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Topical Review

Electron Mobility in Oxide Heterostructures

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Abstract

Next-generation integrated circuit devices based on transition-metal-oxides are expected to boast a variety of extraordinary properties, such as superconductivity, transparency in the visible range, thermoelectricity, giant ionic conductivity and ferromagnetism. However, the realisation of this so-called oxide electronics as well as the study of their unconventional physics is stalled by inferior carrier mobilities compared to conventional semiconductor materials. Over the past 10 years, bulk conducting oxides and oxide heterostructures with superior carrier mobilities have nonetheless seen significant progress. This progress is signifying the approaching era of oxide-based electronic circuits along with novel solid-state phenomena originating from the combination of hybridized oxygen p orbitals, transition-metal d orbitals and electronic correlations. Here, we review the recent advancements and results on high mobility oxide heterostructures based on SrTiO3 and ZnO as well as other prominent oxides.

Section 1: Introduction

The race for achieving high carrier mobility in semiconductor materials has proved beneficial from both a scientific and technological perspective. It has led to electron mobilities exceeding 35,000,000 cm²/Vs in Al0.87Ga0.13As-based material systems [1], the realization of quantum phenomena such as integer and fractional quantum Hall effect [2,3] and new concepts for achieving pure materials with high carrier mobilities [4]. High carrier mobilities are particularly beneficial for: i) the study of quantum phenomena where extended mean free paths are required. ii) an increased control over detrimental scattering mechanisms. iii) applications requiring fast electronic responses (e.g. high-frequency devices). iv) achieving low-resistance materials without increasing the carrier density, which is advantageous for, e.g., thermoelectric materials and transparent conductors. Whereas the development of high-mobility silicon and III-V semiconductors in many cases is very mature, appealing functionalities can be obtained by considering materials beyond these. Some of the new promising materials for these high mobility electronic applications are the oxides and oxide heterostructures.

1.1 – Why oxides?

Heterostructures based on transition-metal-oxides (TMOs) exhibit a variety of extraordinary properties, such as superconductivity, transparency in the visible range, thermoelectricity, giant ionic conductivity and ferromagnetism. The electronic properties of transition-metal oxides are determined by a delicate balance between hybridized oxygen p orbitals, transition-metal d orbitals and electronic correlations. The interfaces based on TMOs can therefore lead to unconventional physics and find use in the present and coming generations of electronic devices. Transparent conducting oxides are for example heavily used in flat panel displays and solar cells.

The interest in these transition-metal oxides has motivated the research in epitaxial growth of layered oxides such as oxide heterostructures. This line of research seeks, in part, to emulate the achievements of previous generations of semiconductor materials researchers who over the past few decades have learned to engineer the electronic properties of epitaxial grown semiconductor materials by exploiting e.g. epitaxial strain and modulation doping. Because of the wider range of phenomena, the implications for physics and technology of advances in transition-metal oxides are likely to be large. The worldwide race between crystal growers from major research laboratories to continuously improve the growth of oxide interfaces has led to high electron mobilities. One of the most extensively studied metal oxide heterointerface systems is the LaAlO3/SrTiO3 (LAO/STO) interface [5]. The initial mobility value of 10³ cm²/V s of this system was surpassed by replacing the perovskite LaAlO3 with spinel structured γ-Al2O3, which resulted in a record-high electron mobility greater than 10⁵ cm²/Vs at 2 K [6]. Another system that is well studied is the ZnO/ZnMgO material system, which shows extremely high mobilities exceeding 10⁵ cm²/Vs at 2 K [7]. These electron mobilities were made possible by improving the quality of the epitaxial oxide thin films prepared with state-of-the-art deposition techniques such as oxide molecular beam epitaxy (MBE) or pulsed laser deposition (PLD), which allow the growth of heterostructures with very sharp interfaces. By fabricating and understanding the high mobility in oxides, we may be able to produce high-quality electrical components with the added functionalities that oxides exhibit.
In this paper, we review the advancements and recent results made in high mobility oxide heterostructure materials. First, we describe the relevant scattering mechanisms and general pathways used to enhance the mobility in Section 2. In Section 3 we review the electron mobility in three-dimensional conducting SrTiO$_3$ (STO) while Section 4 presents a detailed discussion on the high mobility obtained in confined electron gases of representative STO-based systems. We then review the electron mobility of three-dimensional conducting ZnO in Section 5 whereas the subsequent Section 6 covers the development in high-mobility Mg$_x$Zn$_{1-x}$O/ZnO heterostructures. Lastly, in Section 7 we provide a brief overview of the literature concerning mobility in oxides beyond STO and ZnO.

**Section 2: Scattering mechanism & pathways to enhance mobility**

Improving the mobility of any conductor is of importance for fundamental understanding of solid-state physics but also for applications. Therefore, understanding the different carrier scattering mechanisms that limits the mobility in functional materials such as oxides is similarly important.

When attempting to understand electronic transport and optical properties in oxides, one essentially aims at accessing the eigenfunction and total energy of the electrons residing in the particular oxide. The universally employed method for accessing this information is the so-called effective mass approximation where electrons essentially are treated as free electrons with a modified mass, $m^*$.

When an electric field ($E$) is applied to the oxide, charge carriers will accelerate. The acceleration is counteracted by elastic or inelastic scattering events, and the charge carriers end up travelling with an average drift velocity ($v_d$). The electron mobility ($\mu$) is defined as the ratio between the drift velocity and electric field:

$$\mu = \frac{v_d}{E}$$

A large carrier mobility can be obtained by increasing the scattering time ($\tau$) or decreasing the effective mass ($m^*$):

$$\mu = \frac{e \tau}{m^*}$$

The scattering time can vary greatly from oxide to oxide depending on e.g. the defect concentration, carrier concentration and phonon.

The electron or hole mobility is typically measured experimentally using the Hall effect. As charges move in an out-of-plane magnetic field, the current ($I$) will deflect and form a transverse Hall voltage ($V_H$). The Hall voltage can be related to the carrier density ($n$) and mobility ($\mu$) by

$$R_H = \frac{d(V_H/I)}{dB} = -\frac{1}{ne}$$

and

$$\mu = \frac{R_H}{R}$$

with $R$ being the resistivity of the material. This mobility is known as the Hall mobility and is importantly used as a figure-of-merit between different oxide conductors. In this case, $\tau$ in Eq. 2 describes the momentum relaxation time ($\tau_{\text{rel}}$) and corresponds to the average time it takes for the momentum of a charge carrier to reduce to 37% due to scattering events. Another characteristic mobility of materials is the quantum mobility in which the relevant time scale ($\tau_{\text{q}}$) relates to the time it takes loose quantum coherence by the scattering events. This is typically measured using Shubnikov-de Haas oscillations. The quantum mobility may differ significantly from the Hall mobility as established in heterostructures based on GaAs [8], STO [9,10] and ZnO [11].

Table 1 shows examples of $m^*$, Hall mobility and quantum mobility at $T < 2$ K when electrons reside in a heterostructure for two important oxides that will be the focus of this review: STO and ZnO.

<table>
<thead>
<tr>
<th>Material system</th>
<th>$m^*$</th>
<th>$\mu$, Hall (2 K)</th>
<th>$\mu$, quantum (20 mK)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STO [6]</td>
<td>$1 , m_e$</td>
<td>$1.4 \times 10^3 , \text{cm}^2/\text{Vs}$</td>
<td>$7.2 \times 10^3 , \text{cm}^2/\text{Vs}$</td>
</tr>
<tr>
<td>ZnO [7]</td>
<td>$0.3 , m_e$</td>
<td>$1.2 \times 10^3 , \text{cm}^2/\text{Vs}$</td>
<td>$4.8 \times 10^3 , \text{cm}^2/\text{Vs}$</td>
</tr>
</tbody>
</table>

The charge carriers undergo scattering from one or multiple sources and the total mobility ($\mu$) is typically assumed to be composed of the individual scattering contributions ($\mu_i$) according to the Matthiesen rule:

$$\mu^{-1} = \sum_i \mu_i^{-1}$$

In the following we will introduce some important scattering mechanisms for oxides. All subsequent equations are in their generalized form (i.e. not material specific) and describe the case for three-dimensional conducting structures. Here it is important to note that alternative expressions exist for confined electronic systems such as two-dimensional electron gases, but these will not be covered here. Additional information on various scattering mechanisms, and mobility expressions for electronic systems with reduced dimensionality can be found in, e.g., [12,13].

### 2.1 – Ionized Donor and Impurity Scattering

In many cases the dominant scattering mechanism at low temperatures is scattering off ionized point defects or defect clusters. This includes ionized donors that are intentionally introduced to the materials as well as ionized impurities that unintentionally are present in oxides often in high concentrations.

For oxides in the low carrier density regime where the contribution from ionized impurity scattering is largest, it is possible to express the electron mobility as follows:

$$\mu = \frac{3 \pi e^2}{2 \hbar^2 a_p^2} \frac{n}{N_s \ln(\pi k_T a_p)}$$

where $a_p$ is the effective Bohr radius given by $a_p = 4 \pi e \hbar^2 / m^* e^2$, $n$ is the carrier density and $N_s$ is the concentration of ionized impurities.

