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Transparent Conductive Tantalum Doped Tin Oxide as Selectively Solar-Transmitting Coating for High Temperature Solar Thermal Applications

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Abstract

The transparent conductive oxide (TCO) SnO_2 :Ta is developed as a selectively solar-transmitting coating for concentrated solar power (CSP) absorbers. Upon covering with an antireflective layer, a calculated absorptivity of 95 % and an emissivity of 30 % are achieved for the model configuration of SnO_2 :Ta on top of a perfect black body (BB). High-temperature stability of the developed TCO up to 1073 K is shown *in situ* by spectroscopic ellipsometry and Rutherford backscattering spectrometry. The universality of the concept is demonstrated by transforming silicon and glassy carbon from non-selective into solar-selective absorbers by depositing the TCO on top of them. Finally, the energy conversion efficiencies of SnO_2 :Ta on top of a BB and an ideal non-selective BB absorber are extensively compared as a function of solar concentration factor *C* and absorber temperature *T*_H. Equal CSP efficiencies can be achieved by the TCO on BB configuration with approximately 50 % lower solar concentration. This improvement could be used to reduce the number of mirrors in a solar plant, and thus, the levelized costs of electricity for CSP technology.

Keywords: concentrated solar power, solar thermal electricity, solar-selective transmitter, transparent conductive oxide, high-temperature *in situ* characterization

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1. Introduction

Concentrating solar power (CSP) or solar thermal electricity (STE) is a commercially available technology in the emerging field of renewable energies. It is based on the concentration of sunlight onto an absorbing receiver, the transfer of the generated heat to a fluid and the subsequent conversion of thermal energy into electrical energy via a heat exchanger, steam turbine and electrical power generator. Currently, the most important CSP plant systems are parabolic trough collectors and central receiver towers, operated at up to 673 K and 873 K respectively [1]. In order to enhance their energy conversion efficiency, higher operation temperatures are required. Additionally, based on physical limits outlined later on, solar selectivity is indispensable to enhance the performance of solar thermal power plants, as emphasized in the reviews of Kennedy [2], Granqvist [3,4] and Weinstein [5].



Fig. 1. Solar spectrum AM 1.5 and black body spectrum (divided by 4 for better visibility) at 1073 K calculated by Planck's law. The blue dashed line shows reflectance R of an ideal selective absorber. The wavelength at which the ideal selective absorber switches from non-reflective to reflective is called cut-off wavelength $\lambda_{\text{cut-off.}}$

The concept of solar selectivity utilizes the different characteristic wavelength ranges of solar spectrum and thermal radiation. Translated into the terminology of CSP, the absorption of solar light occurs at much shorter wavelengths than the radiative emission of a black body $B(\lambda)$ which it heats (**Fig. 1**). The solar spectrum $G(\lambda)$ with the air mass coefficient (AM) 1.5 represents the yearly average of solar irradiance at mid-latitudes on the surface of the earth [6]. It can be approximated by the emissivity of a black body (BB) at T = 5777 K calculated by Planck's law, taking into account the intensity reduction due to the small solid angle and atmospheric absorption losses [7, 8]. Its intensity maximum at 515 nm occurs at a much shorter wavelength than that of a BB at T = 1073 K ($\lambda_{max} \approx 2700$ nm), the target temperature of current research and development for CSP tower plants (Fig. 1) [9, 10]. The energetic separation of the peak intensities can be used to design solar-selective coatings (SSCs) resembling the ideal characteristics, i.e., very low reflectance *R* in the range of the solar spectrum and high reflectance in the range of the BB radiation (blue dashed line in Fig. 1). The wavelength at which the coating switches from non-reflective to reflective is called cut-off wavelength, $\lambda_{cut-off}$. **Equation 1a** and **1b** show the standard definitions of the two crucial material parameters of CSP, absorptivity α and emissivity ε [9,10].

$$\alpha = \frac{\int_{300 nm}^{2500 nm} [1 - R(\lambda)] G(\lambda) d\lambda}{\int_{300 nm}^{2500 nm} G(\lambda) d\lambda}$$
(1a)

$$\varepsilon = \frac{\int_{1000 nm}^{25000 nm} [1 - R(\lambda, T)] B(\lambda, T) d\lambda}{\int_{1000 nm}^{25000 nm} B(\lambda, T) d\lambda}$$
(1b)

Using α and ε , the efficiency η for the conversion of solar energy into mechanical energy can be calculated by **Equation 2**:

$$\eta = \eta_{opt} \cdot \eta_{Carnot} = \left(\alpha - \frac{\varepsilon \sigma \left(T_H^4 - T_{amb}^4\right)}{I C}\right) \left(1 - \frac{T_{amb}}{T_H}\right)$$
(2)

where σ is the Stefan-Boltzmann constant, $T_{\rm H}$ the absorber temperature, $T_{\rm amb}$ the ambient temperature, I the integrated solar irradiance and C the sun concentration factor, given by the ratio of collecting and absorbing area in a solar plant. The first term of Equation 2 is called optical efficiency; the second term represents the Carnot cycle efficiency. Clearly, a CSP absorber coating should have a high absorptivity and low emissivity at a given temperature. Nevertheless, it has to be noted that the absorptivity has a much larger influence on the efficiency compared to the emissivity. Inserting the values for σ and I = 892 W/m², which corresponds to AM 1.5, and assuming $T_{\rm H} = 1073$ K, $T_{\rm amb} = 303$ K and C = 1000, the ratio of emissivity and absorptivity in Equation 2 is only 0.08. This means, a solar-selective transmitter under these conditions can only be useful if it decreases the bare BB emissivity by at least 13 times the TCO-induced absorptivity loss.

