

The electronic structure of Cu⁺, Ag⁺, and Au⁺ zeolites

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A variety of procedures have been used to prepare d¹⁰-zeolite materials. The electronic structure of these materials can be regarded to a first approximation as a superposition of the framework, of the charge compensating ions, of solvent molecules and of guest species. Zeolite oxygen to d¹⁰-ion charge transfer transitions dominate the electronic spectra if the ions coordinate to the zeolite oxygens. Specific

coordination sites can influence the energy and the intensity of these transitions remarkably. Intra guest transitions dominate in quantum dot materials, as discussed in detail for luminescent Ag₂S zeolite A. The zeolite is not needed for the photocatalytic water oxidation on Ag⁺/AgCl photo anodes with visible light. It can, however, be used to increase the active surface area substantially.

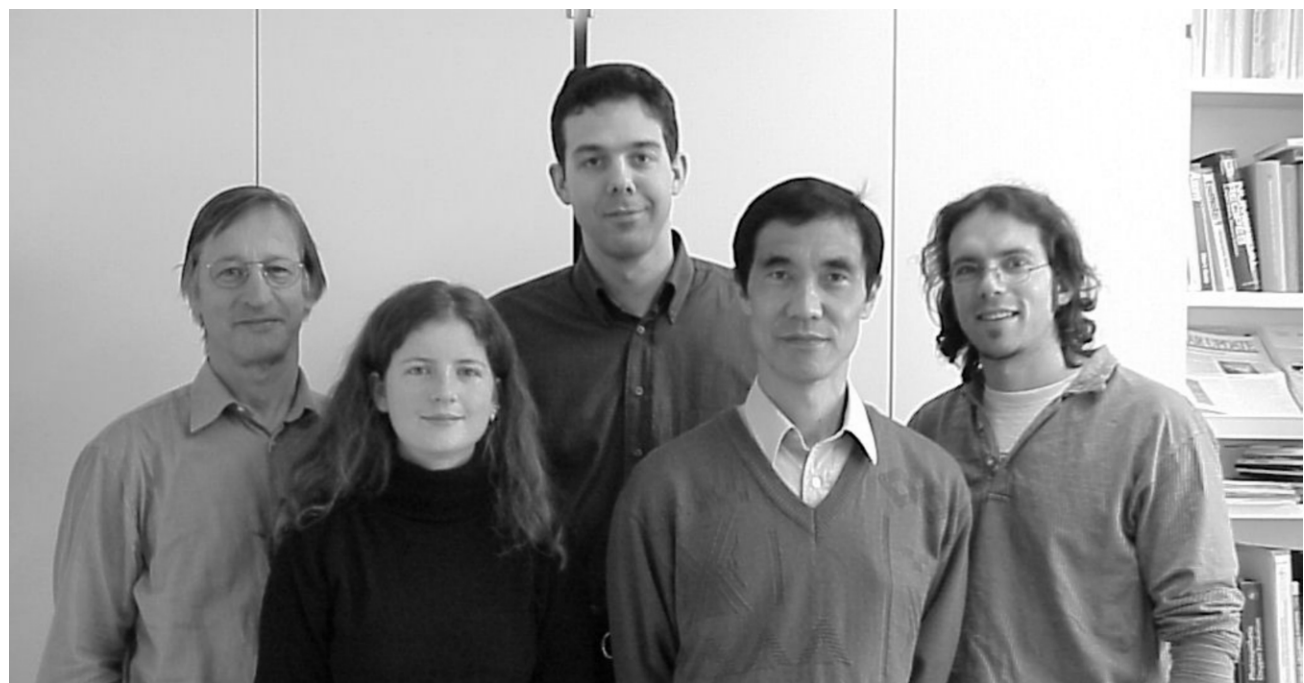
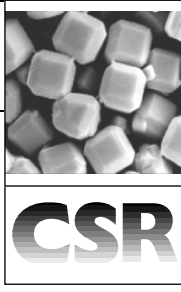
Gion Calzaferri (left) received his PhD degree at the University of Fribourg, Switzerland, 1971. He moved to the University of Berne, after a postdoc period at the University of Basel and physical-chemical research in connection with pharmacy at Ciba-Geigy Basel, Switzerland. He has been professor of physical chemistry in the Department of Chemistry and Biochemistry since 1988. His current research interests concern supramolecularly organized molecules, clusters and complexes in zeolite cavities, dye-zeolite materials as photonic antenna systems, and photochemical transformation and storage of solar energy.

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1 Introduction

The importance of host–guest materials in technology and environmental science can hardly be overemphasized. Basic research on such materials has been less attractive, probably because of the difficulties encountered in rigorous structure determination. Similar difficulties occur in some branches of nanoscience, which have attracted much fundamental research and also put new emphasis on host–guest materials. We are now learning how to describe the often fascinating properties of these materials in a more rigorous way and how to carry out experiments so that artifacts can be avoided.

Zeolites are appealing host materials because they provide a large variety of well defined cavities and channels so that structural identification can concentrate on the shape and position of the guest, provided that care is taken that the framework of the host has not been affected by the synthesis procedure. This is usually easy to check by powder X-ray and vibrational spectroscopy. In this article we focus on the zeolites LTA, LTL, and FAU shown in Fig. 1. These zeolites are crystalline aluminosilicates with cavity and channel structures. Their lattice can be regarded as an enormous crystalline polyanion which contains cations for charge compensation.¹

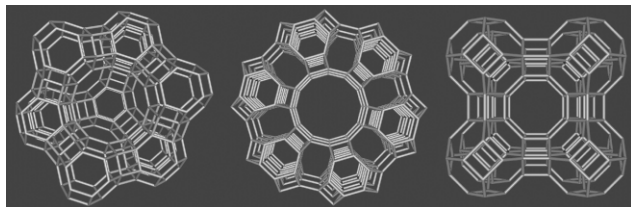


Fig. 1 Framework of zeolite Y (left), zeolite L (middle), and zeolite A (right). The bridging oxygens are not shown.

The electronic structure of a typical zeolite material can be regarded as a superposition of the electronic structure of the framework, of the charge compensating cations, of the solvent molecules and of the guest species. The band gap of about 7 eV and the HOMO position at about –10.7 eV of the zeolite framework are similar to those of α -quartz, despite the fact that the material is less dense.^{2–4}

The charge compensating cations cause new electronic states, some of which may lie within the band gap region. They can interact with each other depending on their nature and the mean distance between them. The presence of solvent molecules, usually water, influences their interaction with the zeolite framework. The electronic structure of the zeolite framework and the charge compensating cations are not influenced by each other, as long as the solvation shell shields the cations from coordinating to zeolite oxygens. A partial removal of the solvent causes incomplete saturation of the coordination sites of the cations which, hence, try to compensate this by sticking closer to the zeolite oxygens. The involved states can be perturbed considerably which causes a change of the optical properties. Apart from charge compensating cations, guests can be present in neutral form, as molecules or as quantum dots or they can be occluded in anionic form. In these cases the electronic structure can be regarded in general as a superposition of the guest and the host with no or only little electronic interaction. However, vibrational interactions can be substantial. This is the underlying general understanding which will guide us in the description of d^{10} -ion zeolites. We concentrate on Cu^{+} , Ag^{+} , and Au^{+} zeolite materials and add only a few remarks concerning other d^{10} -zeolites, *e.g.* such containing Zn^{2+} , Cd^{2+} , and Hg^{2+} ions.

During the last few years the physical and chemical properties of Cu^{+} –zeolites have attracted some interest in the field of catalysis and of physical adsorption or chemisorption of small molecules. For example Cu^{+} –Y can be applied not only as

a catalyst in organic synthesis,⁵ but also for the conversion of NO_x into N_2 and O_2 ,⁶ and for separation procedures like diene/olefin separation.⁷ Ag^{+} –Y was tested for the same separation procedures,⁸ but it exhibited lower separation factors. Ag^{+} -containing zeolites were found to oxidize water in a photochemical reaction,^{9,10} however, the role of the zeolite in these experiments remained obscure. $Ag(CN)_2^-/Au(CN)_2^-$ doped zeolite A was studied as a catalyst for the photochemical decomposition of carbaryl pesticide.¹¹ The understanding of the electronic structure of Ag^{+} –A,^{12,13} recently led to the discovery of the first luminescent silver sulfide quantum dots in zeolite A.¹⁴

Knowledge of the coordination sites of metal cations in zeolites is important for the understanding of these materials. They have therefore been studied by X-ray diffraction methods,^{1,15} by gas adsorption,¹⁶ by different kinds of spectroscopy,¹⁷ and by theoretical means.¹⁸ We show them for the zeolites discussed in this article in Fig. 2, where we have also added SEM pictures to illustrate the morphology of typical materials used in our laboratory.

