MODELLING SPELEOGENESIS IN SOLUBLE ROCKS:
A CASE STUDY FROM THE PERMIAN ZECHSTEIN SEQUENCES
EXPOSED ALONG THE SOUTHERN HARZ MOUNTAINS
AND THE KYFFHÄUSER HILLS, GERMANY

MODELIRANJE SPELEOGENEZE V TOPNIH KAMNINAH:
PRIMER ZECHSTEINSKIH KAMNIN V JUŽNEM DELU GOROVJA
HARZ IN V HRIBOVJU KYFFHÄUSER

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Abstract

Georg Kaufmann & Douchko Romanov: Modelling speleogenesis in soluble rocks: A case study from the Permian Zechstein sequences exposed along the southern Harz Mountains and the Kyffhäuser Hills, Germany

Soluble rocks such as limestone, dolomite, gypsum, anhydrite, and salt can be dissolved by water flowing through voids in the rocks. The removal of the dissolved material from fissures and bedding partings by physical and/or chemical dissolution enlarges the permeability of the soluble rocks within geologically short periods of time, ranging from 100,000 years down to decades. This geologically short evolution time of voids in soluble rocks poses a substantial risk of mechanical instability of the enlarged voids, and possible surface deformation, when enlarged voids start to collapse.

We describe karst and cave features in the rock sequence exposed along the southern part of the Harz Mountains and the Kyffhäuser Hills in Germany, where limestone/dolomite and anhydrite/gypsum are exposed along a kilometer-wide strip following the foothills of the Harz Mountains. The rocks have been deposited during the Permian Zechstein period, buried, and exposed later through tectonic uplift. The exposed part of this soluble sequence is dominated by karst features. But there are also substantial cave voids deeper in the rock, with no obvious entrance to the surface, which have been discovered by chance through mining activities. Often, the sub-surface void evolution is closely linked to surface deformation, creating collapse sinkholes and subsidence. In the city of Bad Frankenhausen at the foothills of the Kyffhäuser Hills, the evolution of sub-

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Soluble rocks from the Permian Zechstein period are present underneath large parts of Northern Germany. These Permian Zechstein deposits comprise sequences of soluble rocks, with limestone along the base of a sequence, followed by anhydrite and salt, and often topped with insoluble marls and/or schists. While in most parts of Northern Germany these Permian Zechstein sequences can be found in several kilometer depth, in some parts the soluble rocks have been moved closer to the surface by salt diapirism and tectonic uplift.

Along the southern Harz Mountains in Germany, the Permian Zechstein sequences are exposed as a result of the tectonic uplift of the Harz along a narrow kilometer-wide strip, with widespread karstification within the outcropping soluble rocks. The uplifted soluble rocks dip gently towards the west / south-west, and karst features such as collapse sinkholes and resurgences are also marking the buried part of the dipping rocks. In the past, the base of the Zechstein sequence, a copper-shale ore (Kupferschiefer), has been mined extensively, with the discovery of numerous karst features such as special type of large mine caves (e.g., Brust & Nash 2013; De Waele et al. 2013), termed Schlotten in german, in the normally impermeable anhydrite layers.

Besides these often spectacular karst features, the dissolution of the soluble rocks (limestone/dolomite, anhydrite/gypsum, salt) is responsible for subsidence of the surface and resulting collapse sinkholes, once the overburden covering the enlarged voids evolved in the soluble rocks becomes too weak. A spectacular example of the ongoing dissolution in the Permian Zechstein sequence underneath the city of Bad Frankenhausen (Kyffhäuser Hills) is the inclined church tower of the Oberkirche, which has been build on a large fault with Permian Zechstein rocks in shallow depth. The titling of this church tower, with an inclination amounting to 4.9°, is caused by dissolution of the soluble rocks in the sub-surface, a still active process in the region.

We are interested in the solution process in the soluble rocks, especially the limestone and anhydrite layers in the shallow sub-surface of the Permian Zechstein rocks. As the timescales for dissolving limestone and anhydrite differ by more than one order of magnitude, interesting interactions between these types of soluble rock occur. We will study, by numerical means, a simplified setup of the inclined Permian Zechstein rocks, with dissolution in the limestone, dissolution in the anhydrite with associated precipitation of gypsum, and a regional water flow providing aggressive solution from below (hypogene settings).

We organize this paper as follows:

In the second section, we introduce the broader setup of the region under consideration, discuss the surface outcrop of the Permian Zechstein sequences with its soluble rock types carbonates, sulfates, and halites, and their continuation in the sub-surface further south. We will point out the hypogenic origin of many karst features in the region.

In the third section, we discuss three caves typical for the region, all developed in the anhydrite layers of the Permian Zechstein sequences, and their relation to surface features such as sinks, resurgences, and sinkholes, the latter related to significant surface deformation and thus a threat to infrastructure.