Selectivity can be achieved by different designs of the absorber coating. The most intensively studied solar selective materials for temperatures up to 673 K – 773 K are metal-dielectric multilayers [13-15], ceramic-metal composites (cermets) [16,17] and semiconductor-metal tandems [18]. Some of these are already commercialized [19]. Very recent research on multilayer [7,20] or composite SSCs [21] addressed even temperatures in the range up to 1000 K. Textured surfaces [22] and selective transmitters [23] can also provide solar selectivity. The application of transparent conductive oxides (TCOs) for low emissivity glasses was studied intensively as shown in the work of Granqvist [4], but only a few studies using TCOs on an absorbing substrate were published. Fan and Bachner reported on In₂O₃:Sn thin films with $\alpha = 0.90$ and $\varepsilon = 0.081$ (at 394 K) as well as on In₂O₃:Sn coated with a 100 nm MgF₂ anti-reflection layer with $\alpha = 0.85$ and $\varepsilon = 0.081$ (at 394 K), assuming a perfect BB substrate [24]. SnO₂:F prepared by spray pyrolysis at 1113 K on black enameled steel was investigated in the work of Haitjema et al. [25]. It exhibited $\alpha = 0.91$ and $\varepsilon = 0.15$, and was thermally stable up to 523 K for 6 weeks and up to 673 K for 8 days in air, respectively. Shimidzu et al. deposited In₂O₃:Sn on a tungsten substrate and reported α = 0.71 as well as emissivities of ε = 0.08 at 353 K and ε = 0.13 at 973 K. The stack was thermally stable at 973 K in vacuum for up to 10 hours when prepared with a SiO_2 interlayer between metal substrate and TCO [26]. Taylor et al. studied a wide range of TCOs including F-, Sb- and Cd-doped SnO₂ (FTO, ATO, CTO), Sn doped In₂O₃ (ITO) and B-, Al-, Ga- and F-doped ZnO (BZO, AZO, GZO, FZO) [23]. In that theoretical study, ITO and FTO showed the best solar selectivity in the temperature range up to 873 K.

However, none of the previous studies reported on any coating with very low reflectance in the solar spectral range and, consequently, $\alpha \ge 0.95$ on a BB substrate in combination with thermal stability up to at least 1073 K. Moreover, the parameter space of $T_{\rm H}$ and C, where TCOs are beneficial for CSP, has not been explored yet. Hence, an investigation of the TCO on BB configuration as alternative concept for CSP plants of the next generation is lacking so far.

In this work, a Ta-doped SnO₂ TCO on top of a BB is developed as selectively solar-transmitting coating for high temperature CSP technology. In addition to its excellent TCO properties, the material selection is based on the demand for thermal, chemical, mechanical and environmental/chemical stability (such as oxidation stability and resistance to moisture), as well as for cost-effectiveness [27]. The results of ex situ optical, electrical and structural characterization by spectroscopic photometry, spectroscopic ellipsometry (SE), Hall-effect measurements and Rutherford backscattering spectrometry (RBS) were used to optimize the deposition temperature and dopant concentration. *In situ* SE and RBS were applied to prove thermal stability of SnO₂:Ta at temperatures up to 1073 K. An anti-reflective (AR) SiO₂ top layer with appropriate thickness allows to achieve an absorptivity of 96% for the TCO on top of a BB configuration. As proof of concept, silicon and glassy carbon were transformed into solar-selective materials by covering them with the TCO. Finally, the advantages and limitations of the TCO on BB concept for high temperature CSP technology are discussed for a ($T_{\rm H}$, C) parameter range that comprises the current and projected future solar plant operation conditions by comparing its efficiency with that of a non-selective BB absorber.

2. Materials and methods

SnO₂:Ta thin film samples were deposited by direct current reactive magnetron sputtering from two metallic 2" targets (Sn (99.99 at.%) and Sn:Ta (6.7 at.% Ta)). Argon was used as sputter gas. The working pressure was varied between 0.9 Pa and 1.0 Pa, and the oxygen partial pressure was fixed at 0.11 Pa. Fused quartz was used as substrate for coating development in order to be able to perform optical measurements in reflection and transmission mode. The Ta concentration was varied from 0 to 1.8 at.% by changing the sputtering powers. In a second sample series, the deposition temperature was varied from 710 K to 907 K. Having optimized the TCO, silicon and glassy carbon substrates were coated for two reasons: i) to investigate the TCO properties on absorbing surfaces and ii) to study to which extend absorbers can be made solar-selective.

For optical characterization, specular reflectance at 5° incidence angle and normal angle transmission were measured in the spectral range from 300 nm to 3000 nm using a SolidSpec 3700DUV spectrometer (Shimadzu). An M-2000 ellipsometer (Woollam) was used for SE measurements in the range from 211 nm to 1688 nm. The WVASE32 software (Woollam) was applied for optical modelling of the measured data to obtain the optical constants, thickness and roughness of the studied coating materials, and moreover, for all the optical simulations in this work [28].

The thermal stability of the optimized SnO₂:Ta thin films was investigated under high vacuum conditions in a cluster tool [29,30] installed at the 6 MV ion beamline of the Ion Beam Center of the HZDR [31]. Temperature-dependent *in situ* SE of SnO₂:Ta on fused quartz was recorded in the spectral range from 371 nm – 1000 nm during a heating ramp of 15 K/min from room temperature (RT) to 1073 K. Temperaturedependent *in situ* RBS was performed using 2 MeV He⁺ ions with a beam current of ~ 20 nA. The heating procedure consisted of three cycles, including heating the sample with 20 K/min from RT to 1073 K, keeping it at this temperature for 30 min (1st and 2nd cycle) and 180 min (3rd cycle) and then cooling it back to RT. In total, the sample was exposed to 1073 K (800 °C) for four hours. The beam spot diameter on the sample was < 1 mm. The backscattered ions were detected by a silicon surface barrier detector under an angle of 155°. The charge carrier density N_e and the electrical charge carrier mobility μ_{el} were determined by Hall-effect measurements using the van der Pauw method [32]. Every sample was measured five times to estimate a realistic error for those values.

3. Results and discussion

3.1 Design and optimization of SnO₂: Ta as solar-selective transmitter

As introduced in section 1, $\lambda_{\text{cut-off}}$ is a crucial parameter for the design of solar-selective coatings. Because the solar spectrum and the high-temperature BB spectrum overlap, $\lambda_{\text{cut-off}}$ separates the spectral ranges of absorption and thermal emission of a selective absorber and thus, it determines the numerical values for α and ε obtained by Equation 1a and 1b (Fig. 2a). Assuming $\lambda_{\text{cut-off}} = 2500$ nm, the sunlight is completely absorbed ($\alpha = 1$), but at the cost of losing 20 % of the BB radiation due to thermal emission ($\varepsilon = 0.2$). For $\lambda_{\text{cut-off}} = 1000$ nm, the thermal emission is completely suppressed ($\varepsilon = 0$), but only 73 % of the sunlight is absorbed ($\alpha = 0.73$).



Fig. 2. a) Absorptivity α and emissivity ε of an ideal selective absorber ($T_H = 1073$ K) in dependence of the cut-off wavelength $\lambda_{\text{cut-off}}$ and b) optical efficiency for four different concentration factors in dependence of $\lambda_{\text{cut-off}}$ calculated with $T_H = 1073$ K, $T_{\text{amb}} = 303$ K, I = 892 W/m². The dashed lines represent the efficiency of a non-selective, perfect BB absorber ($\alpha = 1, \varepsilon = 1$).

The effect of solar selectivity on the optical efficiency of solar absorbers is estimated according to Equation 2 for an absorber temperature of 1073 K and four different concentration factors, which are relevant for CSP plants (**Fig. 2b**). The shape of the obtained functions is determined by i) the increase of α until the integration includes the complete solar spectrum up to 2500 nm, ii) the efficiency-enhancing effect of an increasing concentration factor C and iii) the efficiency-reducing effect of the emissivity ε in the infrared range. The dashed lines represent the efficiencies of a non-selective, perfect BB absorber ($\alpha = 1$, $\varepsilon = 1$) for the different *C* values. The intersection between the dashed and the solid line of each color represents the $\lambda_{cut-off}$ from which an ideal selective absorber is more efficient than the BB absorber alone. In general, solar selectivity improves the optical efficiency of a solar absorber, and hence, the total efficiency of CSP plant systems for a wide range of cut-off wavelengths and concentration factors by a substantial amount (Fig. 2b). An optimal optical efficiency of ≥ 90 % is achieved when $\lambda_{cut-off}$ is between 1500 nm and 2500 nm. The possibility of tuning the cut-off wavelength in order to match the optimized value for a given combination of absorber temperature and concentration factor would therefore be a significant progress for design and development of SSC's for CSP.