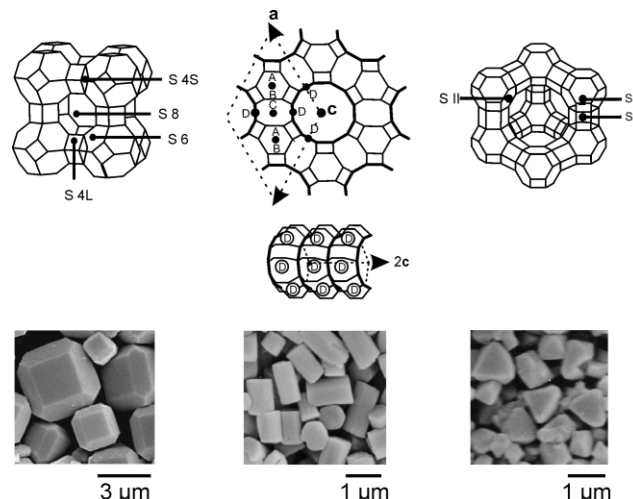


Fig. 2 Frameworks of zeolites A, L, and Y showing the crystallographically identified cation positions and SEM pictures of the corresponding zeolite crystals.

The importance of the occupation of a specific site by a cation has been impressively demonstrated for zeolite A, which possesses 8 S6, 3 S8, 6 S4L, and 12 S4S sites. For Ag^{+} –zeolite A it was proven that only if a silver ion sits on one of the S4L positions, does the zeolite turn to deep yellow under evacuation at room temperature,^{12,13} a reversible color change that was first reported by Rálek *et al.* in 1962,¹⁹ but was misinterpreted for many years. A classical experiment concerns the occupancy of site S8 by means of argon adsorption experiments, which give access to the pore diameter and the inner volume.^{1,20} We show the result of an Ar adsorption experiment at 87 K for a Ca_6A -, a $Ag_{12}A$ -, and a $Na_{12}A$ zeolite in Fig. 3. Ca_6A shows a Langmuir type isotherm, so called type I isotherm. The inner volume of $Na_{12}A$ and $Ag_{12}A$ is, however, not accessible for Ar, because the cation positioned at the S8 ring blocks the entrance of the α -cage.

2 Preparation of d^{10} -zeolite materials

A variety of procedures have been used to prepare d^{10} -zeolite materials. One procedure is ion exchange in some cases followed by selective reduction or other selective reactions. An example is the ion exchange with Cu^{2+} from an aqueous solution, followed by selective reduction with H_2 or CO .²¹ A different procedure is the insertion of molecules or ion pairs

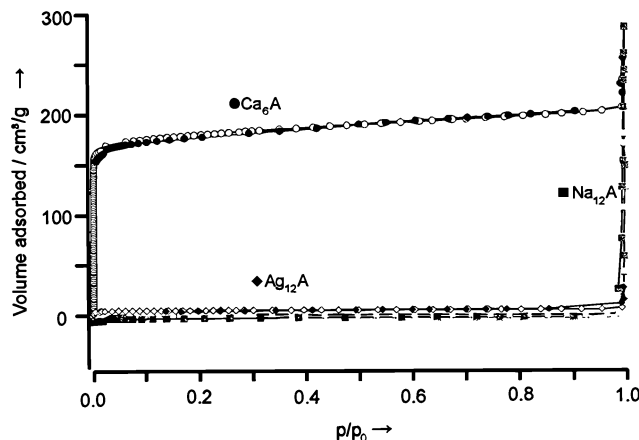


Fig. 3 Argon adsorption isotherm of Ca_6A -, Ag_{12}A -, and Na_{12}A -zeolites, measured at the boiling point of Ar (87 K). p/p_0 is the relative pressure; p_0 is the saturation pressure of Ar.

from concentrated solutions. In this way anions can be occluded which otherwise would not enter the negatively charged framework. Examples are the inclusion of Cl^- , Br^- , I^- ,²² and CN^- ²³ in zeolite A from boiling saturated aqueous solutions of *e.g.* the corresponding Na^+ or Ca^{2+} salts, followed by *e.g.* Ag^+ exchange. $\text{Ag}(\text{CN})_2^-$ and $\text{Au}(\text{CN})_2^-$ loaded zeolite A was prepared by reaction of potassium zeolite A for 12 h at room temperature with 4 M aqueous $\text{KAg}(\text{CN})_2$ and $\text{KAu}(\text{CN})_2$, respectively.¹¹ Another procedure is insertion of molecules from the gas phase into a dried zeolite, sometimes followed by a specific reaction. Examples are the insertion of AuCl_3 followed by reduction with CO ²⁴ and the insertion of CuCl .²⁵

Care has to be taken in any case, that the procedure does not damage the zeolite framework. X-Ray powder diffraction data and the morphology should be checked. A good example is the exchange of Cu^{2+} which works well for zeolites Y and X but which for zeolite A can only be used for low partial exchange, namely 1% to 2.5% by charge, depending on the co-cations. The zeolite framework breaks down at higher Cu^{2+} exchange as can first be seen by the corrugation of the crystal surface in SEM or TEM pictures, and at greater damage also in X-ray diffraction. It is reasonable to assume that strain is induced in the zeolite framework upon exchange of alkali cations by the Cu^{2+} . The cause of this strain is not the charge—materials which are fully exchanged with Ca^{2+} are perfectly stable—but the specific coordination geometry. This strain increases with increasing exchange level until the framework ruptures.²⁶ This problem can be avoided by ion exchange of Cu^+ in liquid ammonia²¹ or by insertion of CuCl from the gas phase.²⁷ In this case nearly a total exchange has been obtained without destroying the zeolite framework.

A general procedure for the preparation of quantum sized Ag_2S ,¹⁴ CdS ,²⁸ and ZnS ²⁸ clusters in zeolites involves first an ion exchange with Ag^+ , Cd^{2+} , or Zn^{2+} , then drying in vacuum and reacting with H_2S . For the preparation of ZnO and CdO nanoparticles, the ion exchange is followed by a calcination procedure.²⁹

Ag^+ loaded zeolites can be easily prepared by ion exchange usually from an aqueous silver nitrate or silver perchlorate solution. This ion exchange is fast and can be quantitatively controlled. The equilibrium and also the kinetics depend of course on the type of cations to be exchanged. It is especially easy to see this for sodium or potassium cations. The experimental procedure for ion exchange in the older literature often recommends exchange times of several days, which is not necessary and which can even damage the morphology of the crystals. We illustrate this by means of a simple kinetics experiment of an Ag^+ ion exchange at 20 °C in Fig. 4. It shows that ion exchange was complete in less than 20 min.

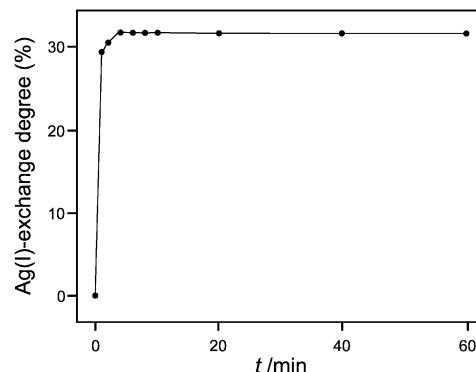


Fig. 4 Ag^+ -exchange degree of zeolite A as a function of the exchange time at 20 °C. A 9.1 mg sample of zeolite A was added to 200 mL of an aqueous 1 M NaNO_3 + 1 mM AgNO_3 solution. The Ag^+ content of the aqueous phase was measured by means of a Ag^+ ion selective electrode.³⁰

The ion exchange isotherm of the system $\text{Na}^+/\text{Ag}^+/\text{A}$ in Fig. 5 illustrates the strong preference for Ag^+ over Na^+ . We observe

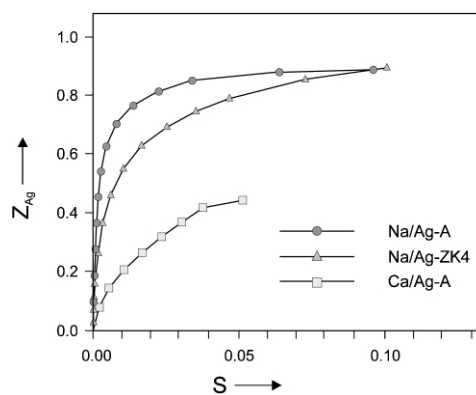


Fig. 5 Ion exchange isotherms for the systems $\text{Na}^+/\text{Ag}^+/\text{A}$, $\text{Na}^+/\text{Ag}^+/\text{ZK4}$ and $\text{Ca}^{2+}/\text{Ag}^+/\text{A}$ at room temperature, where Z_{Ag} is the exchange degree inside the zeolite and S is the equivalent fraction of Ag^+ in solution.¹³

a fast quantitative uptake up to an exchange degree of about 0.5 (that means for zeolite A 6 Ag^+ per α -cage). The system $\text{Ca}^{2+}/\text{Ag}^+/\text{A}$ shows quite a different isotherm. Because of the stronger bonding of the Ca^{2+} ions to the zeolite framework, the Ag^+ ions are less preferred than in the $\text{Na}^+/\text{Ag}^+/\text{A}$ system and also the exchange rate is slower. It takes about 1.5 h for the equilibrium to be reached. The exchange isotherm of the system $\text{Na}^+/\text{Ag}^+/\text{ZK4}$ rises less steeply than the isotherm of $\text{Na}^+/\text{Ag}^+/\text{A}$, but Ag^+ is still preferred over Na^+ .