In the fourth section, we present the chemical reactions for the interaction of water and its dissolved agents with the soluble rock present in the working area. We discuss chemical equilibria, the way towards these chemical equilibria, and present time scales for dissolving rocks types such as the carbonates and the sulfates of the Permian Zechstein sequences. We finally discuss the combined dissolution of anhydrite and the precipitation of gypsum, usually responsible for the sealing of the anhydrite as aquitard.

In the fifth section, we use our chemical derivation and embed it into a three-dimensional flow, transport and reaction model, describing a simplified version of the Permian Zechstein rocks (only one sequence). We will see what is needed to generate Schlotten-type mine caves in the anhydrite rock.

Our key questions in this paper are:

1. Can we explain the evolution of the Schlotten-type...
mine caves with a large-scale karst evolution model, accounting for dissolution in a combined carbonate-sulfate setting?

2. Can we explain the surface deformation observed in Bad Frankenhausen with numerical models of large-scale dissolution in the underlying sulfate layers?

3. Can we estimate the role of aggressive mineral-rich brine water arriving from below through fault systems (hypogene setting)?

GEOLGY

In this section, we introduce the geological setting of the working area and discuss the lithology in the broader context of the soluble rocks exposed along the Southern Harz Mountains.

HARZ MOUNTAINS

The Harz Mountains in Northern Germany (Fig. 1) are mainly composed of old magmatic and metamorphic rocks (e.g., Hohl 1985; Schönenberg & Neugebauer 1987), and are surrounded by soluble rocks, mainly Cretaceous limestone in the North and Permian anhydrite and limestone/dolomite in the South. The mountain ridge, about 110 km long and 30–40 km wide, with its highest peak the Brocken (1141 m), has been uplifted during the Variscian mountain building period (700–500 Ma B.P.), and was almost denuded at the beginning of the Permian Period, when the Permian Zechstein ocean transgressed several times over this area (300–250 Ma B.P.), which was then located on the equator. The repeated deposition from the Zechstein ocean resulted in up to seven depositional cycles (z1/zw-Werra, z2/zs-Staßfurt, z3/zl-Leine, z4/zA-Aller, z5/zO-Ohre, z6/zf-Friedland, z7/zf-Fulda), all starting with carbonates (limestone/dolomite), sulfates (anhydrite, gypsum), halides (halite, sylvite), and finally shales. The bottom of the Permian Zechstein series is the Kupferschiefer, which has been mined in the past extensively.

After the Zechstein ocean finally disappeared, erosion deposited the late Permian and Triassic Buntsandstein rocks, topped by Muschelkalk. During the uplift of the European Alps (100 Ma B.P.), the Harz region got uplifted again, but erosion lowered the surface since then. A final uplift phase in the Tertiary caused an asymptotic uplift of the Harz Mountains, with the northeastern part forming a steep cliff and the southwestern part experiencing less uplift. In the wake of this tilting, the Permian Zechstein rocks, already buried at greater depth, were uplifted to the surface along a narrow stripe following the
Southern Harz. This area nowadays is intensively karstified.

KYFFHÄUSER HILLS

The tectonic evolution of the Kyffhäuser Hills (Fig. 2), a 6x13 km outcrop south of the Harz Mountains, is intimately linked to the Harz Mountains (e.g., Wunderlich 2014). The Kyffhäuser Hills, with its highest peak the Kulpenberg (473 m), experienced a similar asymmetric tilting during Tertiary times, creating a steep northern edge and exposing Permian Zechstein rocks along the southern rim. Additionally, two major faults bound the Kyffhäuser Hills, the northern (NKF) and the southern (SKF) Kyffhäuser Fault. Along the SKF, the Permian Zechstein rocks have been displaced vertically by several hundreds of meters.

GEOLOGICAL CROSS SECTIONS

Next, we discuss the lithological structure of the entire region. In Fig. 2, three profiles are marked with red lines. Geological cross sections of these three profiles are shown in Fig. 3.

The first section (A-A'-A'', Schriel & Bülow 1926b) traverses the southern Harz Mountains close to the city of Sangerhausen. Here, the Kupferschiefer has been mined extensively, and the lithologies are well known from mining. In Fig. 3 (top), the three letters mark mining shafts, with A the location of the Röhrigschacht, A' the location of the Thomas-Mün(t)zer-Schacht, and A'' the location of the Bernhard-Koenen-Schacht. Starting from the north, the Permian Zechstein sequences z1/zW (Werra) and z2/zS (Stassfurt) are exposed along the surface, caused by the tilting of the Harz. The soluble sequences then gently dip towards the south, and are covered by younger deposits, amongst other the Buntsandstein sequence. While the salts are absent in the sequences for the depth range from surface to about -200 m below sea level, they can be found further south (of Thomas-Münzer-Schacht), where the Permian Zechstein rocks are buried deeper and the salt layers have not been removed by circulating water from the surface, which is devoid of dissolved carbonates, sulfates, and salts. This cross section is representative for a location along the southern Harz Mountains, in which the Permian Zechstein layers are more or less undisturbed by further faulting.