It is a well-known fact from semiconductor physics that charge carrier density and mobility are responsible for the wavelength and slope of the reflectance edge in the absorption spectra [33,34]. **Fig. 3a and 3b** show simulated reflectance spectra of a doped SnO₂ coating (thickness 500 nm) on top of a transparent fused quartz substrate as a function of charge carrier concentration and mobility. The optical constants of the substrate have been determined by SE measurements and modelling. They correspond exactly to the SiO₂ model [35] provided by the database of the WVASE32 optical modelling software [28]. The basic dielectric function of SnO₂, described by a single Lorentz oscillator for the interband absorption, which was used for the simulation, is also obtained from the same database. It was extended by a Drude oscillator to account for the contribution of the dopant charge carrier mobility μ_{opt} , were varied in meaningful intervals in accordance to the values given in the work of Sago et al. [36] in order to show their influence on the reflectance. For better comparability, the effective electron mass was fixed at $m^* =$ 0.275 m_0 , although Sago et al. states a slight dependency of m^* on N_e ($m^* = m_0$ (0.275+0.01×10⁻²⁰× N_e)), where m_0 is the free electron mass [36].



Fig. 3. Calculated reflectance *R* of doped SnO₂ (500 nm) on fused quartz as a function of a) charge carrier density N_e at fixed optical charge carrier mobility $\mu_{opt} = 30 \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ and b) μ_{opt} at fixed $N_e = 3 \times 10^{20} \text{ cm}^{-3}$ of a Drude oscillator. The effective mass m^* was set to 0.275 m_0 [37]. The SnO₂ interband absorption was described, according to literature [28], using a high energy Lorentz oscillator with amplitude A = 1.1853, energy position E = 5.7507 eV and broadening B = 0.64275 eV.

The calculated reflectance of doped SnO₂ in the UV, visible (vis) and near IR (NIR) spectral range is of the order of 10 % (Fig. 3). Intensity oscillations due to interferences show the transparency of the coating in the UV-vis range. In contrast, the IR part of the spectrum, above the plasma wavelength λ_P , is characterized by a high reflectance due to excitation of free charge carriers. These are the desired properties for a solar-selective transmitter. The variation of N_e results in changes of $\lambda_{cut-off}$ (Fig. 3a), which for real selective materials could be defined as λ where 50 % of the maximum reflectance is reached. The variation of μ_{opt} on the other hand results in different slopes of the plasma edge and therefore, in a different degree of selectivity of the films (Fig. 3b). Since there exists an optimal $\lambda_{cut-off}$ for a given absorber temperature, the charge carrier density N_e can be optimized in order to match the requirements for a maximum CSP efficiency at a specific operation temperature. However, it should be noted that low values of N_e , even if favored for a suitable $\lambda_{cut-off}$, also lead to lower values of IR reflectance and therefore higher emissivity. The optical mobility of the charge carriers should be as high as possible, as this increases the IR reflectance, and thus, the selectivity of the coating.

Following these considerations, Ta content and deposition temperature T_s of as-deposited SnO₂:Ta samples were optimized with respect to charge carrier density and mobility by preparing two sample series as described in the experimental section (**Fig. 4**).



Fig. 4. Optimization of SnO₂:Ta thin film deposition parameters on fused quartz. a) Charge carrier concentration and charge carrier mobility and b) resistivity as a function of the Ta content (at.%) at a fixed deposition temperature of 693 K. c) Charge carrier concentration and charge carrier mobility and d) resistivity as a function of the deposition temperature at fixed Ta content of 1.2 at.%.

It should be noted that optical (**Fig. 3**) and electrical mobility (**Fig. 4**) do not have to have the same value [38,39]. A discrepancy between them is indicative for different charge carrier scattering mechanisms in a sample. Electrical mobility is measured for a macroscopic sample size and is influenced by intragrain and intergrain charge carrier scattering [40]. It can be lower than the optical mobility, when the free electron path, upon electric field oscillations, is shorter than the sample's grain size. Then, only intragrain scattering occurs [41]. Nevertheless, high values of electrical mobility also imply high optical mobility, and the electrical characterization is a suitable indicator for the quality of a solar-selective TCO.

The variation of the Ta content results in a minimum of resistivity ρ at approximately 1.2 at.% (**Fig. 4b**), which is in perfect agreement with a previous study of radio-frequency magnetron sputtered SnO₂:Ta [42]. At lower Ta concentrations, the charge carrier concentration N_e and mobility μ_{el} approach the values of undoped SnO₂ (Fig. 4a), resulting in a high resistivity (Fig. 4b). Higher Ta concentrations (> 1.5 at.%) do not increase N_e considerably, but lead to decreasing μ_{el} and higher resistivity. This is probably due to a higher scattering probability at the ionized Ta impurities, or clustering of the Ta ions [43,44]. The variation of the substrate temperature, T_s , results in a minimum of ρ at approximately 840 K - 860 K, which is very similar to the value of 873 K reported in the literature [42]. Samples deposited at lower temperatures showed low mobility and charge carrier concentration, probably due to poor crystallinity and small amount of Ta on Sn lattice sites. This behavior was already reported for other TCOs [44]. In conclusion, a Ta content of 1.2 at.% and a substrate temperature of 850 K were established as deposition parameters to prepare optimized SnO₂:Ta TCO films.

3.2 Thermal stability and optical characterization of optimized, selectively-transmitting SnO₂:Ta thin films

The optimized TCO films used for the following investigations had the composition 67.1 at.% O, 31.6 at.% Sn and 1.2 at.% Ta. Their thermal stability up to a temperature of 1073 K was investigated *in situ* by SE and RBS. SE shows marginal changes in the measured ellipsometric angles (**Fig. 5**), which are attributed to small temperature-induced deformations of the sample holder and the resulting slight misalignment. Thus, *in situ* SE provides evidence for the conservation of layer thickness and optical properties of the SnO₂:Ta thin film, both during and after high-temperature treatment.



Fig. 5. Temperature dependence of the spectroscopic ellipsometry angles a) ψ and b) Δ during *in situ* heating of a SnO₂:Ta (1.2 at.% Ta) thin film on fused quartz from RT to 1073 K.

The in-depth compositional stability of the SnO₂: Ta thin films was studied by a temperature-dependent in situ RBS experiment (Fig. 6). Minor reproducible steps of the RBS energies occurred during mapping just when the final temperature of each heating ramp was reached (Fig. 6b). At the highest temperature itself, the energies were constant, and the changes were fully reversed when re-cooling was started. Thus, the steps are attributed to the detector response at the highest temperatures. With this in mind, the pre- and post-heating spectra (Fig. 6a), as well as the *in situ* map (Fig. 6b), reveal the conservation of the elemental depth distribution throughout the entire film thickness over both time and temperature. In conclusion, both in situ experiments demonstrated the excellent optical and chemical stability of the SnO₂:Ta thin films at temperatures up to 1073 K in vacuum. Since the substrate and the antireflective layer have the same chemical composition, SiO₂, these in situ investigations allow at least the well-founded assumption that neither mixing processes nor chemical reactions are to be expected between the anti-reflective layer and the TCO layer. Moreover, possible clustering or segregation processes of the dopant that can occur in doped materials at high temperatures were excluded for SnO₂:Ta (1.2 at.% Ta) by the demonstrated temperature stability. It should be noted that there are no previous records of in situ RBS measurements at such high temperatures in the analysis of solar thermal materials. The in situ techniques applied here are ideally suitable for upcoming CSP technologies and can moreover provide important insight into failure mechanisms of coatings in general (e.g. substrate atom migration or diffusion on layer interfaces) and their dynamics, which is impossible with standard post-heating analysis.



Fig. 6. Temperature dependent in situ RBS study of a SnO₂: Ta (1.2 at % Ta) thin film on fused quartz in the range from RT to 1073 K. a) Single RBS spectra taken at RT before and after the heating cycles to 1073 K (800 °C) and b) in situ RBS map taken during three heating cycles between RT and 1073 K with a total dwell of 4 hours at the highest temperature.

For reliable optical and electrical characterization of the optimized SnO₂:Ta (1.2 at.% Ta) thin films, a series of five samples was deposited using the optimized deposition parameters described in section 3.1. Hall measurements yielded a mean charge carrier concentration of $N_e = (4.3 \pm 0.3) \times 10^{20}$ cm⁻³ and a mean charge carrier mobility of $\mu_{el} = (14 \pm 3) \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$. The dielectric function was determined by means of SE and spectrophotometry, and used to build an optical model comprising the contributions of the SnO₂:Ta layer, a quartz reference substrate and a surface roughness layer on top. The latter is described by an effective medium approximation (EMA) layer, a superposition of the optical constants of a 50/50 vol.% material/void mixture. This is a common approximation to simulate surface roughness [45,46]. The dielectric function of SnO₂:Ta was described by a Tauc-Lorentz oscillator for intraband transitions, which, in comparison to the previously used Lorentz oscillator from the optical database, allows direct access to the optical band gap energy E_{gap} of the material [47]. E_{gap} of SnO₂:Ta (1.2 at.% Ta) is found to be 4.5 eV. Compared to the commonly observed value for undoped SnO₂, $E_{gap} \approx 3.9$ eV, it is increased by doping due to Burstein-Moss shift [41]. Additionally, a Drude oscillator was used to account for the free charge carrier contributions as explained in section 3.1.



Fig. 7. Experimental and simulated reflectance of a 555 nm SnO₂:Ta (1.2 at.% Ta) coating on fused quartz substrate.

The reflectance spectrum shows a distinct solar-selective behavior with $\lambda_{\text{cut-off}}$ at ≈ 1900 nm. The average charge carrier concentration determined by optical analysis of SnO₂:Ta (1.2 at.% Ta) is $N_e = (4.4 \pm 0.2) \times 10^{20}$ cm⁻³, which is in good agreement with the electrically measured value of $N_{\rm e}$. On the other hand, the optical and electrical charge carrier mobilities differ, with $\mu_{opt} = (35 \pm 3) \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$ and $\mu_{el} = (14 \pm 3) \text{ cm}^2 \text{V}^{-1} \text{s}^{-1}$, for reasons explained in the previous section [39,48]. These results further show the films are highly suited to provide solar selectivity (Fig 3. and Fig. 7).

