of Zr-based BMGs has been investigated as function of strain rate and temperature. To capture discrete shear events precisely a dedicated acoustic emission spectrometer has been used in-situ during deformation testing, which allows for extremely high data acquisition rates of the order of MHz. Preliminary results of evaluating the viscosity within shear bands during deformation suggest that the local temperatures are of the same order of magnitude as those measured during homogeneous deformation close to the glass transition temperature. Acoustic emission results indicate that irreversible deformation starts well before reaching the elastic limit and that the Kaiser effect known to crystalline metals is present also in amorphous metals. It is shown that serrated flow and the associated stress drops are a function of temperature and strain rate. The time periods for a shear band to be activated increases with decreasing temperature from milliseconds at ambient temperatures to seconds at temperatures of 200 K.

MM 43.4 Thu 16:45 IFW A

Modelling the mechanical properties of bulk metallic glasses with nanoscale precipitates — ●YVONNE RITTER and KARSTEN ALBE — Institut f. Materialwissenschaft, TU Darmstadt, Petersenstr. 23, D-64287 Darmstadt

Bulk metallic glasses exhibit unique mechanical properties as compared to conventional crystalline metals. Large elastic strains and specific strengths exceeding the strength of crystalline alloys by more than 100% illustrate the high potential for structural applications. On the other hand, metallic glasses have a relatively low tensile ductility and fail catastrophically when reaching the flow stress. Nanoscale precipitates in the glassy matrix were found to improve the plasticity of BMGs. The underlying mechanisms, however, that lead to the improved deformation behavior have not yet been exposed. In this study, we investigate the role of structural inhomogeneities in CuZr by means of molecular dynamics simulations. Nanoprecipitates of crystalline Cu and CuZr as well as amorphous precipitates with varying composition are investigated and their influence on the materials behavior under tensile load and shear is investigated.

MM 43.5 Thu 17:00 IFW A

Deformation-induced martensitic transformation in Cu-Zr-Al(Ti) bulk metallic glass composites — ●RAM BACHCHAN KUMAR, SIMON PAULY, JAYANTA DAS, and JÜRGEN ECKERT — Institut für Komplexe Materialien, IFW Dresden

Plastic deformation of Cu-Zr-(Al, Ti) bulk metallic glass (BMG) composites induces a martensitic phase transformation from the B2 to the B19* CuZr phase. Addition of Ti to binary Cu-Zr increases the temperature above which the B2 CuZr phase becomes stable. This affects the phase formation upon quenching in Cu-Zr-Ti BMG composites. The deformation-induced martensitic transformation is believed to cause the strong work hardening and to contribute to the large compressive deformability with plastic strains up to 15%.