In the second section in Fig. 3 (middle, B-B', Wunderlich 2014), the three first Permian Zechstein sequences are present (z1/zW-z3/zL). While in the western part close to the Kyffhäuser Hills the salts have been dissolved by circulating water, in the eastern part, beyond the Northern Kyffhäuser Fault (NKF), the salts are still present. The lithologies are confirmed by the deep boreholes Ud17/56 and Ud119/64.

The third section in Fig. 3 (bottom, C-C', Schriel & Bülow 1926a,b; Wadas et al. 2016) crosses the southern Kyffhäuser Hills towards the south. Here, north of the Southern Kyffhäuser Fault (SKF) the Permian Zechstein sequences z1/zW and z2/zS are exposed along the surface. South of the SKF, these sequence can be found around 300 m deeper, offset by the Southern Kyffhäuser fault, and here the salt in the z2/zS-Leine Series is still present. Thus, an interesting situation arises: the exposed soluble rocks north of the SKF are intensely karstified by meteoric waters, with numerous dolines, sinkholes, and caves, and along the SKF brackish water from deeper layers ascends and increases the calcium equilibrium concentration in the layers above due to the presence of dissolved salt ions.

FROM GEOLOGY TO KARST ...

We have seen that the entire region (Southern Harz Mountains, Kyffhäuser Hills) is characterized by soluble rocks of the Permian Zechstein Period, which outcrop along a narrow strip due to the tectonic uplift of the Southern Harz Mountains and the Kyffhäuser Hills, and then continue into the sub-surface with a shallow dip angle towards the south-west. Faulting has altered this simple sequence at some locations, with significant impact to the karstification, as we will see in the next section.

Fig. 2: Topographical map of the Kyffhäuser Mountains and its vicinity. The blue area is the Kelbra Reservoir, the black dashed lines the northern and southern Kyffhäuser fault (NKF, SKF) and the Kelbra Fault (KF), and the red lines with markers the profiles shown in Fig. 3. The three caves marked with bold letters and red polygons are: ES: Elisabethsächter Schlotte, BC: Barbarossa-Cave, NC: Numburg-Cave.
In this section, we present evidence for the widespread karstification of the region, with its special mine caves, the Schlotten, entrance-less voids in the anhydrite layers of the Permian Zechstein rocks, often exposed by mining operations. We then link the knowledge about sub-surface karst features to the situation in the city Bad Frankenhausen, where the tilted church tower of the Oberkirche is a spectacular sign of solution processes in the sub-surface.

CAVES

In the soluble Zechstein rocks, numerous caves can be found. A special type of caves are the Schlotten, mine caves in the anhydrite rock just above the limestone or dolomite layer. These Schlotten classify as hypogenic caves (e.g., Klimchouk 2011; Kempe 2008; Bauer & Elste 2016; Kempe et al. 2017), with water supply responsible for dissolution arriving through the underlying limestone/dolomite layers.

Fig. 3: Geological cross sections for the three profiles shown in Fig. 2. Faults are shown as black lines and marked (northern and southern Kyffhäuser fault, NKF, SKF), boreholes as white bars with black outlines. The letters indicate positions marked in Fig. 2.
These caves are often found by mining operations and are mainly located below the local base level, but accessible due to the pumping in the mine galleries. The German term *Schlotte* is a mining term describing natural cavities found during the mining operations, often in the anhydrite layers of the Permian Zechstein sequences. The *Schlotten* have often been used by the miners as a natural drainage route for mine water, and to store waste rock.

Next, we discuss three prominent caves of this type, which are characteristic for the karstification of the region.

**Elisabethschächter Schlotte**

The *Elisabethschächter Schlotte* is a large mine cave in the Werraanhydrit (z1/zW) (Fig. 4), found through mining operations during mining of the Kupferschiefer already in 1755, it has no natural entrance (e.g., Völker & Völker 1982; Brust 2005; Kupetz & Brust 2008; Kupetz 2008; Spilker 2016; Brust & Graf 2016). It is located just south of the exposed soluble rocks along the southern Harz rim (see marker in Fig. 2), in a classical undisturbed situation of the soluble Permian Zechstein rocks.

The cave is around 400 m long, with a succession of large rooms up to 14 m high. The cave follows the inclined bed of the Zechsteinkalk (z1/zW) layer, thus the rooms are dipping towards the south. Cave rooms, however, are developed in the Werraanhydrit (z1/zW). The larger cave rooms, often cupola-shaped, are connected by small crawlyways. