Helium droplet assisted synthesis of plasmonic Ag@ZnO core@shell nanoparticles

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ABSTRACT
Plasmonic Ag@ZnO core@shell nanoparticles are formed by synthesis inside helium droplets with subsequent deposition and controlled oxidation. The particle size and shape can be controlled from spherical sub-10 nm particles to larger elongated structures. An advantage of the method is the complete absence of solvents, precursors, and other chemical agents. The obtained particle morphology and elemental composition have been analyzed by scanning transmission electron microscopy (STEM) and energy dispersive X-ray spectroscopy (EDS). The results reveal that the produced particles form a closed and homogeneous ZnO layer around a 2–3 nm Ag core with a uniform thickness of (1.33 ± 0.15) nm and (1.63 ± 0.31) nm for spherical and wire-like particles, respectively. The results are supported by ultraviolet photoelectron spectroscopy (UPS), which indicates a fully oxidized shell layer for the particles studied by STEM. The plasmonic properties of the produced spherical Ag@ZnO core@shell particles are investigated by two-photon photoelectron (2PPE) spectroscopy. Upon excitation of the localized surface plasmon resonance in Ag at around 3 eV, plasmonic enhancement leads to the liberation of electrons with high kinetic energy. This is observed for both Ag and Ag@ZnO particles, showing that even if a Ag cluster is covered by the ZnO layer, a plasmonic enhancement can be observed by photoelectron spectroscopy.

KEYWORDS
nanoparticle, helium droplet, plasmonics, photoelectrons

1 Introduction
Zinc oxide has become one of the most popular materials for applications in the fields of photocatalysis and optoelectronics due to its intriguing properties. ZnO is a non toxic n-type semiconductor with a wide direct band gap of 3.37 eV, very similar to TiO2 but with a higher absorption efficiency under solar irradiation and a large exciton binding energy of about 60 meV [1]. As a photocatalyst it shows high potential in degradation processes of organic pollutants via the generation of reactive oxygen species and also exhibits antibacterial properties, especially when scaled down to the nano regime [2]. Due to its photoconductivity and photoluminescence characteristics, applications of ZnO in optoelectronics are possible, for example, in ultraviolet (UV) detectors, solar cells, lasers, and various sensing devices [3, 4]. In recent years, miniaturization and engineering of material properties on the nanometer scale have strongly influenced the research on ZnO. Starting with needles and other structures in the micrometer size regime [5], by now the synthesis of ZnO nanoparticles is an important branch of research. An increasing amount of one-, two-, and three-dimensional structures have been synthesized in shapes ranging from nanoparticles, -rods, and -ribbons over nanoplates to exotic flower or snowflake structures [6]. The formation of ZnO-based composite materials opens up further possibilities for the tuning of properties, which permit the addition of new functionalities and, consequently, the doping of ZnO receives increasing attention. Different material groups can be used as dopants, for example, to produce p-type ZnO, enable applications in spintronics, or to improve optical and electrical properties [7]. A modification with Ag can significantly enhance the photocatalytic activity of ZnO [8] because the electron-hole pair concentration is increased by the strong local fields induced by the excitation of the localized surface plasmon in Ag [9].
Most of the synthesis routes for pure or doped nanosized ZnO have in common that they are based on multi-step chemical processes involving solvents or precursors. Though those techniques are well controlled and documented, it is possible that residues from chemical agents or surfactants remain at the particle surface or even at the core–shell interface.
In this study, the helium nanodroplet synthesis approach is employed as a pristine and facile technique for the synthesis of Ag@ZnO core@shell nanoparticles. This process bears the advantage of dispensing no chemicals, additives, or other additional agents besides the high purity metals, which are doped into the superfluid helium droplets where they agglomerate to nanoparticles. Since the whole procedure is conducted under ultra-high vacuum (UHV) conditions, a contamination of the
sample and other influences from environmental sources are excluded. Due to the dependence of the particle size and shape solely on a few well defined and accurately adjustable experimental parameters the approach allows for a high reproducibility. The range of accessible shapes spans from smaller spherical to longer wire-like nanoparticles, as shown in the course of this study. The obtained particle size and shape depends on the helium droplet source conditions, with which the helium droplet size can be controlled [10]. Small helium droplets lead to the formation of spherical nanoparticles, typically with a diameter below 5 nm. As a consequence of their superfluidity, large droplets host quantum vortices [11], along which dopants preferentially agglomerate enabling the formation of wire-like nanoparticles. Even though the scalability of the production is limited because only a few 100 ng per hour can be produced, the method allows for an investigation of fundamental particle properties. Furthermore, for future experiments the synthesis of core@shell species with a wide range of material combinations is accessible with one experimental setup simply by exchanging the dopant metal sources. Electron microscopy has been used to probe the size distribution, morphology, and composition of the deposited nanoparticles, revealing that the Ag cores are covered by a uniformly shaped wurtzite ZnO shell. Further investigations on the oxidation state of the ZnO shell have been conducted by ultraviolet photoelectron spectroscopy and two-photon photoelectron spectroscopy was employed to demonstrate the plasmonic activity of the ZnO covered Ag core.