3.3 Optimization of the SnO₂: Ta thin film as selectively-transmitting coating

In order to optimize the SnO₂:Ta (1.2 at.% Ta) thin film as selectively-transmitting coating, the effect of an AR coating on top of the TCO was explored. According to Fresnel's equations, small differences of refractive indices n at an interface lead to low reflectance. Thus, SiO₂ with 1.42 < n < 1.50 in the solar range is a suitable material for this purpose, if deposited on a SnO₂:Ta layer with a refractive index of 0.5 < n < 2.1, as determined in this work from ellipsometry and reflectivity measurements in the range 0.5 to 5 eV. **Fig. 8a** shows the change of reflectance of a 805 nm thick SnO₂:Ta layer on fused quartz when a 50 nm thick AR layer of SiO₂ is deposited on top. **Fig. 8b** displays the calculated absorptivity of the AR/TCO/ fused quartz system as a function of the AR layer thickness. It is evident that the absorptivity of the layer stack is increased from 0.89 to a maximum of 0.96 for a 90 nm thick SiO₂ AR layer, after which it decreases. The remaining loss of 4 % results from the reflection at the AR layer in the solar range. It cannot be further reduced by optimizing thicknesses [49]. The simulated effect of the AR layer was experimentally confirmed by the absorptivity enhancement from $\alpha = 0.90$ for the bare TCO on BB to $\alpha = 0.94$ for SnO₂:Ta with a 50 nm AR SiO₂ layer (Fig. 8b). The slightly reduced IR reflectance (and therefore higher emissivity ε) of < 5 % (Fig. 8a) is negligible compared to a similar percentage of absorptivity increase. Additionally, the nonreactive SiO₂ layer is very likely to ensure thermal stability even under atmospheric conditions.

Having found the favorable AR layer thickness, the influence of the TCO layer thickness and of the film roughness were studied. Therefore, the absorptivity was experimentally studied as a function of the TCO layer thickness (Fig. **8c and 8d**). The thickness and roughness effect were then simulated for the model system ideal BB ($\alpha = 1$), optimized TCO and AR surface layer with and without roughness as a function of SnO₂: Ta thickness (Fig. 8c and 8d). Simulations of different SnO₂:Ta thicknesses without taking surface roughness into account gave an absorptivity minimum at 300 nm due to interference effects, but the overall variation of α in the considered film thickness range from 200 nm to 10000 nm is less than 1 % and therefore marginal (Fig. 8d, open squares). However, the experimental data show a larger variation of α when the TCO thickness increased from 290 nm to 1285 nm. This can be attributed to a simultaneous roughness increase from 8 nm to 30 nm. Modelling the roughness of the deposited samples gives a much better agreement between experimental and simulated data (Fig. 8d, open circles). In the simulation, the roughness varied from 6 nm to 80 nm for the simulated TCO thickness range from 200 nm to 10000 nm. The roughness layer acts as an anti-reflective coating as it consists of 50 % SnO₂: Ta and 50 % voids and can therefore be thought of as a material with reduced refractive index. Hence, the roughness of the TCO layer should be known in order to apply an AR layer with optimal thickness to maximize absorptivity. The independence of the performance on the film thickness itself, however, is a big advantage of the TCO on BB configuration in comparison to other selective absorbers, which usually have to meet very precise thickness requirements in the designed multilayer stacks.



Fig. 8. Influence of an antireflective SiO_2 top layer on the measured reflectance (a) and the calculated absorptivity (b), and influence of the SnO_2 : Ta layer thickness on the measured reflectance (c) and the calculated absorptivity (d). The absorptivities were obtained assuming a perfect BB absorber beneath the TCO with and without roughness.

3.4 Proof of concept for selectively transmitting coatings on top of a pure absorber

It is now an interesting question, whether non-selective absorber materials can be transformed into solar-selective one's by the developed TCO. Therefore, a 320 nm thick SnO₂:Ta layer was deposited on top of a silicon wafer and a glassy carbon substrate.



Fig. 9. Reflectance of bulk and of SnO₂:Ta-coated (320 nm) a) silicon and b) glassy carbon substrates showing the transformation of the two materials into solar-selective absorbers by depositing the TCO on top of them.

Si and glassy carbon are absorbers with absorptivities of 0.65 and 0.83, respectively. Both have low reflectance in the IR range, and thus, are lacking of solar selectivity (**Fig. 9a and 9b**). After deposition of a SnO₂: Ta thin film on top of them, the layer stacks exhibit distinct solar-selective behavior: enhanced absorptivity of 0.81 (SnO₂:Ta on Si) and 0.86 (SnO₂:Ta on glassy carbon), a $\lambda_{cut-off}$ of \approx 1800 nm and high reflectance, i.e. reduced emissivity in the IR. The absorptivity enhancement is due to the previously described anti-reflective effect of the SnO₂:Ta. In

particular the Si/ air ($n_{\text{Si}, 300 \text{ nm}} = 5.1$; $n_{\text{Si}, 2500 \text{ nm}} = 3.4$ [50])/ $n_{\text{air}} = 1.0$) interface is replaced by a SnO₂:Ta/ air ($n_{\text{SnO2:Ta}}$, $_{300 \text{ nm}} = 2.1$; $n_{\text{SnO2:Ta}, 2500 \text{ nm}} = 0.5$) and a Si/ SnO₂:Ta interface, leading to an absorptivity increase of 25 %. These results show yet another advantage of the TCO on BB absorber concept, as all absorbing materials could be turned into solar-selective absorbers, provided that their surface roughness is low compared to the TCO thickness.