3 The electronic structure of d^{10} - zeolites

The simplest approach, which has proven to be very successful, is to regard the electronic structure of a typical zeolite material to a first approximation as a superposition of the electronic structure of the framework, of the charge compensating cations, of the present solvent molecules (usually water), and of the guest species. We start with the band structure and density of states of the silicon dioxide analogue of zeolite A, which is especially simple because of the absence of co-cations. The result of a band structure calculation of a silicon dioxide analogue of zeolite A displayed in Fig. 6 shows that the bands in the HOMO region are flat which indicates the presence of nonbonding states. Some bands below -14 eV are significantly bent and contribute to the Si-O bonding. Further insight is gained from the density of states DOS(E), defined in such a way that $\text{DOS}(E)dE$ is the number of states in the interval E to $E + dE$. Since we are expressing the crystal orbitals as a linear combination of atomic orbitals (LCAO) we can highlight

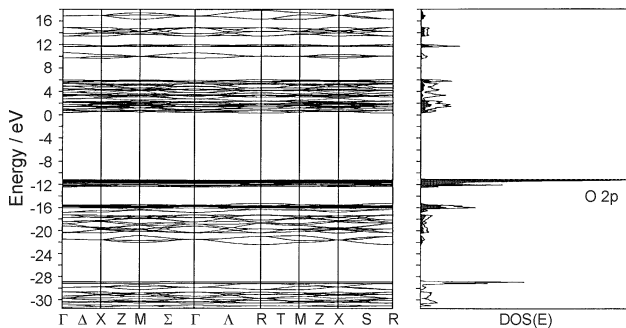


Fig. 6 Band structure (left) and density of states (right) of the silicon dioxide analogue of zeolite A. The oxygen 2p density is projected out (shaded regions).

specific atomic orbitals or linear combinations of them. This is done by shading the oxygen 2p contributions and leaving the 2s oxygen and the silicon contributions blank. This shows that the HOMO region consists of nearly pure oxygen 2p lone pairs which we denote as $|\text{O}\rangle$. The DOS of the LUMO region reflects the loose framework of the material.

The Al^{3+} centers in zeolites cause a negatively charged framework (AlO_2^-). This charge is compensated by exchangeable cations, which influence the band structure to some extent. Calculations show that the HOMO and the LUMO regions are in general similar to those illustrated in Fig. 6. Next we add a solvent. We only discuss water as a solvent molecule. The only level of importance is its lone pair at -12.6 eV, the energy of which is little affected *e.g.* by coordination to silver cations.³¹ We can therefore assume that the HOMO–LUMO region of zeolites containing Cu^+ , Ag^+ , and Au^+ can be drawn qualitatively as illustrated in Fig. 7. We show on the left a

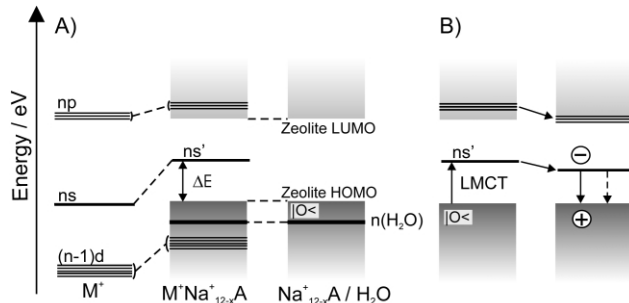


Fig. 7 A) Correlation diagram of $\text{M}^+(\text{d}^{10})$ loaded zeolite. On the left the energy levels of M^+ , on the right the energy levels of a zeolite are shown. The HOMO region of zeolites consists of many narrow lying localized states strongly concentrated on the oxygen atoms. We call this the lone pair region of the silicate and abbreviate it as $|\text{O}\rangle$. The water lone pairs are additionally marked. B) Energy diagram of metal cations in a zeolite framework. Some of the np' levels may reach into the LUMO region of the silicate. Three lines have been added to indicate this.

correlation diagram and on the right a simplified state diagram. It also holds for the alkali ions $\text{Li}^+, \dots, \text{Cs}^+$, but in this case the d-levels can be omitted. The levels consist of the oxygen lone pair region denoted as $|\text{O}\rangle$, the empty ns' level of the metal cations M^+ , and of the LUMO region of the zeolite which may be modified by np' contributions of M^+ . The ns' and np' and also the $(n-1)\text{d}$ levels are shifted to higher energy with respect to the ns and np levels of the free cations because of antibonding interactions with the oxygen lone pairs. This scheme strongly suggests the occurrence of ligand to metal charge transfer (LMCT) transitions of the $\text{ns}' \leftarrow |\text{O}\rangle$ type, where an oxygen lone pair electron is excited to the metal cation coordinated to the zeolite oxygen. The energy ΔE_{CT} needed for this transition is equal to the difference of the ionization potential $I_p|\text{O}\rangle$ of the oxygen lone pair and the first ionization potential $I_p\text{M}$ of the metal M, plus a correction Δ_M which stands for the antibonding interaction of the empty ns' level of the metal ion M^+ with the environment and maybe some shift of the potential energy

curve. The index M of the interaction term Δ_M indicates that the destabilization of the ns levels depend on the nature of the cation. We shall see in the next section, that it also depends on the coordination site, a fact we ignore at the moment.

$$\Delta E_{\text{CT}}(\text{ns}' \leftarrow |\text{O}\rangle) = I_p|\text{O}\rangle - I_p\text{M} + \Delta_M \quad (1)$$

This simple relation allows us to estimate the energy of the charge transfer band for different situations. $\text{ns}' \leftarrow |\text{O}\rangle$ charge transfer transitions have been observed in $\text{Cu}^+ \text{-- A}$, $\text{Cu}^+ \text{-- X}$ and $\text{Ag}^+ \text{-- A}$ zeolites in the region of $28\,000 \text{ cm}^{-1}$ (360 nm). This means that Δ_M is in the order of 4000 cm^{-1} for both metals, because the first ionization potentials of the Cu and Ag atoms are 7.68 and 7.54 eV, respectively. CT transitions for which Δ_M is larger will be discussed in the next section. Ag^+ in water absorbs light at about 225 nm ,³¹ in agreement with eqn. 1. Au has the largest $I_p\text{M}$, namely 9.18 eV, which gives Au its noble character. In general Au^+ is unstable but can be stabilized by specific ligands. This makes it difficult to incorporate Au^+ by conventional ion-exchange methods. However, some researchers have reported the synthesis of Au^+ loaded zeolites by means of ion exchange with gold complexes or by sublimation of $(\text{AuCl}_3)_2$.^{32,33} The electronic spectra of the resulting materials so far reported suggested that Au^+ was still coordinated to chloride and direct interaction to the zeolite could not be observed. Clear assignment of the observed electronic transitions have not been established. More work is needed to get electronic spectra of Au^+ coordinated to zeolite oxygen atoms.

Luminescence of the $\text{Cu}^+ \text{-- A}$, $\text{Cu}^+ \text{-- X}$, $\text{Cu}^+ \text{-- ZSM5}$, and $\text{Ag}^+ \text{-- A}$ zeolites after $\text{ns}' \leftarrow |\text{O}\rangle$ excitation occurs at $400\text{--}700 \text{ nm}$, depending on the samples and the conditions.^{25\text{--}27} This means that the Stokes shift is in the order of 8000 cm^{-1} . Excitation of an electron from the oxygen lone pair level $|\text{O}\rangle$ into the empty ns' orbital of the metal cation causes a formal reduction of M^+ to M^0 . The radius r of M^0 is significantly smaller than that of M^+ ($r(\text{Cu}^+) = 0.96 \text{ \AA}$, $r(\text{Cu}^0) = 1.35 \text{ \AA}$, $r(\text{Ag}^+) = 1.26 \text{ \AA}$, $r(\text{Ag}^0) = 1.6 \text{ \AA}$). This means that the LMCT transition expands the metal by $0.3\text{--}0.4 \text{ \AA}$ which causes a change in its position. As a consequence, the ns' level relaxes to a state of lower energy which we denote as the $(|\text{O}\rangle \oplus \text{ns}')^1$ state. The latter relaxes to the ground state either by emitting a photon with a large Stokes shift or by radiationless processes, as illustrated in Fig. 7 on the right hand side.