2 Experimental

2.1 Nanoparticle synthesis

The ZnO and Ag@ZnO core@shell nanoparticles investigated in this work were produced utilizing the helium nanodroplet approach. This technique employs superfluid helium droplets (HeN) as an inert environment for the synthesis of nanosized metallic particles [12–14]. Details about the experimental design of the HeN apparatus can be found in Ref. [15]. In summary, a closed cycle refrigeration unit (Sumitomo RDK-408D2) is used to cool gaseous helium (99.9999% purity) at a stagnation pressure of 20–60 bar down to only a few Kelvin. The helium is then adiabatically expanded into high vacuum (~10^−6 mbar) through a 5 μm diameter nozzle resulting in a beam of superfluid HeN. The internal temperature of the HeN, thereby, drops to 0.37 K [16, 17]. The droplet sizes can be controlled by the temperature and the stagnation pressure of the helium gas. For this work a setup with a pressure of 20 bar at 8 K was used for the synthesis of spherical nanoparticles resulting in a He droplet diameter of about 70 nm with a mean number of about 5 × 10^10 He atoms [10]. For the larger wire-like particles, the parameters were changed to 20 bar and 5 K leading to HeN diameters of around 1 μm, consisting of about 2 × 10^10 He atoms [10]. Subsequently, the collimated HeN beam passes through another chamber (~10^−7 mbar) where resistively heated tungsten baskets coated with aluminium oxide are used as pickup cells for the evaporation of high purity metals. While passing through the metal vapor, the HeN are doped with the evaporated atoms. The superfluidity of the HeN matrix enables an unobstructed and frictionless roaming of dopant atoms. Upon statistical collisions within the HeN, the dopants accumulate and start to form clusters [18, 19]. Initially, these clusters are grown as multicenters inside a droplet before they further agglomerate to larger structures. The final size of the nanoparticles depends on the doping rate and on the initial HeN size, since both affect the average time between collisions and, thereby, the growth rate [20]. The synthesis of wire-like nanoparticles is enabled by the presence of vortices in larger He droplets with diameters of several hundred nanometers. These vortices form due to the angular momentum acquired by the superfluid droplets during the expansion of the helium through the nozzle into vacuum [11, 21, 22]. The dopant metal atoms and grown clusters inside the droplets can agglomerate along such vortices due to a pressure gradient directed towards the vortex core [23]. By adjusting the size of the droplets accordingly and by doping enough material into them, elongated wire-like structures can be formed along the vortices [20, 24, 25]. As stated above for the two employed parameter settings in this experiment, a change in the helium source conditions directly influences the size of the HeN and allows for a facile switching between the different nanoparticle types. It should be noted that by decreasing the source temperature not only the droplet size is increased, also the total HeN flux and the deposition rate are enhanced. During agglomeration of atoms and clusters, the released binding energy is dissipated by the evaporation of He atoms from the droplet. The attenuation of the HeN beam due to the decrease in droplet size caused by this process is measured in order to monitor the particle growth process [20]. The synthesis of core@shell particles is enabled by the subsequent doping of two different metals in consecutive pickup cells [26, 27]. Before entering the second pickup cell holding the shell material, the particle cores have already formed with the metal provided in the first cell. The metal doped to the droplets in the second pickup cell forms a shell layer around the core [28]. The dopant vapor pressure in each cell and, thereby, the metal atom doping rate and the nanoparticle growth are directly controlled by the temperature of the resistively heated pickup cells. Since the pickup cells are separately controlled, the amount of doped core and shell material can be adjusted independently. This allows for a tailoring of the core diameter and the shell thickness of the synthesized nanoparticles within a certain range, as has been shown for Ag@Au [29] and Fe@Au [30]. For this work, the pickup cells were loaded with Ag and Zn, enabling the formation of pure Ag, Zn and Ag@Zn core@shell particles inside the helium droplets.