At large carrier concentrations, $n$ will become comparable with $N_s$. Beyond this point, the concentration of ionized donors in the oxide will exceed the concentration of ionized impurities and most electrons will scatter on donors. Thus, the mobility in the large carrier density regime is given by:
where $a$ is the lattice size of the given oxide. To reduce the effect of ionized donor scattering, one can employ delta doping or modulation doping strategies [14,15]. In delta doping, the donors are positioned in a thin sheet whereas the charge carriers extend beyond this, and thus part of the charge carriers reside in parts of the oxide without charged donors. In modulation doping, the electrons and donors are spatially separated so that the interaction is only through long-range Coulomb attraction. If the defects remain neutral, the scattering cross-section is much smaller than the ionized defects and therefore generally less detrimental to the overall mobility.

2.2 – Electron-phonon interaction

As the temperature increases, more phonon modes become populated and the interaction between charge carriers and lattice vibrations becomes gradually more important for determining the carrier mobility. Lattice vibrations are particularly important in oxides where the high electronegativity of oxygen creates polar bonds leading to polaron formation and oscillating electric fields that may severely affect the charge flow:

2.2.1 – Polaron formation

As the charged carriers move through the ionic oxide, charge-lattice interactions deform the lattice. If the interaction is sufficiently strong, the charge carriers can no longer be considered to be freely moving, but rather move in the lattice as a heavier polaron. The effect of polaron formation is generally described through a mass renormalization where the band effective mass ($m_b$) is replaced with a polaron effective mass ($m_p$). The increased mass of the polarons leads to a lowering of the mobility as it will be presented in Section 4.2.1 for the case of STO-based structures.

The mass enhancement can be circumvented by lowering the electron–phonon coupling ($\alpha$) according to $m_p = m_b(1 + \alpha/6)$ and $m_p = m_b(1 + (\alpha/8)\alpha + 0.1272\alpha^2)$, which is valid for three- and two-dimensionally confined carriers, respectively, with Fröhlich interactions to a three-dimensional lattice [16,17].

2.2.2 – Acoustic phonon scattering

The coherent motion of acoustic phonons typically scatter charge carriers by locally offsetting the bands. When considering e.g. $n$-type oxides, the acoustic phonon modes change the interatomic distances and lead to local changes in the conduction band that scatters the electrons. The contribution to the mobility ($\mu_{\text{AC}}^{\text{3D}}$) is given by [12,18]:

$$\mu_{\text{AC}}^{\text{3D}} = \frac{2e\hbar v_s^2}{3\pi n_{3D}m^*a_c^2} \ln(1 \exp(E_F/k_B T))$$  \hspace{1cm} (8)

where $\rho$ is the crystal density, $v_s$ is the velocity of sound, $n_{3D}$ is the three-dimensional carrier density, $E_F$ is the Fermi energy, $a_c$ is the conduction band deformation potential, $k_B$ is Boltzmann constant and $T$ is the temperature.

Acoustic phonon scattering is reported to be the dominant scattering mechanism for e.g. $p$-type ZnO from $T = 10 – 300$ K [19]. The acoustic phonon scattering can generally be counteracted by decreasing the temperature, varying the carrier density and modifying the phonon modes.

2.2.3 – Optical phonon scattering

When the longitudinal optical (LO) phonon modes of an ionic lattice are excited, macroscopic electric fields are created that scatter the charge carriers. In the weak/intermediate coupling regime where $\alpha < 6$, Low and Pines derived an expression for the mobility used frequently for three-dimensional conducting STO (as well as two-dimensionally confined carriers in STO-based structures) [16]:

$$\mu_{\text{LO}} = \frac{\hbar}{2\alpha \hbar \omega_{\text{LO}} m_p} \frac{e}{m_b^3} f(\alpha) \exp\left(\frac{\hbar \omega_{\text{LO}}}{k_B T}\right)$$  \hspace{1cm} (9)

where $\omega_{\text{LO}}$ is the angular frequency of the longitudinal optical phonon mode and $f(\alpha)$ is a monotonic function of $\alpha$ that varies slowly from 1 to 1.35 as $\alpha$ increases from 0 to 6 [16]. Other expressions for longitudinal optical phonon scattering have also been derived [12]. Transverse optical (TO) phonon modes, however, create no net macroscopic electric field, and they interact with the electrons through deformation of the bands. The associated mobility contribution ($\mu_{\text{TO}}^{2\text{D}}$) can be written as [12,18]

$$\mu_{\text{TO}}^{2\text{D}} = \frac{\sqrt{2\pi} e \hbar v_s^2}{(m^*)^{5/2} \hbar \omega_{\text{TO}} d_0^2} \exp\left(\frac{\hbar \omega_{\text{TO}}/k_B T}{E_F + \hbar \omega_{\text{TO}}}ight) - 1$$  \hspace{1cm} (10)

where $\omega_{\text{TO}}$ is the angular frequency of the transverse optical phonon mode and $d_0$ is the optical phonon deformation potential.

The general pathways to decrease the influence of optical phonon scattering include lowering the temperature, decreasing the electron–phonon coupling/deformation potential and modifying the phonon modes. The latter can be done e.g. using strain, elemental substitution with lighter or heavier atoms or using interfaces or thin films to break the lattice symmetry [12].

2.2.4 Piezoelectric scattering

Crystals without inversion symmetry can be piezoelectric active, and here the acoustic phonon modes may lead to changes in the electric fields that can scatter charge carriers strongly. The mobility in e.g. $p$-type ZnO is significantly limited by piezoelectric scattering [19,20].

2.3 – Electron-electron scattering

Electron-electron interactions often play an important role in determining the properties of oxides. This is particularly the case for correlated transition metal oxides with the conduction band derived from $d$-orbitals. The electron-electron interaction may also severely affect the carrier mobility. The Hall mobility is only sensitive to changes in the total carrier momentum. This leaves the Hall mobility insensitive to regular electron-electron collisions where the momentum is conserved. However, some electron-electron collisions involve the lattice such as Umklapp scattering events and collisions that promote electrons to another band with a different effective mass. The contribution from electron-electron interactions ($\mu_{\text{ee}}$) is usually described by

$$\mu_{\text{ee}} = AT^{-2}$$  \hspace{1cm} (8)

where, $A$ is the temperature coefficient and $T$ is the temperature. For STO, this contribution is likely to be the dominant scattering mechanism for temperatures between $5$ K $< T < 150$ K [21].

2.4 – Interface roughness scattering

For many devices it is crucial to confine the charge carriers to two or fewer dimensions in order to effectively tune the device properties with a gate. This requires a broken symmetry, which is typically created by making a heterostructure or exploiting the surface of the material. In both cases, the broken lattice symmetry inherently forms imperfections and defects, which serves as scattering sites for the charge carriers. The common procedure to reduce the interface roughness scattering is to lattice match the materials used to form the heterostructure. For additional details on interface roughness scattering in e.g. STO, see reference [22].

2.5 – Additional scattering mechanisms
There are several other scattering mechanisms that are not described in detail here. These includes (but are not limited to) scattering with extended lattice imperfections such as grain boundaries, domain walls and dislocations [12]. Spin-related scattering mechanisms may also be of high importance for materials exhibiting e.g. Kondo effect and colossal magnetoresistance [12].

**Section 3: Electron mobility in bulk SrTiO$_3$**

Electronic transport in bulk conducting SrTiO$_3$ (STO) have been studied for more than half a century. Since the first papers appeared in the early 1960s [23,24], much effort has been spent in understanding how electrons are transported through STO. STO displays both metallic behaviour and superconductivity for a wide range of charge carrier densities remarkably even down to very dilute values [25]. The charge carriers in STO originate either from intentional donors or from a reduction of the STO crystal - i.e. formation of oxygen vacancies through vacuum annealing. Much of the exciting physics in STO arise due to the empty 3$d$ shell of titanium. Due to the crystal field splitting, the 3$d$ levels split into the twofold and threefold degenerate $e_g$ and $t_{2g}$ levels, respectively. When STO is doped, electrons populate the $t_{2g}$ levels, which lead to a bulk $n$-type metallic conductivity of STO [26].

When STO display metallic behaviour, the carrier density changes very little as the temperature is reduced from room temperature to cryogenic temperatures. However, as the dielectric constant of STO, and consequently the degree of electrostatic screening, undergoes dramatic changes in this temperature interval, the electron mobility in STO is profoundly susceptible to temperature changes (see Figure 1). Through this change in electrostatic screening, the electron mobility can increase by almost a staggering four orders of magnitude as the temperature is reduced from 300 K to 2 K, reaching values well in excess of 10,000 cm$^2$/Vs (see Figure 1). This exciting behaviour has fostered a substantial and ongoing effort in trying to understand the underlying solid-state physics of bulk conducting STO. The physical properties of STO are given in Table 2.

![Figure 1](image_url)  
*Figure 1 – Initial studies of bulk conducting STO uncovered a large temperature dependence of the electron mobility. Reprinted figure with permission from [23]. Copyright 1967 by the American Physical Society.*

<table>
<thead>
<tr>
<th>Physical property</th>
<th>Value</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Molar mass</td>
<td>183.49 g/mol</td>
<td>PT</td>
</tr>
<tr>
<td>Appearance</td>
<td>White, opaque crystals</td>
<td>PT</td>
</tr>
<tr>
<td>Density</td>
<td>5.11 kg/m$^3$</td>
<td>PT</td>
</tr>
<tr>
<td>Melting point</td>
<td>2080 °C</td>
<td>PT</td>
</tr>
<tr>
<td>Boiling point</td>
<td>N/A</td>
<td>PT</td>
</tr>
<tr>
<td>Refractive index</td>
<td>2.394</td>
<td>PT</td>
</tr>
<tr>
<td>Direct and wide band gap</td>
<td>Indirect/direct band gap of 3.26 eV/3.75 eV at 2 K</td>
<td>[27,28]</td>
</tr>
<tr>
<td>Large exciton binding energy</td>
<td>220 meV</td>
<td>[29]</td>
</tr>
<tr>
<td>Piezoelectric constants</td>
<td>14 C/m$^2$ (e$<em>{13}$), -0.5 C/m$^2$ (e$</em>{15}$)</td>
<td>[30]</td>
</tr>
<tr>
<td>Effective mass</td>
<td>$m_e$</td>
<td>[6]</td>
</tr>
<tr>
<td>High-frequency dielectric constant</td>
<td>100 at 2 K</td>
<td>[31]</td>
</tr>
<tr>
<td>Static dielectric constant</td>
<td>20000 at 2 K</td>
<td>[32]</td>
</tr>
<tr>
<td>Lattice constant</td>
<td>$a = 3.904$ Å at 2 K</td>
<td>[33]</td>
</tr>
<tr>
<td>Sound velocity</td>
<td>7.9 x 10$^3$ m/s at 300 K</td>
<td>[34]</td>
</tr>
</tbody>
</table>

Table 2 – The physical parameters of STO adopted from the literature or the periodic table (PT).

### 3.1 - Temperature dependence of the mobility

A large number of different scattering mechanisms have been used to try to explain the mobility of STO as a function of the temperature, which vary greatly (see Figure 1). In general, the temperatures are divided into three regimes: At low temperatures (typically $T < 10$ K), scattering with ionized impurities and donors in STO dominate charge carrier transport. This temperature regime will be discussed in Subsection 3.3. At intermediate
temperatures (typically $10 \, \text{K} < T < 150 \, \text{K}$), a $\mu \propto T^{-n}$ behaviour with $n \approx 2$ is observed and generally ascribed to electron-electron interactions [21,35–37] (see Figure 2), although other explanations have also been proposed based on temperature dependent polaron [38] or dielectric [39] properties and scattering with acoustic [18], transverse optical [18] or longitudinal optical phonons [39]. The detailed understanding of the scattering mechanism in this temperature regime remains elusive, and Mikheev et al. indeed recently showed that the scattering rate is independent on the carrier density in contrast to what is expected from Landau Fermi liquid theory [35]. Lastly, as it will be discussed in the next subsection, the dominant electron scattering contribution close to room temperature ($T > 150 \, \text{K}$) is generally attributed to longitudinal optical (LO) phonons [21,23,24]. However, in particular for high carrier densities, the electron-electron interaction may dominate all the way up to room temperature [21].

As mentioned above, LO phonons are thought to be one of the dominant contributions to room temperature electron scattering in STO. The effect of this contribution is reduced in most materials when the charge carrier density is increased. In other words, more charge carriers correspond to less scattering due to an increased effective screening of the phonons. However, for small charge carrier densities, the electron mobility of STO remains essentially unaltered compared to the situation at large carrier densities (see Figure 3). Mikheev et al. suggested that this unconventional behaviour of the electron mobility with carrier density was due to the presence of a significant electron-electron scattering contribution [21]. The effect of this contribution should scale directly with the charge carrier density, i.e. with less charge carriers corresponding to less scattering. This electron-electron scattering mechanism will hence compete with the scattering contribution due to LO phonons. Interestingly, these two effects should overall cancel with each other leading to the observed constant electron mobility with charge carrier density. Mikheev et al. estimated the crossover carrier density where the two contributions are equal to be around $n_{3D} \approx 6 \times 10^{18} \, \text{cm}^{-3}$ (see Figure 4).

3.2 – Electron mobility at room temperature

Unlike most conventional materials, the room temperature electron mobility of STO is almost independent with the carrier density. Specifically, the electron mobility remains between $\sim 1$–$8 \, \text{cm}^2/\text{Vs}$ for carrier densities spanning several orders of magnitude between $10^{16}$–$5 \times 10^{20} \, \text{cm}^{-3}$ (see Figure 3). Note that only samples with $n_{3D} > 2 \times 10^{16} \, \text{cm}^{-3}$ presented in Figure 3 showed metallic behaviour down to $T = 2 \, \text{K}$, whereas samples with $n_{3D} < 2 \times 10^{16} \, \text{cm}^{-3}$ showed a resistance upturn at some point while cooling from room temperature.
3.3 – Electron mobility at low temperature

Opposed to the situation at room temperature, the low-temperature electron mobility in STO is very sensitive to variations in the charge carrier density (see Figure 5).

As mentioned above, the low-temperature electron mobility is limited by scattering with ionized impurities. These charged impurities come in two varieties: ionized background impurities and ionized intentional impurities. In the case of the former, the possible background donors include oxygen vacancies, while common background acceptors include strontium vacancies, aluminium or iron [41–43]. In the case of the latter, the common intentional donors include oxygen vacancies, niobium and lanthanum. As these impurities ionize upon incorporation into the STO lattice due to donor-acceptor compensation, there reside substantial concentrations of scattering centres for charge carriers in STO. Usually, the concentration of background impurities is around $N_i = 5 \times 10^{18} \text{ cm}^{-3}$ whereas the concentration of intentional impurities naturally depends on the doping level [41]. This means that for $n_{3D} \ll N_i$, electrons in STO will primarily scatter on ionized background impurities and consequently represent the intrinsic upper limit of electron mobility in STO. On the other hand, when $n_{3D} \gg N_i$, the low-temperature electron mobility is instead limited by scattering on the intentionally added ionized donors. These different scattering regimes and the corresponding overall electron mobility in STO is given according to as described in Subsection 2.1 and Equations 6 and 7. Using the Matthiessen additive rule to the different contributions, it is possible to express the overall electron mobility for any given charge carrier density. This overall mobility obtained by combination of the different scattering contributions can then be calculated for carrier densities between $1 \times 10^{17} - 5 \times 10^{20} \text{ cm}^{-3}$ and is shown in Figure 5.

3.4 – Electron mobility of strained STO

Until this point, we have been dealing with the electron mobility of STO having an unperturbed and intrinsic lattice. The effect of a strained STO lattice on electron mobility was studied by Jalan et al. [44]. Impressively, the authors found that the electron mobility of La-doped STO thin films under uniaxial stress could be substantially increased along the direction of the stress. For one of the two different carrier densities studied, the electron mobility exceeded a staggering 120,000 cm$^2$/Vs at 1 K (see Figure 6). Importantly, the authors note that through strain engineering it could be possible to realise STO under biaxial stress with a uniform high electron mobility. It is interesting to observe that, in this study, the effect of stress was only affecting the low-temperature electron mobility and for $T > 20 \text{ K}$ did not seem to change the transport properties noticeably. This may indicate that the strain effect is primarily affecting the scattering contribution from ionized impurities.
By applying a uniaxial stress, the electron mobility of STO can be increased significantly. Reprinted from reference [44], with the permission of AIP Publishing.

Conversely, Kobayashi et al. studied the effect of deliberately introducing strontium vacancy clusters in Nb-doped STO [45]. Due to the strontium vacancy clusters, the STO films were remarkably shown to exhibit enhanced electron mobilities exceeding 50,000 cm$^2$/Vs at 2 K (see Figure 7). According to the authors, this behaviour was due to the strontium vacancy clusters creating a compressive strain field throughout the STO crystal. Opposed to the above case of uniaxial strained STO, this strain field manifested itself in an increase of the electron mobility for the entire temperature range studied ($T = 2 – 300$ K). Therefore, it is apparent that these strontium vacancy clusters may affect, not only a single but possibly, several scattering mechanisms.

In order to understand the effect of strain in STO on electron mobility, Janotti et al. modelled biaxially strained STO using first-principles calculations [46]. Here, they found that STO with tensile biaxial stress in the (100) or (110) planes indeed had a lowered effective mass of the lowest energy conduction band along the transport directions. From the above studies, it is clear that strain is an effective pathway to increase the low-temperature electron mobility, and in some cases even increase it above the intrinsic upper limit dictated by ionized background impurities of $\sim 30,000$ cm$^2$/Vs at 2 K. The usual way to design and control the strain of a given material is to place it in proximity with another material in a so-called heterostructure. As this has been studied extensively in the case of STO, the electron mobility of STO-based heterostructures will be the topic of the following Section 4.

**Section 4: Electron mobility in confined STO-based systems**

Besides the strain-approach described in the previous Section, the electron mobility has been enhanced beyond what is usually observed in bulk conducting STO by confining the electrons to a thin sheet. This is achieved by breaking the lattice symmetry either by doping only a confined region with, e.g., La donors ($\delta$-doping) or by forming a heterostructure where STO is interfaced with another material. In both these cases, the electrons are confined to reside near the dopant layer or interface. A careful control of the local scattering landscape underpins the success of modulation doping where the donors are moved to a layer spatially separated from the electrons. The downside of this is, however, that the lattice symmetry breaking may introduce additional scattering mechanisms such as interface roughness scattering as described in Section 2.4.

Specifically, confined conductivity may be formed by doping STO locally either by: i) intentional dopants such as La, ii) electronic reconstructions due to polar discontinuities across the heterostructure or iii) introduction of oxygen vacancies. In STO-based heterostructures, the translation symmetry near the interface is broken and the previously degenerate $t_{2g}$ levels split further in accordance with their specific orbital orientation.
These split $t_{2g}$ levels are comprised of the $3d_{xy}$, $3d_{xz}$ and $3d_{yz}$ levels, which usually all are populated in confined STO-based systems [26].

In particular, the confined conductivity – i.e. a two-dimensional electron (2DEG) gas – found at the LaAlO$_3$/SrTiO$_3$ interface has been studied extensively for the past fourteen years. However, its $p$-type counterpart – the two-dimensional hole gas (2DHG) – has proved elusive. Very recently it was demonstrated that a highly mobile 2DHG is realisable in epitaxially grown SrTiO$_3$/LaAlO$_3$/SrTiO$_3$ heterostructures [47,48]. This provides a platform for future applications using confined electron–hole systems.

In the following, we will review a representative selection of different high-mobility STO-based material systems with confined electrons, and outline the mechanisms proposed to explain the mobility enhancement.

### 4.1 – Temperature dependence of the mobility

The transport properties of a range of high-mobility STO-based material systems are summarized in Figure 8. For the majority of the systems, it is evident from Figure 8 that the carrier densities are temperature independent. At low temperatures, the mobility varies greatly from one material system to another with $\gamma$-Al$_2$O$_3$/SrTiO$_3$ exhibiting the highest mobility exceeding 100,000 cm$^2$/Vs at 2 K. Several other heterostructures also obtains high electron mobilities larger than 20,000 cm$^2$/Vs at 2 K. Similar to bulk conducting STO, the mobility at the intermediate temperatures roughly varies as $\sim T^{-2}$ characteristic for electron-electron scattering up to $T \sim 150$ K where longitudinal optical phonon scattering often becomes dominant. The effect of this electron-electron scattering contribution is similar to that in Figure 2.

![Figure 8](https://example.com/figure8.png)

Figure 8 – The sheet resistance, carrier density and electron mobility of a range of different confined STO-based systems along with bulk conducting STO. Note that the data presented here represent the highest reported electron mobilities of the respective systems. The data is obtained from: $\gamma$-Al$_2$O$_3$/SrTiO$_3$ (GAO/STO) [6]; amorphous-LaAlO$_3$/LaSrMnO$_3$/SrTiO$_3$ (a-LAO/LSM/STO) [49]; CaZrO$_3$/SrTiO$_3$ (CZO/STO) [50]; SrCuO$_2$/LaAlO$_3$/SrTiO$_3$ (SCO/LAO/STO) [51]; bulk SrTiO$_3$ (STO) [52]; $\delta$-doped SrTiO$_3$($\delta$-doped STO) [53]; LaAlO$_3$/SrTiO$_3$ (LAO/STO) [54]; amorphous-LaAlO$_3$/SrTiO$_3$ (a-LAO/STO) [55]. Common for these high-mobility systems are that the carrier density often only is little temperature dependant, whereas the electron mobility can differ by up to four orders of magnitude from room temperature to 2 K. At intermediate temperatures between $T = 10 - 150$ K, the electron mobility follows an overall $\mu \propto T^{-2}$ dependence characteristic for electron-electron scattering.
4.2 – Mobility at room temperature

For both bulk conducting STO and two-dimensionally confined electron gases in STO, the room temperature mobility is generally less than 12 cm²/Vs (see Figure 3 and Figure 8). The room temperature mobility is limited by longitudinal optical phonon scattering or electron-electron scattering. Moderate variations ranging from 2 to 12 cm²/Vs is observed upon varying the growth conditions, confinement, carrier density and material systems. There is a tendency for obtaining a larger room temperature mobility when increasing the carrier density as for example observed in γ-Al₂O₃/SrTiO₃.

![Figure 9](image)

**Figure 9** – Electron mobility ($\mu$), electron-phonon coupling ($\alpha$) and polaron mass ($m_p$/$m_e$) as a function of the Hall carrier density ($n_s$) for γ-Al₂O₃/SrTiO₃ heterostructures with varying carrier densities obtained by varying the deposition conditions or performing post-annealing in 1 bar oxygen at 200 °C. Reprinted figure with permission from [56]. Copyright 2018 by the American Physical Society.

4.2.1 – γ-Al₂O₃/SrTiO₃

The interface between spinel γ-Al₂O₃ and perovskite STO becomes $n$-type conducting as oxygen vacancies are formed in the near-interface region of STO [6,57]. The sheet carrier density ($n_s$) can be tuned by varying the deposition conditions or performing a post-annealing step in an oxygen-containing atmosphere after the deposition [57]. The room temperature mobility was found to be enhanced as the carrier density increased, reaching a value up to 12 cm²/Vs at $n_s \geq 8 \times 10^{13}$ cm⁻² (see Figure 9). When decreasing the carrier density, the mobility reduces to 2 cm²/Vs at $n_s = 4 \times 10^{12}$ cm⁻². Curiously, the same trend is observed for bulk conducting STO (see Figure 3).

Similar to the case in bulk STO, the limitations of the room temperature mobility in γ-Al₂O₃/SrTiO₃ was attributed by Christensen et al. to interaction with longitudinal optical phonons, which scatter both the electrons and increases the effective mass by polaron formation [56]. Using the expression for the phonon scattering (see Section 2.2 and Equations 8-10), the electron-phonon coupling ($\alpha$) and the polaron effective mass enhancement ($m_p/m_e$) was calculated from the mobility as presented in Figure 9. From this, it was deduced that the mobility enhancement at high carrier densities is caused by electron screening which weakens the electron-phonon coupling and lowers the effective polaron mass.

4.3 – Mobility at low temperature

Analogous to bulk conducting STO, the low-temperature electron mobility of confined STO-based systems is likewise strongly carrier density dependent. The subsequent subsection will introduce the different confined STO-based systems, previously studied at low temperature, and provide an overview of this dependence.

4.3.1 – δ-doped STO

Through state-of-the-art deposition techniques, it is possible to meticulously engineer the dopant profile in oxides such as STO. This degree of lattice engineering has provided the possibility to realise so-called δ-doped STO where all charge carrier dopants are embedded in a layer with thickness less than 10 nm between two slabs of insulating and nominally un-doped STO. This confinement of the electrons, result in a symmetric quantum well with essentially the same physical properties as observed for electrons confined at oxide heterointerfaces. More importantly, this method, perfected for semiconductors and simple oxides such as ZnO (see Section 5), is a commonly used pathway to augment the electron mobility of a given system. δ-doped STO has been realised experimentally by use of pulsed laser deposition (PLD) [58] and molecular beam epitaxy (MBE) [53,59]. In these previous studies, the dopant used in PLD and MBE growth of δ-doped STO were niobium [58] and lanthanum [59],[53], respectively. Figure 10 shows the available data for electron mobility of δ-doped STO at $T = 2$ K realised by PLD and MBE. It is evident from the figure that the lowest observed electron mobilities of $\mu = 20,000$ cm²/Vs coincide with the highest observed carrier densities at $n_s = 10^{13}$ cm⁻². This behaviour seems at first glance to qualitatively agree with what is the case for bulk conducting STO (see Figure 5). However, the electron mobility at larger carrier densities where $n_s > 10^{14}$ cm⁻² seem not to reduce as rapidly as is the case for bulk STO but rather saturate around a mobility of $\mu = 1000$ cm²/Vs. The origin of the high electron mobility in δ-doped STO was in all previous studies attributed to the absence of background impurities in the δ-doped STO layer due to the purity of the growth sources [53,58,59].
2\text{n} and a strong bluish cathode 

\[ \text{AO/STO} \]

growth at a higher oxygen partial pressure, which reduces the amount of 

The mobility of interface-confined electrons in LAO/STO has been 

Here, a decrease in the growth pressure from 900 °C to 650 °C resulted in 

an increase of the electron mobility from 600 to 8,000 cm²/Vs (see Figure 

The mobility enhancement was found to occur together with an order of 

magnitude decrease in the total carrier density reaching a value of \(5 \times 10^{12} \text{ cm}^{-2} \). Interestingly, the observation of high electron mobilities 

coinciding with low carrier densities appears to be a general trend for ungated 

LAO/STO [63–65] as shown in Figure 13. The same figure also shows that 

surface treatments using conducting atomic force microscopy, solvent 

deposition and heating can be used to enlarge the mobility up to 20,000 

cm²/Vs at a low carrier density of \(1.5 \times 10^{12} \text{ cm}^{-2} \).

Several other pathways for enhancing the electron mobility in LAO/STO 

have been reported in the literature, including electrostatic gating [66,67] 

as well as inserting a La₀.₅SrMnO₃ buffer layer [49] between LAO and STO 

or a SrCuO₂ capping layer [51] on top of LAO/STO during growth. The latter 

two pathways will be discussed further in the next two subsections.

Figure 10 – The low-temperature electron mobility of δ-doped STO is 
gradually decreased as the carrier density is increased. This behaviour 

qualitatively resembles the trend in bulk conducting STO. The figure is 
adapted from reference [53] and the data is obtained from Kozuka et al. 
[58], Jalan et al. [59], Matsubara et al. [53].