The luminescence intensity of Cu^+ and Ag^+ zeolites is remarkably influenced by the degree of hydration of the samples. Fig. 8 shows the luminescence spectra of

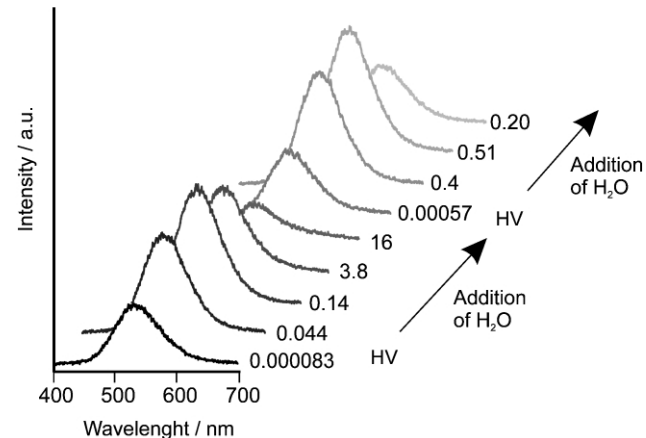


Fig. 8 Luminescence spectra of $\text{Cu}_{9.3}\text{H}_{9.3}\text{Na}_{67.4}\text{X}$ as a function of the degree of hydration measured at room temperature after excitation at 355 nm . Two evacuation–hydration cycles are shown. The numbers on the right side of the spectra are the pressure in the sample compartment in mbar. HV means 30 min of evacuation at room temperature.²⁶

$\text{Cu}_{9.3}\text{H}_{9.3}\text{Na}_{67.4}\text{X}$ as a function of the degree of hydration. This sample was prepared by a partial ion exchange of $\text{Na}^+ \text{-- X}$ with

an aqueous CuCl_2 solution at room temperature and selective reduction of Cu^{2+} to Cu^+ with CO .²⁶ At a water vapor pressure of 16 mbar the sample can be regarded as fully hydrated. Maximum emission is achieved at a well defined range of water content. While in the case of $\text{Cu}^+ \text{--} \text{A}$ and $\text{Cu}^+ \text{--} \text{X}$ the emission maximum only shifts 10 nm during rehydration, the luminescence of $\text{Ag}^+ \text{--} \text{A}$ shows more dynamics. In both cases, however, the optimum amount of water depends on the co-cations and on the type of zeolite. For example, while $\text{Ag}^+ \text{--} \text{Na}^+ \text{--} \text{A}$ doesn't show luminescence in its fully hydrated state, a weak luminescence can be observed in $\text{Ag}^+ \text{--} \text{Ca}^{2+} \text{--} \text{A}$ and $\text{Ag}^+ \text{--} \text{Mg}^{2+} \text{--} \text{ZK4}$. An explanation for this phenomenon can be attempted by considering the stronger electrostatic interaction between divalent cations and water than between monovalent cations and water. Thus in zeolites with divalent co-cations like Ca^{2+} and Mg^{2+} , less water is coordinated to Ag^+ than in *e.g.* $\text{Ag}^+ \text{--} \text{Na}^+ \text{--} \text{A}$. There are not many reports about luminescence of Au^+ -zeolites. However, luminescence of $\text{Au}(\text{CN})_2^-$ loaded zeolite A occurs at 390–445 nm and is assigned to trimers and tetramers of the complex. This also suggests that the Au^+ ions do not have strong interactions with the zeolite.

4 The electronic structure of Ag^+ zeolite A

Rálek *et al.* reported in 1962 that hydrated colorless zeolite $\text{Ag}^+_{x} \text{Na}^{+12-x} \text{A}$ turns yellow to brick-red on activation.¹⁹ No explanation of this phenomenon was given at that time. Later it was believed that the color change was due to formation of silver clusters (Ag_n^0) in the cavities of silver zeolite A. These neutral silver species were assumed to be formed at elevated temperatures *via* an auto-reduction process in which O_2 from the zeolite framework was released.³⁴ We studied the vibrational spectra of Ag^+ zeolite A materials in some detail,³⁵ and we showed that activation at room temperature under high vacuum is already sufficient to produce the yellow form of $\text{Ag}^+_{x} \text{Na}^{+12-x} \text{A}$. The fully reversible color change, which depends on the hydration state of the silver zeolite, was attributed to electronic charge transfer transitions from the oxygen lone pairs of the zeolite framework to the empty 5s orbital of the Ag^+ ions, denoted as $\text{Ag}^+(5s) \leftarrow \text{O}(n)$.¹² Silver-containing sodium zeolite A is colorless in its fully hydrated form. In activated silver zeolite A materials, the Ag^+ is forced to coordinate zeolite oxygen because an insufficient number of available water molecules are present. The question remained if specific coordination sites which act as yellow and/or red 'color centers' can be identified. We answered this question by studying the UV/vis spectra of $\text{Ag}^+_{x} \text{Na}^{+12-x} \text{A}$ and of $\text{Ag}^+_{x} \text{Ca}^{2+6-0.5x} \text{A}$ materials in their fully hydrated, in HV room temperature dehydrated, and in HV elevated temperature dehydrated states. The possible ion positions in $\text{Ag}^+_{x} \text{Ca}^{2+6-0.5x} \text{A}$, probed by gas adsorption experiments, offered the unique possibility of investigating different coordination sites of Ag^+ ions in zeolite A.¹³ Pure sodium and calcium zeolite A do not absorb light within the spectral range from 200–1000 nm we have investigated. This means that any absorption band or colors observed in silver zeolite A materials are due to the presence of silver ions. Fig. 9 shows the different colors of a $\text{Ag}^+_{6} \text{Na}^{+6} \text{A}$ zeolite in its fully hydrated state (white), after activation under high vacuum (2×10^{-7} mbar) at room temperature (deep yellow) and after heating in the vacuum to 200 °C (red). The spectrum of the yellow sample is fully reversible. After readsorption of water the yellow samples turn white again and the absorption spectrum is the same as before the activation.

We found that 6- and 8-ring coordinated Ag^+ give rise to electronic transitions in the near UV region. An absorption in the visible, namely at 450 nm, was only observed in materials where 4-ring coordinated Ag^+ was present and only they

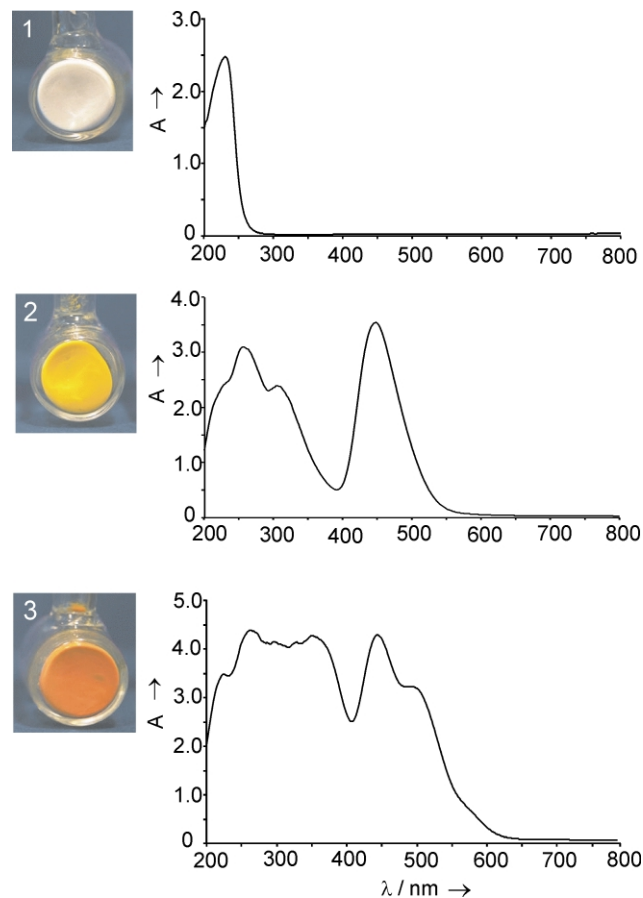


Fig. 9 Samples (left) and the corresponding UV/Vis spectra (right) of $\text{Ag}^+_{6} \text{Na}^{+6} \text{A}$ in its fully hydrated state (1), after activation in high vacuum at room temperature (2), and after activation in high vacuum at 200 °C (3).

showed the typical deep yellow color. We also observed that Ag^+ avoids the 4-ring sites as long as possible in $\text{Ag}^+_{x} \text{Ca}^{2+6-0.5x} \text{A}$, namely as long as x is smaller than 10. In the case of $\text{Ag}^+_{x} \text{Na}^{+12-x} \text{A}$ either a Na^+ or a Ag^+ is forced to coordinate a 4-ring site because all other places are occupied. The presence of the 450 nm absorption responsible for the yellow color, already at $x < 0.2$, proves that isolated Ag^+ ions are sufficient to cause it and that the 4-ring coordination of Ag^+ is significantly stronger than that of the Na^+ . The red color of elevated temperature activated samples is caused by a strong absorption band at 520 nm. We observed that samples which remained colorless after room temperature activation never turned red, that samples with lower silver content than one Ag^+ per α -cage never turned red, and that room temperature dehydration under our experimental conditions was not sufficient to produce red colored samples. These observations strongly indicate that only samples with 4-ring coordinated Ag^+ can give rise to the 520 nm band and this only occurs if a second Ag^+ is not too far away at a 6-ring site, so that they can interact to develop a corresponding low lying state.