3.5 Comparison of the solar conversion efficiency: selectively transmitting TCO on a black body absorber vs. bare black body absorber

According to Equation 2, the evaluation of the optical efficiency of a solar absorber also requires the knowledge of the emissivity. Simulations of the IR reflectance based on the optical model presented in section 3.2 were applied to calculate ε of the TCO on top of a perfect BB substrate. According to Equation 1b the emissivity of SiO₂ (90 nm)/ SnO₂:Ta (1000 nm) at 1073 K is $\varepsilon = 0.265$. As a conservative estimation $\varepsilon = 0.30$ and $\alpha = 0.95$ were used to compare the solar energy to mechanical energy conversion efficiencies of the bare BB and the TCO on BB concept. The following configurations were compared: a) SiO₂ (90 nm)/ SnO₂:Ta (1000 nm) coated BB ($\alpha = 0.95$, $\varepsilon = 0.3$) and b) uncoated non-selective BB ($\alpha = 1$, $\varepsilon = 1$). The efficiencies, η , of the two model systems were calculated using Equation 2 and are displayed for a relevant (*C*, *T_H*) parameter space in **Fig. 10**. Numeric efficiency values for selected (*C*, *T_H*) combinations are listed in **Table 1** and **Table 2**.



Fig. 10. Solar energy to mechanical energy conversion efficiency (η) maps of a SiO₂ (90 nm)/SnO₂:Ta (1000 nm) coated BB absorber ($\alpha = 0.95$, $\varepsilon = 0.3$), (left panel) and a non-selective BB absorber ($\alpha = 1$, $\varepsilon = 1$), (right panel) as function of concentration factor *C* and absorber temperature *T*.

For the discussion of this comprehensive data set, i) the efficiency η for a given parameter combination of *C* and $T_{\rm H}$ and ii) the maximum efficiency achievable for a given *C* value, $\eta_{\rm max}$, are used. While the former value represents a pure performance parameter, the latter can be associated with the number of mirrors necessary to achieve a certain solar plant efficiency. As such, the maximum efficiency is of predictive value for the design of CSP plants. The most general conclusion from Fig. 10 is that the TCO on BB configuration leads to higher efficiencies for lower concentration factors. Going through the results from high to low concentration factors, for 10000 > *C* > 4000 the emissivity reduction due to the TCO cannot compensate the lowered absorptivity and thus, the TCO coating even worsens the overall CSP efficiency. In the range of 4000 > *C* > 1000 the TCO coating can improve the efficiency at temperatures above 1100 K. However, there are no commercial CSP plants operating at these

concentration factors and temperatures yet. At 1000 > C > 100, which are typical values for central receiver tower plants in operation or under construction, the TCO coated system achieves similar or higher efficiencies in the whole temperature range. At the target temperature, $T_{\rm H} = 1073$ K, the improvement of the absolute efficiency is 0.36 for C = 100 (**Table 1**). However, the maximum efficiencies for C = 100 are achieved at even lower temperatures: $\eta_{max, \text{TCO/BB}} = 0.58$ ($T_H = 870$ K) and $\eta_{max, \text{BB}} = 0.53$ ($T_H = 700$ K). This corresponds to a relative efficiency improvement of >9 % due to the TCO. For 100 > C > 10, which covers the typical concentration factors of parabolic trough systems, a large improvement in efficiency can be achieved for all temperatures considered in this work. The maximum efficiencies at C = 10 are $\eta_{max_0 \text{TCO/BB}} = 0.40$ ($T_H = 570$ K) and $\eta_{max_0 \text{BB}} = 0.30$ (T = 460K), respectively, corresponding to a TCO-caused relative improvement of 33 %.

Table 1. Comparison of efficiencies of a SiO₂ (90 nm)/SnO₂:Ta (1000 nm) coated BB absorber ($\eta_{TCO/BB}$, $\alpha = 0.95$, $\varepsilon = 0.3$) and a non-selective BB absorber (η_{BB} , $\alpha = 1$, $\varepsilon = 1$) at the targeted temperature $T_{\rm H} = 1073$ K for different concentration factors (*C*).