4.3.2 – LaAlO₃/SrTiO₃

The active field of confined electron gases in STO-based heterostructures 

was initiated in 2004 when it was found by Ohtomo & Hwang that the 

interface between the insulators LAO and STO could be made conducting 
[5]. Moreover, when the LAO film was deposited at an oxygen partial 

pressure of approximately \(10^{-6} \text{ mbar} \), an electron mobility in excess of 

10,000 cm²/Vs could be realized. This enabled the observation of clear 

Shubnikov-de Haas quantum oscillations of the resistance as a function of 

the magnetic field. The oscillations were observed both when the magnetic 

field was applied perpendicular and parallel to the interface. This 

suggested that the highly mobile electrons obtained by growth at this low 

oxygen partial pressure were not confined to a narrow sheet at the 

interface. The three-dimensional conductivity was in later studies by 

Herranz et al. and Kalabukhov et al. confirmed to be due to oxygen 

vacancies homogeneously distributed in the bulk of STO formed during the 

deposition of LAO [60,61]. This is shown in Figure 11 where the LAO/STO 

heterostructure grown at a low oxygen pressure exhibits a very high sheet 

electron density, a high electron mobility and a strong bluish cathode 

luminescence stemming from oxygen vacancies [61]. The confined nature of 

the electrons can be obtained either by post-annealing in oxygen or by 

growth at a higher oxygen partial pressure, which reduces the amount of 

oxygen vacancies in the bulk of STO. It is still debated whether these 

confined electrons stem from, e.g., residual oxygen vacancies in STO or 

Figure 11 – The Hall carrier density \(n_s\) and mobility \(\mu_H\) as a function of 
temperature for (i) STO vacuum annealed at 800 °C in an oxygen pressure of 
\(10^{-6} \text{ mbar} \), (ii) STO exposed to argon ion bombardment with an energy of 
300 eV and a current density of 0.2 mA/cm² and (iii) LAO/STO deposited 
at 800 °C in an oxygen pressure of \(10^{4}\) or \(10^{6} \text{ mbar} \) with or without a two-

hour postannealing step at 600 °C in 500 mbar oxygen pressure during 
cool-down after the deposition. Reprinted figure with permission from 
[61]. Copyright 2007 by the American Physical Society.
4.3.3 – Amorphous-LaAlO$_3$/LaSrMnO$_3$/SrTiO$_3$

One pathway to increase the electron mobility beyond what typically is observed in LAO/STO of around ~1,000 cm$^2$/Vs is to insert a buffer layer of La$_{1-x}$Sr$_x$MnO$_3$ between LAO and STO [49]. Inserting a La$_{1-x}$Sr$_x$MnO$_3$ layer with a thickness of two unit cells or more leads to insulating interfaces, however, inserting a single unit cell reduces the carrier density and increases the electron mobility up to 70,000 cm$^2$/Vs (see Figure 14). The reduction in the electron density were by Chen et al. attributed to occur by the La$_{1-x}$Sr$_x$MnO$_3$ layer acting as an electron sink thus localizing part of the electrons leaving the residual electrons to populate the conduction band in STO (see Figure 14). The low oxidation potential of La$_{1-x}$Sr$_x$MnO$_3$ was thought by the authors to minimize the oxygen vacancy formation in STO and hence reduce the ionized donor scattering. As the oxygen vacancy concentration in STO were lowered, the conducting electrons were suggested to originate from the LAO layer. This was expected to result in a separation of donors and charge carriers in accordance with the modulation doping strategy, leading to the observed mobility enhancement.

4.3 – Amorphous-LaAlO$_3$/LaSrMnO$_3$/SrTiO$_3$

One pathway to increase the electron mobility beyond what typically is observed in LAO/STO of around ~1,000 cm$^2$/Vs is to insert a buffer layer of La$_{1-x}$Sr$_x$MnO$_3$ between LAO and STO [49]. Inserting a La$_{1-x}$Sr$_x$MnO$_3$ layer with a thickness of two unit cells or more leads to insulating interfaces, however, inserting a single unit cell reduces the carrier density and increases the electron mobility up to 70,000 cm$^2$/Vs (see Figure 14). The reduction in the electron density were by Chen et al. attributed to occur by the La$_{1-x}$Sr$_x$MnO$_3$ layer acting as an electron sink thus localizing part of the electrons leaving the residual electrons to populate the conduction band in STO (see Figure 14). The low oxidation potential of La$_{1-x}$Sr$_x$MnO$_3$ was thought by the authors to minimize the oxygen vacancy formation in STO and hence reduce the ionized donor scattering. As the oxygen vacancy concentration in STO were lowered, the conducting electrons were suggested to originate from the LAO layer. This was expected to result in a separation of donors and charge carriers in accordance with the modulation doping strategy, leading to the observed mobility enhancement.

4.3.3 – Amorphous-LaAlO$_3$/LaSrMnO$_3$/SrTiO$_3$

One pathway to increase the electron mobility beyond what typically is observed in LAO/STO of around ~1,000 cm$^2$/Vs is to insert a buffer layer of La$_{1-x}$Sr$_x$MnO$_3$ between LAO and STO [49]. Inserting a La$_{1-x}$Sr$_x$MnO$_3$ layer with a thickness of two unit cells or more leads to insulating interfaces, however, inserting a single unit cell reduces the carrier density and increases the electron mobility up to 70,000 cm$^2$/Vs (see Figure 14). The reduction in the electron density were by Chen et al. attributed to occur by the La$_{1-x}$Sr$_x$MnO$_3$ layer acting as an electron sink thus localizing part of the electrons leaving the residual electrons to populate the conduction band in STO (see Figure 14). The low oxidation potential of La$_{1-x}$Sr$_x$MnO$_3$ was thought by the authors to minimize the oxygen vacancy formation in STO and hence reduce the ionized donor scattering. As the oxygen vacancy concentration in STO were lowered, the conducting electrons were suggested to originate from the LAO layer. This was expected to result in a separation of donors and charge carriers in accordance with the modulation doping strategy, leading to the observed mobility enhancement.
4.3.4 – SrCuO$_2$/LaAlO$_3$/SrTiO$_3$

It was demonstrated by Huijben et al. that depositing STO with additional top layers after the normal growth of LAO could enhance the electron mobility [51]. Specifically, when a usually grown LAO/STO heterostructure was capped with one unit cell of SrCuO$_2$ and an additional two unit cells of STO, the resulting conducting LAO/STO interface was shown to display enlarged electron mobilities exceeding 50,000 cm$^2$/V s at 2 K (see Figure 15). The authors attributed this large enhancement in electron mobility to a reduction in the amount of oxygen vacancies at the LAO/STO interface that in turn would correspond to a reduced ionized impurity scattering of electrons [51]. The highest observed electron mobility exceeding 50,000 cm$^2$/V s at 2 K was realised when growing the SrCuO$_2$ capped LAO/STO heterostructure at the lowest studied oxygen partial pressure of 10$^{-9}$ mbar.

![Figure 15](image)

Figure 15 – (a) The electron mobility, $\mu_n$, as a function of temperature, T, for SrCuO$_2$ capped LAO/STO heterostructures (filled markers) and identically prepared structures without the SrCuO$_2$ capping layer (open markers). Evidently, inclusion of the SrCuO$_2$ capping layer results in greatly enlarged mobility values. The highest observed electron mobility was realised in the SrTiO$_3$(2u.c.)/SrCuO$_2$(1u.c.)/LaAlO$_3$(10u.c.)/SrTiO$_3$ heterostructure 2(1/10) grown at 10$^{-9}$ mbar. (b) The proposed mechanism for the enhanced interface electron mobility, where the SrCuO$_2$ is thought to prevent the formation of interfacial LAO/STO oxygen vacancies and thereby reduce electron ionized impurity scattering. The figure is adapted from reference [51].

4.3.5 – γ-Al$_2$O$_3$/SrTiO$_3$

The heterostructure composed of spinel γ-Al$_2$O$_3$ epitaxially grown on perovskite STO exhibits the, to date, largest electron mobility of 140,000 cm$^2$/Vs at 2 K observed in any STO-based electronic system [6]. The high electron mobility was observed to peak at a sheet carrier density of around 4×10$^{14}$ cm$^{-2}$ [see Figure 16] in contrast to LAO/STO where the highest mobilities are observed at much lower carrier densities of around ~10$^{12}$ cm$^2$. Interestingly, it was shown by Christensen et al. that a gentle ex-situ annealing at 20-100 °C following growth enhanced the mobility without affecting the carrier density [68]. On the contrary, a harsher annealing at 200 °C was shown to lead to a decrease in both carrier density and electron mobility [56]. Photoemission studies and density functional theory calculations allowed the authors to suggest that the oxygen vacancy donors are more stable at the γ-Al$_2$O$_3$/SrTiO$_3$ interface [68]. Based on this, it was proposed that the high mobility is caused by a lowering of the electron-donor scattering from having oxygen vacancies preferentially located at the interface with the electrons extending deeper into STO (see Figure 16).

![Figure 16](image)

Figure 16 – (a) Electron mobility ($\mu$) as a function of carrier density ($n_s$) for γ-Al$_2$O$_3$/SrTiO$_3$ heterostructures with varying carrier densities obtained by varying the deposition conditions or performing post-annealing in 1 bar oxygen at 200 °C. Reprinted figure with permission from [56]. Copyright 2018 by the American Physical Society. (b) Schematic illustration of the proposed mechanism for the high-mobility in which a lowering of the electron-donor scattering occurs as oxygen vacancies primarily are thought to locate at the interface whereas electrons extend deeper into STO. Reprinted figure with permission from [68]. Copyright 2017 by the American Physical Society.
4.3.6 – Overview of low-temperature electron mobility

The previous Subsections 4.3.1–4.3.5 have provided a representative overview of the available literature for confined STO-based electronic systems with high mobility at low temperatures. Herein, the origin of enhanced electron mobility in the different systems was presented as suggested by the authors in the original papers. We have compiled the data into an overview of the low-temperature electron mobility as a function of carrier density for both bulk conducting STO and confined STO-based systems (Figure 17). The figure also includes the scattering model presented in Section 3.3. Note that in order to plot the electron mobility data for bulk conducting STO together with the confined electron systems, the three-dimensional carrier density was translated into the equivalent two-dimensional density following the expression derived by Trier et al. [65]. At first glance, it is clear from Figure 17 that some confined systems seem to follow a similar trend in $\mu(n_s)$ as bulk conducting STO over two orders of carrier densities for $10^{12}$ cm$^{-2} < n_s < 10^{14}$ cm$^{-2}$. In particular, only δ-doped STO and γ-Al$_2$O$_3$/SrTiO$_3$ with carrier densities larger than $n_s > 10^{14}$ cm$^{-2}$ falls markedly outside this overall electron mobility trend. At lower carrier densities, the observed electron mobilities of both these systems seem to merge with the trend of the other confined STO-based systems in addition to the trend of bulk STO. Although a few reported electron mobility values for STO-based systems at low carrier density significantly differ from the remaining data points, most electron mobility values seem to follow a common trend. In order to fully understand why the few outlying data points are falling outside this common trend, further investigations are needed to clarify the mechanism of their reduced scattering rates.

Since there, nonetheless, seem to exist a common behaviour of electron mobility between most confined systems and bulk STO, it points towards that the same scattering mechanisms in bulk STO are governing low-temperature transport in confined STO-systems. In the low carrier density regime of $n_s < 10^{12}$, this suggest that the electron mobility in confined STO-systems overall is limited by scattering with ionized background impurities, i.e. the inherent purity and crystal quality of STO. In the large carrier density regime of $n_s > 10^{13}$ the electron mobility reduction seems consistent with increased ionized donor scattering. In bulk STO these additional ionic scattering centres are formed by doping STO with e.g. oxygen vacancies or lanthanum. Therefore, the reduction of electron mobility observed for the confined STO-based systems having $n_s \sim 10^{13}$–$10^{14}$, with the exception of δ-doped STO and γ-Al$_2$O$_3$/SrTiO$_3$, may be consistent with the dominant scattering mechanism being due to ionized donors. For some confined STO-based systems, this behaviour seems in agreement with how the authors in the original papers attributed the doping mechanism. In particular, confined STO-based systems where charge carriers are expected to originate from oxygen vacancies in STO is in good agreement with their trend in electron mobility. In these cases, control of the location and amount of the oxygen vacancy donors are generally believed to be essential for obtaining a high mobility. Conversely, charge carriers in LAO/STO are often attributed to arise due to the polarity discontinuity between LAO and STO within the so-called polar catastrophe model ([62,69]). Although this model is expected to provide electrons to the LAO/STO interface, these electrons are not hypothesised to lead to ionized donors located in STO, but rather from the LAO surface similar to modulation doping. Therefore, the electron mobility trend of LAO/STO observed in Figure 17 therefore brings forth an interesting perspective on the polar catastrophe model as discussed by Trier et al. [65].

Concerning δ-doped STO and γ-Al$_2$O$_3$/SrTiO$_3$ there seem to be other origins for the scattering mechanism once the carrier density becomes large with $n_s > 10^{14}$. However as presented in Subsection 4.3.5, recent transport, photo-electron spectroscopy and density functional studies of γ-Al$_2$O$_3$/SrTiO$_3$ have shed some light on the ionized donor distribution at this particular interface [68]. Specifically, the authors came to the conclusion that oxygen vacancies in γ-Al$_2$O$_3$/SrTiO$_3$ were highly concentrated in STO right at the interface [68]. Whereas for δ-doped STO there are less concrete suggestions in the literature why the electron mobility does not follow the remaining confined STO-based systems having $n_s > 10^{14}$ and should be the subject of further investigations.

Figure 17 – The dependence of the low-temperature electron mobility, $\mu$, on the sheet carrier density, $n_s$, as well as corresponding bulk carrier density, $n_{3D}$, for bulk conducting STO and confined STO-based systems. The scattering model is obtained by interpolating between the three different scattering regimes described in Section 3.3. The data is obtained from: γ-Al$_2$O$_3$/SrTiO$_3$ (GAO/STO) [6]; amorphous-LaAlO$_3$/LaSrMnO$_3$/SrTiO$_3$ (δ-LAO/LSM/STO) [49]; CaZrO$_3$/SrTiO$_3$ (CZO/STO) [50]; SrCuO$_2$/LaAlO$_3$/SrTiO$_3$ (SCO/STO) [51]; bulk SrTiO$_3$ (STO) [23,25,18,40,69,70]; δ-doped SrTiO$_3$ (δ-doped STO) [53]; LaAlO$_3$/SrTiO$_3$ (LAO/STO) [63,64]; ion liquid-gated LaAlO$_3$/SrTiO$_3$ (liquid-gated LAO/STO) [67]; amorphous-LaAlO$_3$/SrTiO$_3$ (δ-LAO/STO) [55].
4.4 – One-dimensional confined conductivity

The previous subsections 4.1-4.3 reviewed the literature concerning electron mobility for two-dimensionally confined STO-based heterointerfaces. Confining electrons in one dimension rather than the two-dimensional interfacial area seems to have a profound impact on the measured electron mobility as studied by Irvin et al. [70] In that study, LAO was deposited on STO at two different deposition conditions using a LAO thickness below the critical thickness for obtaining metallic conductivity. When a positively charged c-AFM tip was scanned at the surface of the insulating LAO/STO heterostructure, conductivity was formed locally underneath the tip (see Figure 18). In this way, the authors were able to write conducting lines and devices with nanoscale dimensions in a highly customizable way. When conducting squares with a side length ranging from 20 nm to 1 μm were measured in the van der Pauw (vdP) geometry, the room temperature mobility was found to be on the order of 6-7 cm²/Vs (see Figure 18). This is consistent with the mobilities of unconfined and two-dimensional confined STO-based systems as discussed in the previous subsections 3.2 and 4.2. However, when conductivity was formed in the shape of 1.2-8 μm long Hall-bar, the mobility was found to increase significantly for Hall bar devices with widths below 100 nm. The highest mobility of 350 cm²/Vs was obtained when the Hall bar width was 10 nm. This mobility is astoundingly 60 times larger than what typically is observed in STO.

Cooling the Hall-bar structures also increased the mobility beyond what is usually observed in LAO/STO heterostructures, reaching a value exceeding 20,000 cm²/Vs at 20 K (see Figure 19). Curiously, deviations from the typical $\mu \propto T^{-2}$ behaviour were also observed (see Subsections 3.1 and 4.1). Although the origin of this mobility enhancement needs further investigation, the authors suggested that it is due a long-range coherence phenomenon occurring when electrons become confined in a quasi one-dimensional structure.

Figure 19 – Mobility (\(\mu\)) as a function of temperature for two-dimensional confined conductivity at the LAO/STO interface as well as for conductivity formed in two Hall bar devices (H5 and H6) with a width of 10 nm. Reprinted with permission from reference [70]. Copyright 2013 American Chemical Society.

Section 5: Mobility of Zinc Oxide: Bulk and thin-films

5.1 – Zinc Oxide (ZnO)

Zinc oxide (ZnO) is a semiconductor compound, which have attracted considerable interest in recent years. ZnO belongs to the class of II–VI semiconductors having the most stable B4 (wurtzite) structure. Moreover, has ZnO a direct band gap of 3.4 eV and a high exciton binding energy of 60 meV, which is much higher than that of e.g. GaN (21–25 meV). Therefore, devices made from ZnO can operate more efficiently at high temperatures making it highly desirable for optoelectronic devices, such as light-emitting diodes, UV sources/sensors and room temperature lasing applications. The physical properties of ZnO are given in Table 3 [71].
There is growing interest for the possibility of realizing active electronic devices exclusively made from transparent oxide materials, the so-called “transparent electronics” [76]. The discovery in 1997 of the oxide CuAlO₂ with a high p-type (hole) conductivity [77] opened up for the first time for the possibility of combining transparent semiconducting n-type oxides with its p-type oxide counterpart. Having both n- and p-type oxides opened the door for a wide range of possible applications such as p-n junction diodes and UV-emitters. Another key device for realizing transparent circuits is the transparent field-effect transistor (TFET). For this application, one of the central challenges in the development of TFETs is getting a charge carrier channel with sufficiently high mobility. Once developed, these TFETs could in turn potentially be utilized in both existing display technologies and future integrated circuits [78]. When planning a thin film transistor layout and considering possible active layer material, amorphous semiconductors are generally preferred over their polycrystalline counterparts owing to their low processing temperature and uniformity of device characteristics. For example, when comparing the carrier mobility of e.g. amorphous-Si:H (µ ~ 1 cm²/Vs) with the crystalline one (µ ~ 200 cm²/Vs), it is clear that the mobility of the amorphous sample is still significantly lower. The low mobility is associated with the intrinsic nature of the difference in the chemical bonding. In contrast, a degenerate band conduction with relative large carrier mobility values (µ > 10 cm²/Vs) is possible in amorphous oxide semiconductors containing post-transition-metal cations. The electron transport in these materials is radically different from those of the covalent semiconductors. The conduction band bottom in oxide semiconductors is usually originating from hybridization between neighbouring atomic orbitals, which have direct overlap between them. Within these materials, doped ZnO plays a major role due to the quality of ZnO single-crystal substrates and films that one can achieve today. In these type of materials, a transparent amorphous oxide semiconductor from the ZnO system acting as the active channel in transparent thin-film transistors (TTFTs) show a relative high carrier mobility [79]. ZnO is also one of the systems which provide the exciting possibility of making thin-film transistors on flexible substrates with relatively high electron mobilities when compared with amorphous silicon or organic semiconductors [80]. In the following, we will review the carrier mobility of electrons in bulk conducting ZnO followed by discussion on the mobility of its derivatives. Unlike STO, thin films of ZnO plays an important technological role and we have therefore devoted a part of Section 5 on

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<tr>
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</tr>
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</table>

Table 3 – The physical parameters of ZnO adopted from the literature or the periodic table (PT).