Molecular orbital calculations carried out on a sufficiently large zeolite part consisting of 1296 atoms allowed us to address questions about the nature of the HOMO and of the LUMO region, about the contributions of the zeolite framework atoms to the electronic transitions, about the influence of the local symmetry of the Ag^+ at 4- and at 6-ring sites, and about the importance of $\text{Ag}^+ \text{--} \text{Ag}^+$ interactions. We found that 6-ring coordinated Ag^+ give rise to electronic transitions in the near UV and that the 4-ring coordinated Ag^+ is responsible for the deep yellow color of the room temperature activated material. This implies that similar $\text{Ag}^+(5s) \leftarrow \text{O}(n)$ LMCT transitions are to be expected in other Ag^+ exchanged zeolites. Ag^+ exchanged zeolite Y can be used as a test. We therefore measured UV/vis spectra of pure $\text{Na}^{+6} \text{Y}$, of room temperature HV dehydrated

Ag^{+69}Y , and rehydrated Ag^{+69}Y . The main result is that an intense band at about $34\,000\text{ cm}^{-1}$ appears upon dehydration which vanishes upon rehydration.¹³ Analogously one would expect a similar type of LMCT transitions in Cu^{+} zeolite materials. $\text{Cu}^{+}(4s)\leftarrow\text{O}(n)$ LMCT transition, reversible upon HV hydration/dehydration, have indeed been observed in Cu^{+} zeolite A and X.²⁶

Based on all the information which has been collected over the last few years, we can now draw the schematic state diagram in Fig. 10 for Ag^{+} containing zeolites.

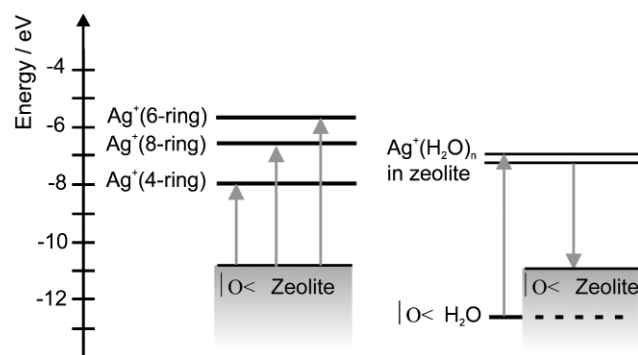


Fig. 10 State diagram of Ag^{+} -loaded zeolite A. On the left side we show the levels observed in room temperature activated zeolites in which all three sites are occupied by silver ions, while the scheme on the right side corresponds to situations typically observed in $\text{Ag}^{+}\text{xCa}^{2+}_{6-0.5x}\text{A}$ materials containing some water.

5 Quantum dots embedded in zeolites

The well defined cavities and channels of zeolites provide a convenient environment for preparing clusters and cluster arrays with a narrow size distribution as discussed in a review by G. Stucky and Mac Dougall.³⁶ d^{10} -Metal ion clusters of ZnO , ZnS , CdO , CdS ,^{28,29,37,38} of silver halides^{22,39} and more recently of silver sulfide¹⁴ were reported. The preparation of the latter was based on the observation discussed in Section 4 that Ag^{+} loaded zeolites can be reversibly activated at room temperature which allows the application of very mild synthesis conditions. This synthesis principle is expected to be applicable for the preparation of many other zeolite based host–guest materials. We therefore focus on the description of this most recent material.

$\alpha\text{-Ag}_2\text{S}$ is a semiconductor with a monoclinic structure and a band gap of approximately 1 eV at room temperature. Fig. 11 on

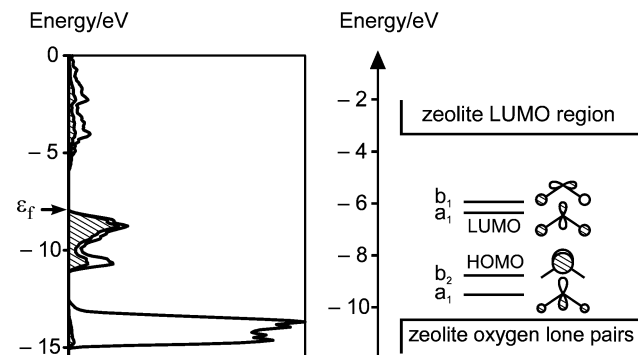


Fig. 11 *Left:* Density of states (DOS) plot of $\alpha\text{-Ag}_2\text{S}$. The hatched region indicates the contribution of sulfur 3p-states. The Fermi level ϵ_f is marked by an arrow. *Right:* HOMO–LUMO region of a Ag_2S molecule in comparison to the HOMO–LUMO region of zeolite A.

the left shows the calculated density of states (DOS) of bulk $\alpha\text{-Ag}_2\text{S}$. The electronic transition from valence band to conduc-

tion band is essentially a charge transfer from 3p(S) to 5s(Ag). This property is also observed in a Ag_2S molecule. The HOMO–LUMO region of such a molecule, and presumably also of larger silver sulfide clusters, fits well into the badgap of zeolite A (see right side of Fig. 11), therefore giving rise to a variety of electronic transitions.

Bulk silver sulfide has been considered for photoimaging and photodetection in the IR,⁴⁰ while small clusters are known to play an important role in photographic sensitivity.^{41,42} It has been reported that silver sulfide clusters with sizes ranging from 23 Å to 76 Å can be synthesized in reverse micelles. Another method utilizes the rapid expansion of a AgNO_3 solution in supercritical ammonia into an ethanol solution of Na_2S . An average diameter of 73 Å was found for particles formed in this way after stabilizing with a suitable polymer. Dosed addition of a AgNO_3 solution to a gelatin solution containing Na_2S was reported to yield silver sulfide clusters in the size regime between 30 Å and 100 Å. Other methods use thin nylon films, Nafion membranes or capping with cysteine/glutathione to stabilize the silver sulfide clusters.⁴³ Some of the results obtained by the above mentioned methods suggest the presence of a quantum size effect for silver sulfide clusters with a diameter between 20 Å and 100 Å while other reports clearly negate the presence of such an effect; for references see ref. 14.

Quantum sized silver sulfide clusters have been synthesized in the cavities of sodium and calcium zeolite A and ZK-4 microcrystals by exposing the room temperature activated Ag^{+} -loaded zeolite to H_2S . The formation of Ag_2S was also detected by X-ray photoemission spectroscopy (XPS) in zeolite Y after exposure of $\text{Ag}^{+}\text{-Y}$ to H_2S .⁸ The growth of the silver sulfide clusters in zeolite A and ZK-4 during rehydration of the samples was investigated by means of diffuse reflectance spectroscopy. Monomers of Ag_2S are formed at low Ag^{+} -loading through reaction of AgSH molecules, yielding colorless composites, which exhibit a characteristic blue-green photoluminescence. Yellow colored samples showing an orange-red luminescence with an average decay time of 81 μs ($-160\text{ }^{\circ}\text{C}$) are obtained at medium silver sulfide content. Further increasing the silver sulfide loading and therefore the cluster size causes a bathochromic shift of this emission which is accompanied by a shortening of the luminescence lifetime. The samples generally exhibit large Stokes shifts, which can be attributed to HOMO–LUMO transitions with small oscillator strengths. The Ag_2S –zeolite host–guest system constitutes a three-dimensional array of silver sulfide clusters. The experimental data and quantum chemical calculations show that the optical absorption and luminescence properties of this material are mainly due to the presence of isolated silver sulfide clusters inside the zeolite cavities. The characteristics of the composites are thereby to a certain extent influenced by the co-cations. Most remarkably, stronger luminescence, easily visible at room temperature, could be observed for silver sulfide clusters in calcium zeolite A as compared to clusters in sodium zeolite A. We show some recent observation on $\text{Ag}_2\text{S}\text{-CaA-}x$ materials to illustrate this: x denotes the average number of silver atoms per pseudo unit cell. Samples with low silver loading (e.g. $x = 0.01\text{ Ag}^{+}$) are white but show green luminescence. Fig. 12 displays as an example the spectra of $\text{Ag}_2\text{S}\text{-CaA-}0.01$.

Analysis of our data shows that these are the spectra of isolated Ag_2S molecules (quantum dots) in the three dimensional framework of zeolite A. This is the first time that the spectrum of an isolated Ag_2S molecule has been observed; see also Fig. 4 of ref. 14. A second luminescence band is observed when the silver loading is increased. Samples in the loading range $x > 0.1$ up to $x = 1$ show deepening of the yellow color. They exhibit both luminescence bands: the 490 nm and the new 620 nm emission; see Fig. 13. With increasing silver sulfide loading the 620 nm emission gets more intense and the 490 nm emission disappears. This means that the ratio of the amount of

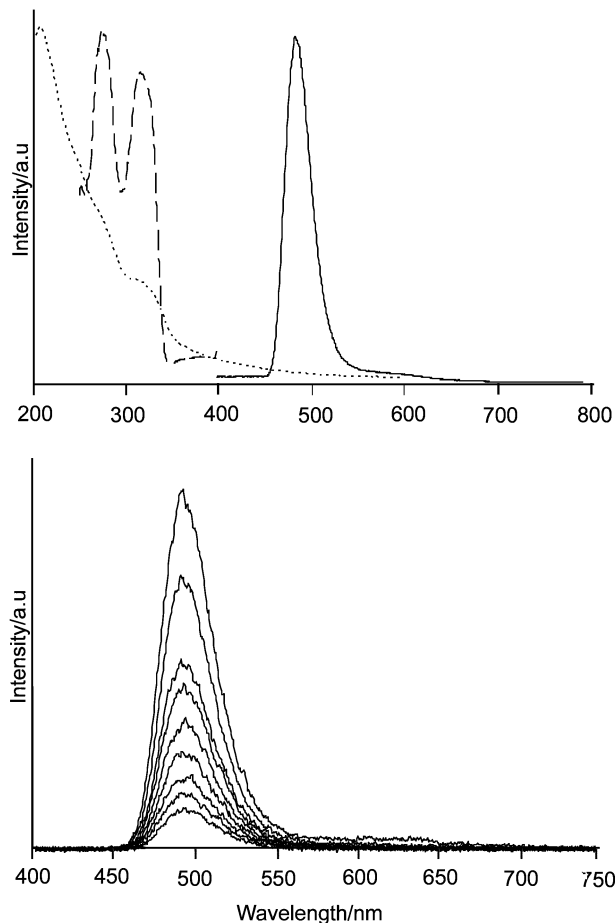


Fig. 12 *Top*: Emission (solid), excitation (dashed), and diffuse reflectance (points) spectra of $\text{Ag}_2\text{S}-\text{CaA}-0.01$. Luminescence was observed after 320 nm excitation while the excitation spectrum was measured at the maximum of the emission which is at 490 nm, both at $-190\text{ }^\circ\text{C}$. The diffuse reflectance spectrum (Kubelka–Munk) was measured at room temperature. *Bottom*: Luminescence spectrum of the same sample after excitation at 320 nm, observed at different times after excitation (time window 100 μs). Mono exponential decay was observed with a luminescence lifetime of 300 μs .

Ag_2S molecules and larger silver sulfide clusters in the zeolite can be tuned by choosing different initial amounts of Ag^+ . The two bands show a different intensity dependence on the temperature. This means that *e.g.* an $x = 1$ material could be used as a responsive micro temperature sensor, by comparing the intensity ratio of the two bands.

The luminescence spectrum in Fig. 13 can be broken down into two components. The 620 nm emission can be specifically excited at $\lambda > 400\text{ nm}$. It has a decay time of about 100 μs . This is in contrast to the 490 nm band which decays much slower. The 490 nm band corresponds in every respect to the luminescence shown in Fig. 12 which means that it stems from Ag_2S dots. We have good reasons to assign the 620 nm emission to $(\text{Ag}_2\text{S})_2$ clusters and we are now investigating if and to what extent radiationless excitation energy transfer from excited Ag_2S to $(\text{Ag}_2\text{S})_2$ cluster does occur. Similar results were observed by using zeolite ZK4 as a host, instead of zeolite A. Increasing loading shifts both the absorption and the emission bands to longer wavelengths, which is most probably due to the formation of larger clusters. The synthesis conditions and the material's properties are influenced by the co-cations (Li^+ , Na^+ , K^+ , Mg^{2+} , Ca^{2+} , Sr^{2+}), which is not astonishing in principle. However, the reasons for this are not well understood and more systematic work is needed. This is possible because the results are reproducible, which means that the system can be controlled.

We summarize that the reaction of activated Ag^+ -loaded zeolite A with H_2S leads to the formation of silver sulfide clusters in the zeolite cavities. The size of the clusters can be

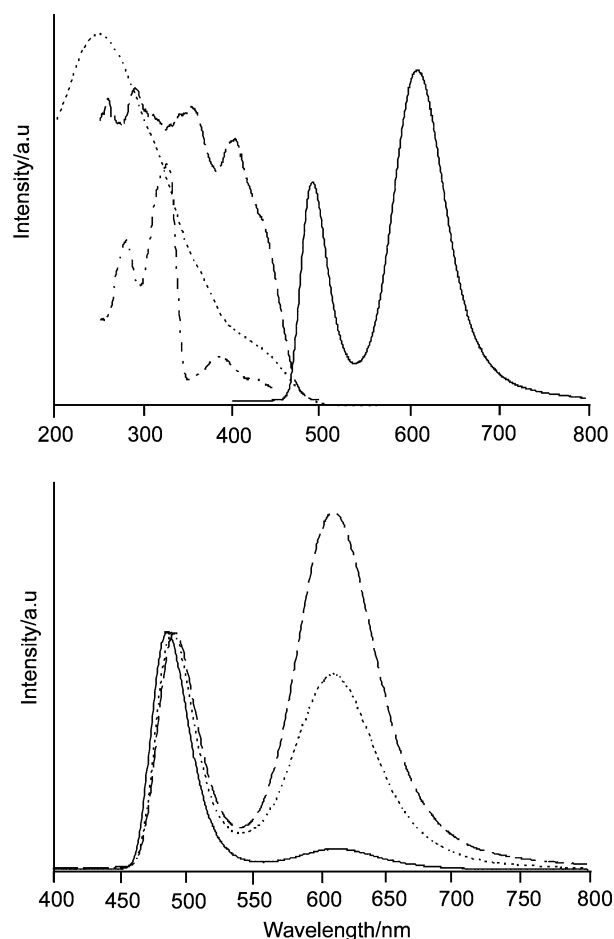


Fig. 13 *Top*: $\text{Ag}_2\text{S}-\text{CaA}-1$: luminescence (solid) and excitation spectra (dashed observed at 620 nm, dash dot observed at 490 nm) at $-190\text{ }^\circ\text{C}$, and diffuse reflectance spectrum (Kubelka–Munk) at room temperature. *Bottom*: Luminescence spectra of $\text{Ag}_2\text{S}-\text{CaA}-x$ with $x = 0.1$ (solid), $x = 0.75$ (dot), and $x = 1$ (dashed) after excitation at 320 nm. The spectra are scaled to the same height of the short wavelength band.

controlled by adjusting the initial Ag^+ -loading. Rehydration of the reaction products contributes to the cluster growth by enhancing the mobility of small silver sulfide and silver hydrosulfide species. The reaction between AgSH molecules, which are formed at low Ag^+ -loading levels, produces Ag_2S and H_2S . The formation of Ag_2S monomers and also of larger silver sulfide clusters is discernible by a band above 300 nm in the diffuse reflectance spectra of the composites. Considering the dimensions of an α -cage of zeolite A, the largest silver sulfide cluster that can be synthesized in such a cavity would have a stoichiometry of approximately Ag_8S_4 . Monomers of Ag_2S have to be considered as the smallest clusters. These species are responsible for the blue-green luminescence exhibited by samples of low silver sulfide content. At higher silver sulfide content a luminescence band around 620 nm is observed. The red-shift of this band with increasing silver sulfide loading can be attributed to a quantum size effect. The comparatively long average luminescence lifetimes observed for the orange-red emission in $\text{Ag}_2\text{S}-\text{CaA}$ (*e.g.* 81 μs for $\text{Ag}_2\text{S}-\text{CaA}-2$ at $-160\text{ }^\circ\text{C}$) and the large Stokes shifts are in agreement with the results of the MO calculations, which yield small oscillator strengths for the HOMO–LUMO transitions, but very large oscillator strengths for transitions of higher energy. According to these results, the qualitative state diagram shown in Fig. 14 can be constructed. Because of the inversely proportional relation between the oscillator strength of an electronic transition and the corresponding intrinsic emissive lifetime, the intense absorption transitions give rise to short-lived excited states. Non-radiative relaxation into the lowest excited state is followed by the observed long-lived luminescence. The $\text{Ag}_2\text{S}-$