After leaving the pickup region, the HeN beam reaches a third chamber at UHV conditions (~5 × 10^−10 mbar), where the deposition rate is monitored by termination of the beam on a quartz crystal microbalance. For deposition of the nanoparticles the substrates are inserted into the beam path with a wobblestick transfer system coupled to a vacuum suitcase (Ferrovac GmbH, NexGeneration UHV Suitcase). The suitcase allows for the storage and transport of the samples at UHV conditions (~1 × 10^−10 mbar). Due to the different employed analysis methods, support grids for scanning transmission electron microscopy (STEM) (Ted Pella Inc., Ultra-thin Carbon Film on Lacey Carbon Support Film, 300 mesh, Au) as well as glass coverlaws coated with indium tin oxide (ITO) (spi Supplies, 30–60 Ω, 18 × 18 mm, No. 1) were used as substrates. As the HeN beam is terminated at the substrate, the particles are cushioned by the surrounding helium and, therefore, keep their shape upon deposition under soft landing conditions [31, 32] as observed by atomic resolution electron tomography [33]. To start the oxidation of the Zn within the nanoparticles, the vacuum chamber with all the decorated substrates inside, was flushed with pure oxygen up to a pressure of 1 × 10^−6 mbar for 1 h. A single carbon substrate has been decorated with both fabricated species, i.e. the spherical and wire-like nanoparticles, for the STEM experiments. During transport to the STEM, this sample was exposed to ambient air for about 1 hour. The spherical particles deposited on the ITO substrates for the
UPS and 2PPE experiments have initially not been exposed to ambient air due to the use of the vacuum suitcase for the transport. Only later, in the course of the experiment, these samples have been deliberately exposed to air in order to achieve complete oxidation.

2.2 Scanning transmission electron microscopy

A detailed analysis of the nanoparticle structure and composition was conducted with a probe-corrected FEI Titan® 60-300 STEM with an X-FEG field emission source operated at a beam energy of 300 keV. The STEM system is equipped with an FEI Super-X four-quadrant detector for energy dispersive X-ray spectroscopy (EDS). For the acquisition of data, the number of channels was set to 1,024 with a dispersion of 10 eV per channel and an exposure time of 100 ms was used. For high-angle annular dark-field (HAADF) images the dwell time per pixel was set to 2.4 μs, while the size of the spectrum images (SI) was adjusted for each individual acquisition.

2.3 Photoemission electron spectroscopy

Photoelectron spectroscopy was conducted with a NanoESCA (FOCUS GmbH & Scienta Omicron) energy filtered photoemission electron microscope (EF-PEEM) at UHV conditions (~ 3 × 10⁻¹¹ mbar). A helium discharge lamp (HIS) operated at 21.2 eV and the second harmonic of an Indigo-S Ti:Sapphire laser system (Coherent) tuned to 3.02 eV (410 nm) were used as photon sources for the acquisition of ultraviolet photoelectron spectra (UPS) and two-photon photoelectron (2PPE) spectra, respectively. The pulse duration of the laser was 30 ns with a repetition rate of 5 kHz. The spot size of the HIS on the sample surface was about 330 μm, while the laser was focused to a spot size of about 50 μm (full width at half maximum) at a laser power of 0.5 mW. With the pass energy for the energy filter set to 50 eV and an entrance slit of 1 mm, an energy resolution of 200 meV was reached. An extractor field of 12 kV was applied with a sample distance of 1.63 mm. The energy filtered photoelectrons were collected and counted with a channeltron (CT), the measurement area was restricted to a spot size with (7 ± 1) μm diameter using an iris aperture.

3 Results and discussion

3.1 Nanoparticle structure and morphology

For STEM analysis, an amorphous carbon TEM grid was decorated with two different types of nanoparticles in order to give an example of the capabilities of the helium nanodroplet synthesis approach in terms of possible particle shapes. These two selected types encompass small, spherically shaped Ag@ZnO core@shell nanoparticles (source conditions: 20 bar and 8 K) deposited for 40 min, as well as wire-like Ag@ZnO core@shell nanoparticles (source conditions: 20 bar and 5 K) deposited for 3 min. Note that both types of nanoparticles were deposited on a single TEM grid, hence both species can be seen in the images presented in the following. During transport and transfer into the STEM, the sample was exposed to ambient conditions for about 1 h in total, which is enough time for the Zn shell to completely oxidize. This finding is in agreement with previous studies, which found that an oxide layer with a thickness up to 3 nm is formed directly upon exposure to air [34].

Figure 1(a) shows a selected HAADF image in which the two particle types can be identified. The bright Ag cores within the particles can be clearly distinguished from the surrounding ZnO since the HAADF image intensity is proportional to the square of the atomic number (Z²) [35]. It can be seen that the synthesis route based on the formation of Ag@Zn particles and subsequent oxidation of the Zn to ZnO results in a very uniform shell around the core, covering the Ag in a ZnO layer of constant thickness. This is true for the smaller spherical particles as well as for the larger wire-like structures as the presented STEM image shows. The inset plot (c) shows the fast Fourier transform (FFT) of the STEM image, yielding periodicities of (0.30 ± 0.02) nm and (0.18 ± 0.01) nm marked by the dashed orange circles. The sixfold symmetry of the FFT pattern is clearly observable and all visible features are assigned. The schematic in inset plot (d) explains the origin of the two observed lengths within the 2D projection of the hexagonal unit cell of wurtzite ZnO in [001] direction.

![Figure 1](image-url)
In order to gain information about the size distribution of the Ag cores are covered by a uniform ZnO shell.

A structural analysis of the uniformly grown ZnO within a selected particle is presented in Fig. 1 (b), which shows a detailed STEM HAADF image of the surface of a nanoparticle oriented in [001] direction. The contrast is enhanced and the observable hexagonal structure is highlighted by a red hexagon to guide the eye. The inset plot in Fig. 1(c) shows the corresponding fast Fourier transform (FFT), which allows conclusions on the crystal structure of the ZnO shell layer. The FFT image yields lattice distances of (0.30 ± 0.02) nm and (0.18 ± 0.01) nm, marked by dashed orange circles. The observed pattern clearly exhibits sixfold symmetry. All visible features are assigned with the corresponding Bravais-Miller indices. The six diffraction spots on the inner circle and the six spots on the outer circle show permutation symmetry among themselves and originate from the {100} and {200} planes of wurtzite ZnO, respectively. The obtained distances are in agreement with literature bulk values for the lattice spacing of 0.2814 and 0.1625 nm, taken from the Crystallography Open Database (COD) [36]. Deviations from the tabulated values most likely originate from the fact that the shell is only several atomic layers thick, which can lead to a lattice expansion as reported for several oxides on the nanoscale [37]. In the inset plot Fig. 1(d) the two periodically occurring lengths found in the FFT image are assigned to the two-dimensional projection of the hexagonal structure of ZnO in [001] direction.

Figure S3 in the ESM shows additional evaluated FFT images of Ag@ZnO nanoparticles in varying orientations, confirming the findings from Fig. 1. Overall, with the obvious sixfold symmetry and the lattice parameters in good agreement with literature, it is concluded that the ZnO shell layer prefers a wurtzite unit cell structure in case of the synthesized nanoparticles.

The chemical composition of the nanoparticles was determined by means of EDS. Figure 2(a) shows a HAADF image of several spherically shaped core@shell nanoparticles. An EDS map acquired from this area was used to create the spatially resolved elemental maps for Ag, Zn and O, which are shown in Figs. 2 (b)–2(d). In particular, the event counts around the Lα1 peak of Ag at 2.984 keV, the Lα peak of Zn at 1.012 keV, and the Kα peak of O at 0.523 keV were extracted from the EDS data and plotted according to their respective position in the image. It is evident that the positions of the bright core structures in the HAADF image are matching those with high abundance of Ag. The same applies for the whole particles including the shell and the respective EDS data for Zn. The slight offsets between particle positions in the HAADF image and the EDS maps may originate from a drift of the sample or a mobilization of the nanoparticles due to the exposure to the electron beam for longer times during the acquisition. These effects can hardly be avoided since the HAADF image and EDS data are acquired in independent scans. Note that the position of oxygen is not as precise as for the metals because it is also present in the background in form of hydrocarbons on the STEM substrate. Nonetheless, there is an increased abundance of O in the background corrected color map at the positions where Zn is also found. In combination with the findings of the structural investigation shown in Fig. 1, it is concluded that the Ag cores are covered by a uniform ZnO shell.