For many applications, such as light-emitting diodes, it is necessary to obtain both the n-type and the p-type materials. Realizing bulk conducting p-type ZnO, however, still poses a challenge, which remains a major hindrance for future applications. The p-type doping is difficult to achieve in ZnO [89] due to the existence of background n-type dopants including H

![Figure 20](image-url)
impurities [90], O vacancies [91] and Zn interstitials [92]. Despite the difficulties in obtaining bulk p-type ZnO, several laboratories have demonstrated thin film p-type ZnO using group V (N [93], P [94], As [95], and Sb [96]) and group I (Li [97]) elements. Co-doping with two potential acceptors (N and As) [98] or acceptor and donor (N and Al) [99] have also been studied. The room temperature hole mobility of these epitaxial thin films was slightly higher than 7.7 cm$^2$/Vs and achieved by careful controlling of the residual impurity densities. Although this hole mobility is not as high as in p-type GaN it still represents a significant advantage towards utilizing ZnO in future applications. The hole transport characteristics are calculated using the “relaxation time approximation” as a function of temperature [19]. As mentioned in Section 2.2.2, the acoustic phonon scattering is the most important mechanisms in p-type ZnO, limiting the hole mobility over a wide range of temperature. For temperatures above 300 K, the polar optical phonon scattering mechanism is the dominant factor (see Figure 21).

![Graph showing theoretical scattering model for electron mobility of p-type ZnO thin films.](image)

Figure 21 – Theoretical scattering model for electron mobility of p-type ZnO thin films. Here, the different scattering contributions in ZnO are: acoustic (dp) and optical phonons (npop), polar optical phonon (pop), piezoelectric deformations (pe). The figure is adapted from reference [19]. Copyright 2006 The Japan Society of Applied Physics.

5.3 – Carrier mobility in ZnO based thin films

The most widely studied transparent conductive oxide (TCO) thin film material is Sn-doped indium oxide (ITO), where carrier densities are exceeding 10$^{21}$ cm$^{-3}$ and resistivities are below 10$^3$ Ω cm. However, the high cost and scarcity of indium has stimulated the search for a replacement material. Recent work has therefore been concentrated on finding alternative materials including promising candidates within the binary oxides ZnO, SnO$_2$ [100], CdO [101], and Ga$_2$O$_3$ [102], and within multi-component oxides such as indium–gallium–zinc–oxide (IGZO) [103] and cubic spinel oxide compounds such as CdIn$_2$O$_4$ [104] and SnZn$_2$O$_4$ [105]. Such alternatives have been reviewed by Ellmer [106]. Here, obtaining a high carrier mobility at room temperature is crucial as it allows for a large conductivity while not sacrificing the transparency by adding additional dopants.

To date, the highest room temperature mobility observed for ZnO was in a bulk single crystal with values approaching $\mu \sim 200$ cm$^2$/Vs and carrier density $n \sim 1 \times 10^{17}$ cm$^{-3}$. Doping thin film ZnO with either Al [107] or Ga [108], was found to increase the carrier density while the resistivity reduced to $10^{-4}$ Ωcm. Introducing other potential dopants into the ZnO thin film have previously been studied for e.g. In-doped ZnO (IZO) [109] and In- and Ga-doped ZnO (GZO) [110]. The Al-doped ZnO (AZO) has been considered to be one of the best candidates as a transparent conductor due to the relative low cost, the natural abundance of Zn and Al as well as the good optical transmission of this compound in the visible range. However, these materials are still facing the following challenges [103]: (1) their room temperature resistivity still remains high, and (2) the transport properties are strongly dependent on film thickness. For these films the additional dopants furthermore reduce the mobility to a relatively low range of $\mu \sim 30-50$ cm$^2$/Vs for a carrier density of around $n \sim 10^{20}$ cm$^{-3}$, primarily because of ionized impurity scattering (see Figure 22). In Figure 22 we show a representative collection of different compositions with their respective mobility and carrier densities measured at room temperature. Here, the carrier density and mobility at room temperature are between $n \sim 10^{14}-10^{18}$ cm$^{-3}$ and $\mu \sim 70-150$ cm$^2$/Vs, respectively.

![Graph showing dependence of room temperature electron mobility, $\mu$, on the corresponding bulk carrier density, $n_{3D}$, for bulk conducting ZnO-based systems.](image)

Figure 22 – The dependence of the room temperature electron mobility, $\mu$, on the corresponding bulk carrier density, $n_{3D}$, for bulk conducting ZnO-based systems. The data is obtained from: Al-doped ZnO (AZO)/Glass [111]; In- and Ga-codoped ZnO (IGZO) on yttrium-stabilised zirconium (YSZ) [79]; Ga-doped ZnO (GZO)/ZnO [112]; ZnO/SCAM [109]; ZnO/Sapphire [109] Ga-doped ZnO/Sapphire (Ga-ZnO/Sapphire) [108].

While traditional TCOs are often highly crystalline, a new class of materials, amorphous indium-doped ZnO (a-IZO) with an unusually high electron mobility, $\mu \sim 54$ cm$^2$/Vs and carrier density of $n \sim 1.3 \times 10^{20}$ cm$^{-3}$ was recently reported by Leenheer et al. [111]. A general observation of these type of materials is that the mobility seems to depend on the carrier density, which in turn can be tuned by annealing in oxygen.
Section 6: Mobility of Zinc Oxide: Two-dimensional electron gases

6.1 – Mobility in ZnO based heterostructures

Much effort has been made in designing two-dimensional electron gases in ZnO-based systems. The first report on high carrier mobility at the (0001)-oriented Mg$_2$Zn$_{1-x}$O/ZnO heterostructures was realized in 2007 using Pulsed Laser Deposition (PLD) by Tsukazaki et al. [11]. The Mg$_2$Zn$_{1-x}$O layer in their heterostructure acts as a potential barrier for the two-dimensional electron gas (2DEG) in the adjacent ZnO layer by gradually adjusting the Mg content. In this study, the authors found that the carrier density and the mobility depended on the amount of magnesium doping (i.e. $x$). The authors demonstrated sheet carrier densities of $n_s \sim 0.66 \times 10^{12}$ and $3.7 \times 10^{12}$ cm$^{-2}$ with electron mobilities of $\mu \sim 5500$ and 2700 cm$^2$/Vs at 1 K for $x = 0.15$ and $x = 0.2$, respectively. Although PLD provided the initial means to achieve a high interfacial mobility, molecular-beam epitaxy (MBE) now serves as the thin-film growth method of choice to achieve exceptional high purity and cleanliness of MgZnO interfaces. Soon after the initial results obtained with PLD, Tsukazaki et al. showed that electron mobilities higher than the one obtained by PLD could be attained in Mg$_{x}$Zn$_{1-x}$O/ZnO heterostructures grown by MBE [112] i.e. electron mobility in the order of 14,000 cm$^2$/Vs. The doubling of the electron mobility of samples obtained by MBE as compared to PLD grown sample was attributed to the growth direction and strain [112]. Further improvement of the growth conditions resulted in even higher mobility values of MgZnO/ZnO heterostructures grown by MBE with an electron mobility exceeding 180,000 cm$^2$/Vs [113]. This high electron mobility allowed observation of the fractional quantum Hall effect for the first time in oxides. The observation underlined the cleanliness and extremely smoothness of the MgZnO/ZnO interface as they both are prerequisites for the observation of this effect.

Impressively, this was the first demonstration of fractional quantum Hall effect in an ionic material with much stronger correlation between electrons compared to conventional semiconductor heterostructures [113]. Further improvements of the electron mobility to a value of 300,000 cm$^2$/Vs and a magnetic field up to 26 T at 40 mK made it possible to further study lower filling factor fractional quantum states in oxide materials [114]. This was due to a reduced charge carrier density compared to the samples studied earlier [113]. Additional increase of the mobility to a value of 700,000 cm$^2$/Vs for the Mg$_{0.07}$Zn$_{0.93}$O/ZnO heterostructures grown by MBE was achieved by a reduction of impurity levels associated with the use of pure distilled ozone and avoiding interface roughness scattering [115]. The dependence of electron mobility with 2DEG sheet carrier density measured at temperatures below 2 K is shown in Figure 23.

The electron mobility values shown in Figure 23 represent the highest reported mobility values for different ZnO-based heterostructures. The continuous improvement in the mobility over the years stems from the refinement of the MBE growth conditions while using ozone to reduce the residual impurity content. The electron mobility improvement is, however, also a result of carefully controlling the carrier density with a lower Mg content of samples. Care was likewise taken to prepare single epitaxial layers using MBE on a ZnO single crystal substrates (Fabricated by hydrothermal method), tuning the Mg content and improving the purity of the Zn source material. In addition, the use of ozone as an oxygen source offers the benefit of an expanded temperature window for uniform growth. More details of the preparation method can be found in reference [7]. This above mentioned careful procedure enabled a record-high electron mobility exceeding a staggering 1,000,000 cm$^2$/Vs [7]. In general, these high electron mobilities occur only for a narrow range of charge densities. Figure 24 show the evolution of electron mobility improvements as a function of sheet carrier density over the years. It is evident from Figure 24 that although the electron density continuously was improved, two distinct scattering mechanisms are limiting the electron mobility depending on the carrier density regime. For low carrier densities, charged impurity scattering is dominating ($\mu \propto n$, see Section 2.1) while interface/ alloy scattering limits the electron mobility at higher electron densities ($\mu \propto n^{-3/2}$, see Section 2.4).