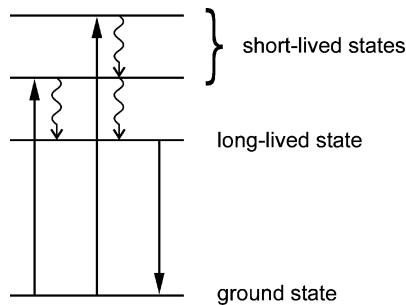


Fig. 14 State diagram for small luminescent silver sulfide clusters. This is valid for Ag_4S_2 and Ag_4S_2 .

zeolite host–guest systems constitute three-dimensional quantum dot arrays or quantum lattices. In the case of Ag_4S_2 clusters, coupling over the whole lattice is expected to occur at an average intercluster distance below 10 Å (center to center).¹⁴

6 Photocatalytic water oxidation

The splitting of water into hydrogen and oxygen driven by solar energy is considered as the ‘Holy Grail’ towards a clean and renewable energy source.⁴⁴ One of the key problems in photocatalytic water splitting has always been the oxidation of water to oxygen driven by visible light, mainly because of the difficulty in accumulating the four oxidative equivalents required. For several years it was known that aqueous dispersions of Ag^+ zeolites are able to produce oxygen under illumination, however, the conditions were not clearly defined. We observed that self-sensitization of photo-oxygen evolution occurs in this system. This means that dispersions which were initially insensitive to visible light become photoactive after they have been illuminated for a few minutes with near UV light.¹⁰ This system, however, was not catalytic and the role of the zeolite was not understood. We later found that aqueous dispersions of very pure Ag^+ exchanged zeolites were not photoactive, but that zeolites containing silver chloride were strongly photoactive provided that an excess of Ag^+ was present. Oxygen evolution was observed in a pH range of about 4 to 8. In fact the zeolite was found not to be necessary for the photochemical oxygen evolution from silver chloride. But the system was still not photocatalytic. Fortunately we found means to prepare silver chloride based electrodes in which the AgCl acts as a catalyst in the presence of a small excess of silver cations with a maximum oxygen evolution rate between pH 4 and 6. Its photoactivity extends from the UV to the visible region due to a process known as self-sensitization. The photocatalytic reaction consists of two main processes: water is oxidized to oxygen plus protons, and silver cations are reduced to silver upon irradiation. The reduced silver species are reoxidized which makes the system catalytic.^{45,46} The self-sensitization of the photochemical activity of AgCl is due to silver clusters adsorbed on its surface. A comparison of the energetic positions of the AgCl band gap⁴⁷ and the silver clusters shows that the silver clusters have empty energy levels below the conduction band of AgCl (see Fig. 15).⁴⁸

In the absence of silver clusters, AgCl does not absorb light below the indirect band gap transition, which is in the near UV at 3.3 eV (380 nm). Their presence enables a new electronic transition from the AgCl valence band to empty silver cluster energy levels. The energy for this transition is lower than the energy needed for an optical transition from the AgCl valence band to the AgCl conduction band. Nevertheless, this new optical transition in the visible spectral range still initiates the oxidation of water, because the conduction band is not directly involved in water oxidation.³¹ Thus, the photocatalytic oxidation of water on AgCl is extended from the near UV into the visible range of the spectrum. Self-sensitization as observed by

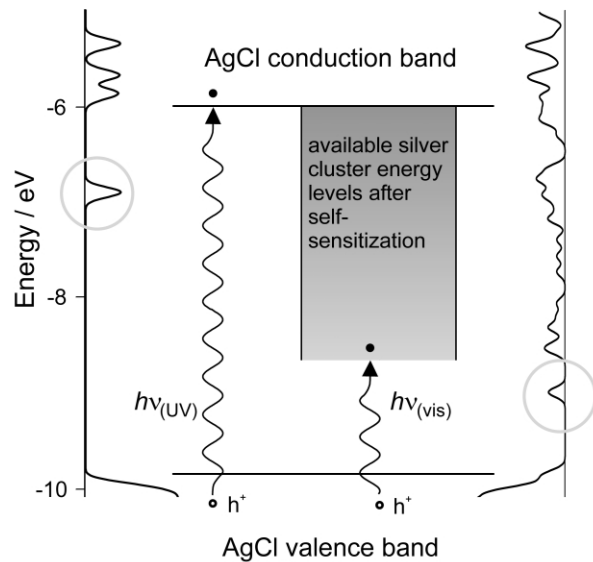


Fig. 15 Mechanism of self-sensitization. The density of states (DOS) of a silver chloride cluster is shown on the left and the *local DOS* of silver–silver chloride cluster composite is shown on the right. We observe new induced levels below the LUMO of the AgCl cluster. A schematic view is shown in the middle. The lowest lying levels into which electrons can be excited are marked with a circle (left: surface states, right: metal induced gap states). The position of the valence band is not markedly influenced by the presence of the silver cluster.⁴⁸

us should not be confused with the spectral sensitization effected by silver clusters, also termed photographic Becquerel effect. The latter observation has been attributed to an electron injection from electron-donating silver clusters into the AgCl conduction band. The first investigation of the photo-electrochemical properties of AgCl -coated electrodes was made by Becquerel in 1839,⁴⁹ a few years after Talbot had realized the first photographic pictures on paper based on AgCl .⁵⁰ The photo current observed by Becquerel was reductive.

Our experimental results show, that appropriately prepared silver chloride electrodes can fulfil this task and we therefore consider this material as a suitable choice in a photocatalytic water splitting device.^{31,45,46} However, the sensitivity of the silver chloride based photon anode is not satisfactory. We know that it can be improved, if the electrode is first covered with a monolayer of a Ag^+ exchanged zeolite A as described in ref. 30 and then modified with a thin layer of AgCl . An experiment which shows the photo response of such an electrode is shown in Fig. 16. The experiment was carried out under similar

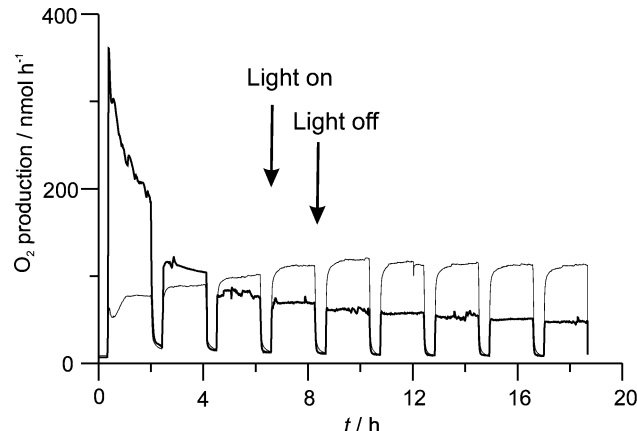


Fig. 16 Photo electrochemical experiment, illustrating the difference of the oxidation of water to O_2 for a AgCl surface modified zeolite-A electrode (bold), and for a AgCl layer on a gold modified conducting glass electrode. The corresponding current, about 8 μA during light period for the layer on gold, 0 for the zeolite layer, is not shown; see refs. 45 and 46.

conditions as reported in ref. 45. The AgCl modified zeolite layer has a maximum O_2 production rate that is almost 4 times

that of a AgCl layer on a gold modified conducting glass electrode which is likely due to its increased active surface area. However, the O₂ production rate is not sustained and falls to almost half its initial value by the end of the first cycle. One reason for the abrupt fall is the lack of electrical contact between the reduced silver on the surface of the zeolite (a product of the photo-reaction) and the conducting substrate. Without reoxidation of these reduced silver species the amount of photoactive AgCl on the surface cannot be maintained, hence the drop in O₂ production rate. That notwithstanding, the results show that there is much possibility for improvement by modifying the AgCl layer structure. Impressive progress on the preparation of zeolite monolayers has been made over the last few years, which should make it possible to combine the more reactive AgCl modified zeolite electrode with the stability and reversibility of a AgCl layer on a gold modified electrode.