3.2 Size distribution and surface coverage

In order to gain information about the size distribution of the deposited particles, the overall diameters and Ag core diameters of 208 spherically shaped nanoparticles (synthesized at 20 bar and 8 K) were individually determined from high-resolution STEM images. Examples for the STEM images can be found in Fig. S2 in the ESM. The results are shown in the histograms in Fig. 3, containing the measured overall diameters of the particles in Fig. 3(a) and the core diameters in Fig. 3(b). For both datasets the diameters are, in good approximation, normally distributed. A fit yields a mean overall particle diameter of 5.84 nm with a standard deviation of 0.93 nm and a mean core diameter of (3.03 ± 0.64) nm. The width of both size distributions is inherent to the He synthesis process and originates from the broad droplet size distribution [14]. Over 100 of the wire-like nanoparticles were measured and 70% of the particles

![Figure 2](image-url)  
**Figure 2** (a) Detailed STEM HAADF image of a selected region with spherically shaped Ag@ZnO core@shell nanoparticles. The spatially resolved EDS maps reveal the spatial distribution of (b) the Ag core material as well as (c) the Zn and (d) the O in the shell of the particles.

![Figure 3](image-url)  
**Figure 3** From high-resolution STEM images, the particle and core diameters of 208 spherically shaped Ag@ZnO nanoparticles were measured. (a) Histogram of the measured particle diameters with a fitted normal distribution yielding a mean particle diameter and standard deviation of $\bar{d}_p = (5.84 \pm 0.93)$ nm. (b) Histogram of the measured Ag core diameters with the normal distribution resulting in a mean of $\bar{d}_c = (3.03 \pm 0.64)$ nm.
have a length between 20 and 50 nm. Their diameter ranges from 6 to 8 nm, however, can extend to over 10 nm for agglomerated structures. Note that the diameter can vary over the length of a single particle and that most of them are kinked or branched, which makes an exact characterization of their size difficult.

For the determination of the surface coverage for the deposited nanoparticles the overview STEM images shown in Fig. S1 in the ESM were used. With the ImageJ 1.52p software package [38] it was possible to separately evaluate the spherical (deposition time: 40 min) and wire-like (deposition time: 3 min) particles due to a discrimination in size. This is shown in Fig. S4 in the ESM for one of the STEM images. The evaluation yields an estimation for the obtained surface coverage of about (9.3 ± 0.5)% per hour for the spherically shaped particles and about (110 ± 30)% per hour for the elongated structures. The difference can be explained by the different helium droplet source conditions. For colder temperatures and larger droplets the deposition rate is enhanced, but a precise control of the particle shape is no longer possible and a variety of different shapes is formed.

The determined surface coverage rate for the spherically shaped Ag@ZnO nanoparticles is used to estimate the coverage of the glass coverslips coated with ITO prepared for analysis in the NanoESCA system. All three samples (pure Zn, pure Ag, and Ag@ZnO) were produced under the same experimental conditions (20 bar at 8 K) with a deposition duration of 5 h each, resulting in an estimated surface coverage of about (45.0 ± 2.5)%. Since the decorated ITO substrate cannot be analyzed by STEM, a scanning electron microscopy (SEM) overview image has been recorded, shown in Fig. S5 in the ESM. It confirms that the particles are distributed uniformly on the surface of the prepared samples, though the nanoparticles appear broadened at the given magnification due to the lower resolution of SEM.

3.3 Ultraviolet photoelectron spectroscopy

For the UPS and the 2PPE spectroscopy experiments with the NanoESCA system three different nanoparticle samples were prepared under the same conditions (20 bar at 8 K for 5 h) to guarantee comparability: pure Ag particles, ZnO particles and Ag@ZnO particles. All three samples are decorated with spherically shaped particles only, since it is of importance for the interpretation of the acquired spectra that the shape is uniform and the size distribution narrow. The elongated wire-like structures, as shown in Fig. 1 (a), are kinked and exhibit a wider range of lengths and shape. Therefore, they are not further evaluated by the employed spectroscopic methods that yield results integrated over an extended surface area. Furthermore, the spherical Ag particles typically exhibit a localized surface plasmon resonance (LSPR) around 400 nm, which is accessible by the Ti:Sapphire laser system [39]. Glass coverslides coated with ITO, which were also characterized which is accessible by the Ti:Sapphire laser system [39]. Glass coverslides coated with ITO, which were also characterized which is accessible by the Ti:Sapphire laser system [39]. Glass coverslides coated with ITO, which were also characterized.

The metallic nature of the partially oxidized Ag@ZnO is also reflected by the shape of the spectrum in the region that displays the energy range around the 3d core level peak of Zn. The orange line has a distinct and broad feature situated in this region with a double-peaked shape. A set of two Gaussians was used to evaluate the maxima of the convoluted peak structure. Thereby, the peak position at lower binding energy was found to be at (−9.7 ± 0.1) eV, marked by the vertical red line. The peak is attributed to the 3d level of pristine Zn. The second maximum, marked with a vertical black line, is located at (−10.3 ± 0.1) eV and represents the 3d level of the Zn atom in ZnO. A comparison with literature confirms the drawn conclusions about the origin of the peaks, though there is a certain spread in the published data [40–42]. The peak shift due to the chemical composition allows for a qualitative assessment of the oxidation state of the Zn atoms in the sample [43]. To ensure that the Zn is fully oxidized and the particles are comparable with the ones produced for the acquisition of the STEM data, the sample was exposed to ambient air for 1 h and then transferred back into the UHV chamber of the PEEM. The green line in Fig. 4 shows the UPS spectrum of the Ag@ZnO particles after the exposure to ambient conditions. The double peak structure is not present anymore, instead there is a single peak centered at (−10.8 ± 0.2) eV, which is again attributed to the 3d level of Zn in the oxidized state shifted to higher binding energies by 0.3–0.5 eV. This energy shift is an undesired side effect of the exposure to ambient air and the subsequent photo-hydroxilation of the sample surface driven by the extreme ultraviolet radiation during the acquisition of UPS data. The observed artifact was investigated in more detail by Gutman et al. in Ref. [44] for nano-crystalline zinc oxide thin films. Within the uncertainty of the measurement the energy shift found for the ZnO covered nanoparticles in this study is in good agreement with their results.

The important conclusion is that there is no sign of any metallic Zn left in the shell. The inelastic mean free path of electrons in inorganic compounds, considering the 21.2 eV photon energy provided by the HIS, is at least ~ 1.5 nm [45]. As this value exceeds the shell thickness of the spherically shaped particles with (1.33 ± 0.15) nm, it is concluded that the shell is fully oxidized after the sample has been exposed to ambient air.

The metallic nature of the partially oxidized Ag@ZnO is also reflected by the shape of the spectrum in the region that contains information about the population at the Fermi edge.
close to $E_{\text{band}} = 0$ eV. A pronounced Fermi cutoff, the spectral signature of a metal, is clearly visible for the corresponding orange spectrum. For the conductive plain ITO substrate the cutoff is barely visible and it seems to be even more suppressed for the Ag@ZnO sample, which has been exposed to air. Considering that at a surface coverage of 45% the recorded spectra correspond to a superposition of both substrate and nanoparticles, the absence of the Fermi cutoff indicates the fully oxidized, and thus semiconducting, ZnO as the dominant species.

Though this study aimed at the full oxidation of the Zn shell, there are intriguing possibilities for future experiments concerning the synthesis and formation of triple layer Ag@Zn@ZnO core@shell nanoparticles. The addition of a third layer, as already shown in literature, can potentially add new characteristics or improve existent properties of a layered nano-scaled system [46–48]. Furthermore, oxidation and reduction cycles could be applied to switch back and forth between Zn and ZnO as was already done e.g. for the Ni/NiO system [49]. Photoelectron spectroscopy offers a reliable method to discriminate between the metallic and semiconducting state and may be employed in future experiments to explore such systems and processes.

