this phenomenon has been a long-lasting enigma in the interpretation of 
geological signatures of the Earth’s mantle and the geochemistry of 
core-mantle differentiation. This inconsistency has been explained 
either by a differentiation model with a chemical equilibrium at the 
hottom of a silicate magma ocean while a metallic core forms at a depth 
of 1100 km [2] or by the so called ‘late veneer’ hypothesis. The later 
postulates an enrichment of HSE in the Earth’s mantle by an intensive 
meteorite bombardment 1000-50 My after the Earth’s accretion [3]. 
So far, all the metal-silicate partitioning studies make use of ‘classical’ 
HP-HT techniques, e.g., multi-anvil press, and therefore are limited to 
PT conditions of the Earth’s mantle (max. 15-20 GPa/2200°C). There is 
urgent needs for experiments at much higher pressures and temperatures 

Keywords: in situ, partitioning, siderophile elements

MS.76.1

Structure and electronic properties of free, supported and protected clusters

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Clusters can be viewed as small representations of the crystals 
formed by metal. Due to finite size effects caused by their special 
electronic and/or ionic configurations, the properties of clusters can be 
substantially different from the bulk however. One example is the well 
known catalytic activity of small gold particles in sharp contrast to the 
inertness of bulk gold.

In our work, we explore the world of small metal clusters with the 
help of density functional theory. In particular small gold clusters are 
governed by strong electronic confinement effects that can be understood 
in a simple jellium picture. The corresponding shell closings give a 
unified explanation of the magic stability in ligand protect clusters [1]. 
This includes an alternative model for the well known Schmid “Au54g” 
cluster different from the models used in the literature. Electronic 
effects explain STM signatures of supported clusters and even give the 
possibility to determine the cluster’s charge directly (see the figure) [2]. 
The work is extended to mixed clusters and clusters of other metals.

Figure: Experimental and simulated STM signatures for MgO supported gold 
clusters [2].

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Keywords: cluster, gold, superatom

MS.76.2


Pushing all-electron DFT past old limits for structure of surfaces and 
molecules

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The prediction and our understanding of structural and chemical 
processes of matter benefit tremendously from the strength of today’s 
first-principles total-energy methods. The obvious and ongoing 
limitations of these methods are:
- the affordability and accuracy of the physical approximations [in 
density functional theory (DFT), primarily the exchange-correlation 
functional]
- but in addition, the affordability and accuracy of all other numerical 
approximations that lead to a total energy for a given structure.

For few-atom structures, serial structure screening, exhaustive 
structure predictions and meaningful time scales in molecular dynamics 
are now routinely possible, but the challenge still grows immensely for 
larger system sizes. The present work describes some recent efforts 
to push out these limits further, using all-electron density functional 
theory as implemented in the numeric atom-centered orbital based 
code package FHI-aims [1]. We show how accurately the classic large-
scale surface reconstruction of Au(100) and Pt(100) are captured in 
terms of structure and energetics by current all-electron DFT - if large 

enough structure approximants (up to ~1000 atoms in a slab model) 
are considered [2]. We then comment on the additional challenges 
encountered for the structure and dynamics of biochemical molecules 
at the other end of the periodic table, addressing both the structure 
and high-temperature stability of a series of polyalanine-based peptide 
molecules (up to 180 atoms) in the gas phase. In particular, we find 
that van der Waals interactions are of unexpectedly large relevance to 
stabilize specifically the “classic” α-helical structure over competing 
conformations (even in the gas phase) [3]. Our results are directly 
verified by comparing theoretical infrared spectra from basis-set
converged molecular dynamics to room-temperature experimental data [4].


Keywords: surface, polypeptide, theory

MS.76.3

Optimization of materials with time dependent density functional theory

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The theory of Quantum Optimal Control (QOCT) aims to the theoretical design of external fields capable of controlling quantum mechanical systems. Nowadays, experimentally, the so-called pulse shaping techniques can be used to design pulses that produce a desired effect on atoms or molecules. These may react in complex manners when they are irradiated with high-intensity electromagnetic pulses: multi-photon, tunneling and over-the-barrier ionization, laser driven photo-induced isomerizations or fragmentations, and high harmonic generation are some of the non-linear effects that are observed.

We wish to investigate the control of these complex phenomena from first principles, by making use of time-dependent density-functional theory (TDDFT) to describe the electrons, a real-space numerical representation, and various optimization techniques.

In this talk, I will present this marriage of the QOCT techniques with TDDFT, along with various application examples.

Keywords: control, laser, theory

MS.76.4

Density functional calculations of electron energy loss data and inelastic mean free paths in elemental and binary materials

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Density functional theory is used to determine joint density of states and optical energy loss data for elemental metals and compounds. A novel transform algorithm is then employed to produce electron energy loss spectra and inelastic mean free paths. These results are compared with recent high profile experimental measurements in the low-energy region of strong plasmon resonances.

We present calculations of electron energy loss spectra and inelastic mean free paths determined using transformed optical loss data from density functional theory. Our novel approach enables investigation of plasmon excitation and electron scattering without the use of empirical fitting parameters or experimental data - common requirements of previous theory [1]. We consider elemental metals copper, molybdenum, zinc and selenium, and the binary semiconductor zinc selenide. The technique is extensible to any non-crystalline or crystalline uniform material.

Results are compared with recent high precision measurements of the inelastic mean free path [2, 3] derived from x-ray absorption experiments [4, 5]. Surprising outcomes of these investigations include accurate dynamic bond lengths [6] and complementary approaches to single crystal structure [7,8]. Broadening mechanisms in the momentum-dependent energy loss function, and the energy evolution of plasmon peaks are investigated as potential sources of error in established theory. We consider the effects of crystal structure on the energy loss spectrum, and critically investigate the correlation between inelastic scattering in polyatomic systems and their corresponding elemental systems. This work focuses on electron energies below 100 eV – the region of strong plasmon resonance – where published data is highly uncertain and incomplete.

Keywords: density functional theory, IMFP, photoelectron

MS.76.5

Topological analysis of charge densities obtained from experiment and plane-wave calculations

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Accurate electron density distribution functions (ρ(r)) for crystals can be obtained from high resolution experiment and calculated by means of density functional (DFT) methods. To recover the experimental electron density one should carry out multipole refinement procedure on the dataset collected with high redundancy at low temperature. Such an “experimental” electron density function usually is suffer from systematic errors caused by thermal motion or/and absorption effects etc. Thus, DFT calculations should be “method of choice” to obtain very accurate ρ(r) function in the case of molecular crystals. Indeed, it does not carry out long-time experiments and multipole refinement. Unfortunately, the convergence between experimental and optimized atomic coordinates and cell parameters is still not ideal for the most of the modern DFT functionals. Besides, all-electron treatment of molecular crystals can be achieved only for small cells which are not contain heavy atoms.

The most abundant type of DFT calculations of crystals is based on the usage of plane wave basis set and pseudopotentials. Pseudopotentials are usually considered as very bad choice to obtain the ρ(r) function suitable for the topological analysis in terms of R. Bader’s “Atoms in molecules” theory. In many cases the critical points (CP) (3,-1) could not be localized. To overcome this problem the pseudopotentials with small radius of core region (r_c) (hard pseudopotentials) should be chosen.

The suitable pseudopotentials can be generated by hand, but they are also can be obtained with quantum chemical programs. Our choice is the VASP code which is supplied with rich library of hard non-local