7 Concluding remarks

A variety of procedures have been used to prepare d¹⁰-zeolite composites. The electronic structure of these materials can be regarded to a first approximation as a superposition of the framework, of the charge compensating ions, of solvent molecules and of guest species. Zeolite oxygen to d¹⁰-ion charge transfer transitions dominate the electronic spectra if the ions coordinate to the zeolite oxygens. Specific coordination sites can influence the energy and the intensity of these transitions remarkably. The reversible color change from colorless to deep yellow of Ag⁺-A and Ag⁺-ZK4 materials, which is very specific for Ag⁺ ions coordinated to the 4-ring, is a beautiful demonstration of this fact. Intra guest transitions dominate in quantum dot materials. We discuss this in detail for luminescent Ag₂S zeolite A. The possibility of preparing highly organized quantum dot materials opens a challenging world of possibilities in the assembly of highly organized macroscopic structures. We have discussed that the zeolite is not needed for the photocatalytic water oxidation on Ag⁺/AgCl photoanodes with visible light but that it can be used to increase the active surface area substantially. The challenge of developing reversible Ag⁺/AgCl photoanodes for water oxidation to di-oxygen based on this fact remains to be explored.

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9 References

- 1 D. W. Breck, *Zeolite Molecular Sieves*, John Wiley, New York, 1974.
- 2 D. L. Griscom, *J. Non-Cryst. Solids*, 1977, **24**, 155–234.
- 3 G. Calzaferri and R. Hoffmann, *J. Chem. Soc., Dalton Trans.*, 1991, 917–928.
- 4 T. Jüstel, D. U. Wiechert, C. Lau, D. Sendor and U. Kynast, *Adv. Funct. Mater.*, 2001, **11**, 105–110.
- 5 S. T. King, *J. Catal.*, 1996, **161**, 530–538.
- 6 M. Anpo, M. Matsuoka, K. Hanou, H. Mishima, H. Yamashita and H. H. Patterson, *Coord. Chem. Rev.*, 1998, **171**, 175–184.
- 7 A. Takahashi and R. T. Yang, *Langmuir*, 2001, **17**, 8405–8413.
- 8 A. Takahashi, R. T. Yang, C. L. Munson and D. Chinn, *Ind. Eng. Chem. Res.*, 2001, **40**, 3979–3988.
- 9 P. A. Jacobs, J. B. Utterhoeven and H. K. Beyer, *J. Chem. Soc., Chem. Commun.*, 1977, 128–129.
- 10 G. Calzaferri, S. Hug, T. Hugentobler and B. Sulzberger, *J. Photochem.*, 1984, **26**, 109–118.
- 11 S. M. Kanan, C. P. Tripp, R. N. Austin and H. H. Patterson, *J. Phys. Chem. B*, 2001, **105**, 9441–9448.
- 12 R. Seifert, A. Kunzmann and G. Calzaferri, *Angew. Chem.*, 1998, **110**, 1603 (*Angew. Chem., Int. Ed.*, 1998, **37**, 1521–1524).
- 13 R. Seifert, R. Rytz and G. Calzaferri, *J. Phys. Chem. A*, 2000, **104**, 7473–7483.
- 14 D. Brühwiler, C. Leiggner, S. Glaus and G. Calzaferri, *J. Phys. Chem. B*, 2002, **106**, 3770–3777.
- 15 S. B. Jang, U. S. Kim, Y. Kim and K. Seff, *J. Phys. Chem.*, 1994, **98**, 3796–3800.
- 16 D. W. Breck, W. G. Eversole, R. M. Milton, T. B. Reed and T. L. Thomas, *J. Am. Chem. Soc.*, 1956, **78**, 5963–5977.
- 17 K. H. Lim and C. P. Grey, *J. Am. Chem. Soc.*, 2000, **122**, 9768–9780.
- 18 F. M. Higgins, N. H. De Leeuw and Stephen C. Parker, *J. Mater. Chem.*, 2002, **12**, 124–131.
- 19 M. Rálek, P. Jíru, O. Grubner and H. Beyer, *Collect. Czech. Chem. Commun.*, 1962, **27**, 142–146.
- 20 P. E. Hathaway and M. E. Davis, *Catal. Lett.*, 1990, **5**, 333–348.
- 21 D. H. Strome and K. Klier, *J. Phys. Chem.*, 1980, **84**, 981–984.
- 22 R. M. Barrer and A. J. Walker, *Trans. Faraday Soc.*, 1964, **60**, 171–184.
- 23 R. J. Taylor, R. S. Drago and J. P. Hage, *Inorg. Chem.*, 1992, **31**, 253–258.
- 24 S. Qiu, R. Ohnishi and M. Ichikawa, *J. Chem. Soc., Chem. Commun.*, 1992, 1425–1427.
- 25 C. Lamberti, S. Bordiga, M. Salvalaggio, G. Spoto, A. Zecchina, F. Geobaldo, G. Vlaic and M. Bellatreccia, *J. Phys. Chem. B*, 1997, **101**, 344–360.
- 26 G. Calzaferri, R. Giovanoli, I. Kamber, V. Shklover and R. Nesper, *Res. Chem. Intermed.*, 1993, **19**, 31–57.
- 27 S. Bordiga, G. Turnes Palomino, D. Arduino, C. Lamberti, A. Zecchina and C. Otero Arean, *J. Mol. Catal. A: Chem.*, 1999, **146**, 97–106.
- 28 M. M. Garcia, H. Villavicencio, M. Hernandez-Velez, O. Sanchez and J. M. Martinez-Duart, *Mater. Sci. Eng., C*, 2001, **15**, 101–104.
- 29 M. Wark, G. Schulz-Ekloff and Nils I. Jaeger, *Bulg. Chem. Commun.*, 1998, **30**, 129–141.
- 30 J.-W. Li, K. Pfanner and G. Calzaferri, *J. Phys. Chem.*, 1995, **99**, 2119–2126.
- 31 S. Glaus and G. Calzaferri, *J. Phys. Chem. B*, 1999, **103**, 5622–5630.
- 32 Y.-M. Kang and B.-Z. Wan, *Catal. Today*, 1995, **26**, 59–69.
- 33 T. M. Salama, S. Takafumi, R. Ohnishi and M. Ichikawa, *J. Phys. Chem.*, 1996, **100**, 3688–3694.
- 34 T. Sun and K. Seff, *Chem. Rev.*, 1994, **94**, 857–870.
- 35 J. Baumann, R. Beer, G. Calzaferri and B. Waldeck, *J. Phys. Chem.*, 1989, **93**, 2292–2302.
- 36 G. D. Stucky and J. E. Mac Dougall, *Science*, 1990, **247**, 669–677.
- 37 Y. Wang and N. Herron, *J. Phys. Chem.*, 1987, **91**, 257–260.
- 38 J. E. Mac Dougall, H. Eckert, G. D. Stucky, N. Herron, Y. Wang, K. Moller, T. Bein and D. Cox, *J. Am. Chem. Soc.*, 1989, **111**, 8006–8007.
- 39 A. Stein, G. A. Ozin and G. D. Stucky, *J. Am. Chem. Soc.*, 1992, **114**, 8119–8129.
- 40 S. Kitova, J. Enevá, A. Panov and H. Haefke, *J. Imaging Sci. Technol.*, 1994, **38**, 484–488.
- 41 T. Tani, *J. Imaging Sci. Technol.*, 1995, **39**, 386–392; T. Tani, *J. Imaging Sci. Technol.*, 1998, **42**, 135–143.
- 42 R. C. Baetzold, *J. Imaging Sci. Technol.*, 1999, **43**, 375–381.
- 43 M. C. Brelle, J. Z. Zhang, L. Nguyen and R. K. Mehra, *J. Phys. Chem. A*, 1999, **103**, 10194–10201.
- 44 A. J. Bard and M. A. Fox, *Acc. Chem. Res.*, 1995, **28**, 141–145.
- 45 M. Lanz, D. Schürch and G. Calzaferri, *J. Photochem. Photobiol. A: Chem.*, 1999, **120**, 105–117.
- 46 G. Calzaferri, D. Brühwiler, S. Glaus, D. Schürch, A. Currao and C. Leiggner, *J. Imaging Sci. Technol.*, 2001, **45**, 331–339.
- 47 S. Sumi, T. Watanabe, A. Fujishima and K. Honda, *Bull. Chem. Soc. Jpn.*, 1980, **53**, 2742–2747.
- 48 S. Glaus, G. Calzaferri and R. Hoffmann, *Chem. Eur. J.*, 2002, **8**, 1786–1794.
- 49 E. Bequerel, *C.R. Acad. Sci.*, 1839, **9**, 561.
- 50 L. J. Schaaf, *Out of the Shadows: Herschel, Talbot, and the Invention of Photography*, Yale University Press, New Haven, CT, 1992.