С	$\eta_{ m TCO/BB}$	$\eta_{ m BB}$	$\eta_{ m TCO/BB}$ - $\eta_{ m BB}$
100	0.54	0.18	0.36
200	0.62	0.47	0.15
300	0.65	0.56	0.09
500	0.67	0.63	0.04
1000	0.69	0.69	0
5000	0.70	0.73	-0.03
10000	0.71	0.74	-0.03

Table 2. Comparison of efficiencies of a SiO₂ (90 nm)/SnO₂:Ta (1000 nm) coated BB absorber ($\eta_{\text{TCO/BB}}$; $\alpha = 0.95$, $\varepsilon = 0.3$) and a non-selective BB absorber (η_{BB} ; $\alpha = 1$, $\varepsilon = 1$) for different operation temperatures (T_{H}) and two concentration factors, C = 100, and C = 1000.

$T_{\rm H}[{ m K}]$	$\eta_{\mathrm{TCO/BB}}$ (C = 100)	η_{BB} (C = 100)	$\eta_{\text{TCO/BB}} - \eta_{\text{BB}}$ (C = 100)	$\eta_{\mathrm{TCO/BB}}$ (C = 1000)	$\eta_{\rm BB}$ (<i>C</i> = 1000)	$\eta_{\text{TCO/BB}} - \eta_{\text{BB}}$ (C = 1000)
400	0.30	0.31	-0.01	0.30	0.32	-0.02
600	0.51	0.50	0.01	0.52	0.54	-0.02
800	0.58	0.51	0.07	0.62	0.64	-0.02
1000	0.57	0.31	0.26	0.68	0.69	-0.01
1200	0.46	-	0.46	0.71	0.68	0.03
1400	0.24	-	0.24	0.71	0.62	0.09

Looking at the results of Fig. 10 from another perspective, equal efficiencies are achieved at lower concentration factors by the TCO/ BB system, with the consequence of operating the absorber at higher temperatures. For example, $\eta = 0.55$ is reached with C = 60 instead of C = 135, accompanied by increasing $T_{\rm H}$ from 740 K to 790 K. Taking the target temperature ($T_{H} = 1073$ K) of this work as a maximum operation temperature, $\eta_{\rm max} = 0.66$ can be achieved with C = 335 compared to C = 620 for the non-selective system at $T_{H} = 970$ K. This corresponds to a reduction in *C* by more than 45 %.

4. Conclusion

The concept of using a solar-selective TCO on top of a black body was explored for mid- and high-temperature CSP technology. Optimized films of SnO₂:Ta were developed, whose cut-off wavelength $\lambda_{\text{cut-off}}$ and spectral selectivity were shown to be tunable in the relevant solar range. In situ SE and RBS measurements demonstrated the compositional, optical and structural stability of the TCO up to 1073 K in vacuum. Its performance was shown to be thickness-independent in a range from several hundred nanometers to microns. Hence, the effort of manufacturing very precise layer thicknesses that is required for complex multilayer structures of state-of-the art SSCs is avoided. The optimized SnO_2 :Ta/ SiO₂ layer stack of this study has reflective losses of < 5 %. When applied on a non-selective BB substrate ($\alpha = 1, \varepsilon = 1$), the BB's emissivity decreases to $\varepsilon \leq 30$ %. The comparison of the energy conversion efficiency of the entire BB/TCO/AR layer stack with conservative values for absorptivity and emissivity, $\alpha = 0.95$ and $\varepsilon = 0.30$, with that of an ideal BB revealed potential application ranges of the developed coating for a wide parameter space of solar concentration factor, C, and absorber temperature, $T_{\rm H}$. Systems with 1000 > C > 100 can be improved by up to 0.05 considering absolute maximum efficiencies η_{max} . The highest absolute efficiency improvements of 0.05 - 0.10 are possible for C < 100, which is typical for parabolic trough systems. Equal efficiencies in combination with higher absorber temperatures can be obtained with a significant reduction of the sun concentration factor, C. At absorber temperatures from 570 K up to the targeted value of this work ($T_{\rm H} = 1073$ K), this reduction in C amounts to 46 % to 60 %. The improvement could be translated into significantly lower levelized costs of electricity produced by solar power plants by reducing the number of mirrors, the areal footprint, management costs and last but not least the costs of mirror maintenance. It has to be emphasized at this point that the comparison with a perfect black body is the worst-case scenario for the SnO₂:Ta system, since the BB absorptivity is inherently less than 1 for real materials with substantial negative consequences for its efficiency.

The universality of the selective transmitter concept was experimentally proven by the deposition of the TCO on bare absorbing surfaces, namely silicon and glassy carbon, which showed distinct solar-selective optical response thereafter. Hence, non-selective absorber materials can be turned into selective absorbers. Additionally, the performance of the TCO is independent on its thickness. Both these properties are very attractive from an application point of view. The excellent TCO properties open the opportunity of applying this concept as an alternative to currently used non-selective absorbers and even to solar-selective coatings based on complex multilayer structures of oxides or nitrides. Finally, compared to currently used pigment paints which have a higher initial efficiency but suffer from relatively fast degradation over time, the robust thermal and structural stability of our films is another strong argument to their deployment in the concentrated solar power plant industry.

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