![Figure 23 - The dependence of the low-temperature electron mobility, $\mu$, on the sheet carrier density, $n_s$, for confined ZnO-based systems. The data is obtained from: Mg$_{0.15}$Zn$_{0.85}$O/ZnO, PLD, Tsukazaki, et al. [11]; Mg$_{0.09}$Zn$_{0.91}$O/ZnO, MBE, Tsukazaki et al. [112]; MgZnO/ZnO, MBE, Tsukazaki et al. [113]; MgZnO/ZnO, MBE, Kozuka et al. [114]; Mg$_{0.07}$Zn$_{0.93}$O/ZnO, MBE, Falson et al. [115]; ZnMgO/ZnO, Radical source MBE, Tampo et al. [116]; MgZnO/ZnO, Ozon-molecular MBE, Falson et al. [7].](image-url)
Section 7: Beyond strontium titanate and Zinc Oxide

Although STO and ZnO exhibit the largest electron mobilities in oxides at low temperature, a large number of oxide structures beyond these have a comparable or even larger mobility at room temperature. The room temperature electron mobility versus carrier density for thin films of key oxide compounds such as SnO₂ [118], TiO₂ [119], ZnO [120], Ga₂O₃ [121], In₂O₃ [122], CdO [123], BaSnO₃ [124] and SrRuO₃ [125] are summarized in Figure 25.

For comparison, we have included additional data for traditional high mobility semiconductors such as GaN [126] and GaAs [127]. Another type of perovskite oxides that recently have gained attention is the Alkaline-earth stannates with the generic composition ASnO₃ (A = Ca, Sr, and Ba). Among ASnO₃, the cubic BaSnO₃, which is a n-type semiconductor stable at temperatures up to 1000 °C, has been intensively investigated [128]. Earlier work showed that the electron mobility of thin film and polycrystalline doped BaSnO₃ is relatively low (µ < 1 cm²/Vs) [128]. However, BaSnO₃ single crystals can now display electron mobilities of µ ~ 320 cm²/Vs [124] at room temperature with a doping level of 8×10¹⁹ cm⁻³. Therefore, there has been high demand for the growth of high-quality single crystals of doped BaSnO₃. Raghavan et al. reported MBE-grown (Ba,La)SnO₃ thin films on PrScO₃(110) substrates with room temperature mobilities of µ ~ 150 cm²/Vs by use of preoxidized Sn as a precursor [129]. This is the highest-known electron mobility in doped BaSnO₃ thin films at room temperature. The previously reported mobility values of BaSnO₃ and doped BaSnO₃ on different substrates is presented in Figure 26. In addition, the mobility was enhanced to around 300 cm²/Vs at 50 K by varying the carrier density in an electric-double-layer setup where an electric field was applied to a BaSnO₃ thin film through an ionic liquid [130]. A more comprehensive review on the high electron mobility of doped BaSnO₃ for TCOs can be found in reference [131].

Figure 24 – The improvements in electron mobility (µ) over the years as function of sheet carrier density (n) for temperatures T ≤ 2 K. To guide the eye the mobility scaled by µ ∝ n and µ ∝ n⁻¹/β are indicated at each data set corresponding to charged impurity scattering and interface/alloy scattering, respectively. The figure shows the evolution of µ over the years at 2007 [11], 2009 [117], 2010 [113], 2011 [115] and 2016 [7]. The figure is adapted from reference [7].

Figure 25 – The dependence of room-temperature electron mobility, µ, on the carrier density, n, for different bulk conducting oxide-based systems either as a single crystal or as a thin film. The data is taken from: SnO₂ [118], TiO₂ [119], ZnO [120], Ga₂O₃ [121], In₂O₃ [122], CdO:Ga,In [123], GaN [126], GaAs [127], BaSnO₃ [124] and SrRuO₃ [125].
Figure 26 – The dependence of room-temperature ($T = 300K$) electron mobility, $\mu$, on the carrier density, $n$, for La-doped BaSnO$_3$. Republished with permission of Annual Reviews, from reference [131]; permission conveyed through Copyright Clearance Center, Inc.

7.1 – Confined systems beyond STO- and ZnO-based heterostructures

Integrating two-dimensional oxide electron gases with semiconductors opens prospects of new functionalities arising from the potential of coupling the 2DEG carriers with the semiconductor. This was studied by Kornblum et al. where the authors prepared (001) GdTiO$_3$/SrTiO$_3$ (GTO/STO) heterostructures on (001) GaAs[132]. The sheet carrier density seems to be independent on the temperature with a carrier density value of $\sim 4 \times 10^{14}$ cm$^{-2}$ and electron mobility of $\sim 4$ cm$^2$/Vs from 60 K to 300 K.

Another successful demonstration is the formation of 2DEG at the interface between a-LaO and TiO$_2$ (Anatase) as well as TiO$_2$ (Rutile) substrates [133]. The TiO$_2$ (R)-based heterostructure presents a comparable carrier density to the intensively investigated system of a-LaO/STO, but with a much lower mobility, i.e. $\sim 77$ cm$^2$/Vs at 30K, and in contrast, the a-LaO/TiO$_2$ (R) gives rise to a low mobility of $\sim 24.5$ cm$^2$/Vs at 2K. The authors of this work concluded that the interfacial conduction comes from redox reactions, and that the differences between the materials systems result mainly from variations in the activation energies for the diffusion of oxygen vacancies at substrate surfaces.

In another recent work, a new 2DEG system at the interface between a Mott insulator, LaTiO$_3$, and a band insulator, KTaO$_3$ was reported [134]. A metallic conductive interface was observed from 2 K to 300 K. The mobility of these interfaces seems to be independent of the thickness of LaTiO$_3$ and found to be $\sim 10$ cm$^2$/Vs at 300 K and $\sim 100$ cm$^2$/Vs at 2 K. This study provides alternative oxide heterostructures that host 2DEGs.

There have also been theoretical studies for potential confined electronic systems where the conduction bands are comprised of highly dispersive non-degenerate $s$ orbitals. In a recent work, first-principles density functional theory calculations were used on oxide materials with $s$-like conduction bands [135]. Here, the interface studied was between nonpolar perovskite BaSnO$_3$ and polar perovskite LaScO$_3$ stacked along the [001] direction. BaSnO$_3$ and LaScO$_3$ single crystals are perovskite with band gaps of $\sim 3$ eV and 5.7 eV, respectively [136]. The conduction band electrons were found to reside in the highly dispersive Sn-$s$ orbitals, which have a large band width and a low effective mass. The result of the calculation indicate that the predicted 2DEG is expected to be highly mobile even at room temperature due to the reduced electron-phonon scattering via the inter-band scattering channel [135].

In another theoretical work, Wang et al. attempted to substitute STO with other nonpolar perovskite oxide whose lowest energy conduction bands are comprised of $s$ or $p$ orbitals [136]. The authors used first-principles electronic structure calculations and explored the possibility of producing a high-mobility 2DEG in the following two systems: LaAlO$_3$/SrGeO$_3$ and LaGaO$_3$/BaSnO$_3$ i.e. with SrGeO$_3$ and BaSnO$_3$ as substrate materials [136]. Under the assumption of constant scattering time approximation, the calculations indicated that the room temperature electron mobility and the electrical conductivity of LaAlO$_3$/SrGeO$_3$ and LaGaO$_3$/BaSnO$_3$ systems should be nearly 2.5 and 2.0 times larger, respectively, than in the LAO/STO 2DEG [136]. These theoretical predictions are however yet to be demonstrated experimentally.

Section 8: Perspectives

The majority of oxides still suffers from a low carrier mobility due to structural imperfections and a large number of impurities. However, the recent progress in achieving high carrier mobility in oxides and oxide heterostructures has led to improvements in, e.g., transparent conductors and realization of quantum phenomena in oxides. In particular, material systems based on STO and ZnO have undergone a large development and now reach electron mobilities exceeding 100,000 and 1,000,000 cm$^2$/Vs, respectively. Such high electron mobilities open up various avenues:

1) Many oxides exhibit a large range of remarkable properties. It is of high interest to achieve new functionalities by combining these properties with the quantum phenomena that can be observed in high-mobility materials.

2) Many properties in functional oxides stem from intentional or unintentional defects, but the properties can be hard to control and reproduce as the purity of oxides generally falls much below that of silicon. In the process of achieving high carrier mobilities, unintentional impurities are frequently reduced drastically, which allows for a better control by introducing intentional impurities in a controlled manner into a clean system.

3) Oxides are promising for various applications, which benefit from a high carrier mobility. These include thermoelectricity due to the chemical stability and low price tag of oxides as well as use as transparent conductors where the high band gap in oxides assures transparency in the visible range. Oxides may also be used in, e.g., magneto-resistive devices where the large positive magnetoresistance intrinsically found in some oxides may be combined with a positive magnetoresistance stemming from the extraordinary magnetoresistance [56].

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