### 3.4 Two-photon photoelectron spectroscopy

2PPE spectroscopy is employed in order to investigate the plasmonic activity of the Ag cores within the Ag@ZnO nanoparticles. The 2PPE experiments are also carried out on ITO substrates covered only with spherically shaped nanoparticles and the conclusions in the following are drawn by comparing 2PPE spectra recorded for pure Ag and ZnO to Ag@ZnO samples. Note that the species containing Zn were already exposed to ambient air such that the shell is fully oxidized. The synthesis of Ag and Ag@ZnO nanoparticles took place at similar conditions ensuring that the plain Ag particles have about the same size as the Ag cores, with diameters in the range of 3–4 nm. For bare Ag nanoparticles in this size regime, it is known that the employed laser with a photon energy of 3.02 eV (410 nm) can excite localized surface plasmons [50, 51]. Note that the exact position of the LSPR also depends on the substrate and is, thus, not exactly known for the prepared particles on ITO. For particles produced by He droplet synthesis on fused silica, however, a relatively broad LSPR was found with a maximum around 450 nm [29].

In photoelectron spectroscopy, the electrons originating from the Ag core have to pass the ZnO shell in order to reach the detector. However, in contrast to the UPS measurements, the inelastic mean free path of the electrons after excitation with the laser beam is up to several tens of nm, which ensures that the acquired data contains information about the core material [45].

In order to compare the kinetic energy of the collected electrons, the work function ($\Phi$) difference between the samples has to be accounted for. $\Phi$ was determined by intersecting the abscissa with a tangent through the turning point of the rising edge of the secondary electron peak in the collected spectra. In addition, the Schottky effect, which emerges due to the presence of the 12 kV extractor field at a sample distance of 1.6 mm, has been considered [52] and introduces a reduction of $\Phi$ by $\Delta \Phi = 104$ meV. The obtained work functions correspond to $(3.7 \pm 0.2)\text{ eV}$ for ZnO, $(4.2 \pm 0.2)\text{ eV}$ for Ag, and $(4.0 \pm 0.2)\text{ eV}$ for Ag@ZnO. Since these values exceed the photon energy of the laser, it can be assumed that the spectra plotted as a function of $E_{\text{kin}}$ in Fig. 5 correspond to electrons excited in a two-photon process with a maximum overall available energy of 6.04 eV (at the Fermi cutoff). For a comparable graphic presentation the signal intensities are normalized to the maxima of the secondary electron peaks of the spectra.

In Fig. 5 it can be seen that the 2PPE spectrum of ZnO, depicted in blue, shows a prominent secondary electron peak that decays fast with increasing electron kinetic energy with only a very weak signal intensity towards the Fermi cutoff. However, in the case of the Ag spectrum (orange) a significant enhancement of electron yield at higher kinetic energies is observed. Such an increase in high energy photons has been attributed to an enhancement of the surface electric field by the excitation of plasmons in Ag nanoparticles [39, 53]. Intriguingly, the Ag@ZnO spectrum closely resembles the form of the plain Ag spectrum. Consequently, it is concluded that also in this case a localized surface plasmon is excited. The precise origin of the electrons, however, remains elusive. Plasmon excitation in the Ag core leads to an enhancement of the electric field close by the nanoparticle such that the photoelectrons may either be created in the Ag or the ZnO shell. Furthermore, in contrast to the plain Ag particles that were kept under UHV conditions, the Ag@ZnO particles were exposed to ambient air. The ZnO shells provide a protective layer preventing a contamination of the Ag cores with water or other hydrocarbons, while still preserving their plasmonic properties.

### 4 Conclusions

In the course of this study, we showed a novel way to produce high quality Ag@ZnO core@shell nanoparticles by employing the helium nanodroplet synthesis approach. Compared to other synthesis routes, this technique has the advantage of utilizing superfluid helium as pristine “nanolabs” in which the particles are grown without the addition of any solvents or other chemical agents. Thorough STEM investigations reveal the uniform and evenly distributed shape of the wurtzite ZnO shells surrounding the Ag cores, not only for spherically shaped particles but also for larger, wire-like core@shell structures. Thereby, the ZnO provides a very stable and reliable protective cover for the plasmonic Ag cores. This is indicated by 2PPE spectroscopy, where a significant plasmon enhancement of the photoelectron yield is observed for plain Ag as well as Ag@ZnO nanoparticles. Furthermore, the oxidation state of the Zn/ZnO shell system was investigated by accessing the Zn 3d core level with UPS, opening up possible routes for the synthesis and in situ monitoring of triple layer Ag@Zn@ZnO core@shell systems on the nanoscale. The stability of the Ag@ZnO nanoparticles and the preservation of important functional features upon exposure to ambient conditions sets the ground for possible future applications outside of the laboratory environment.
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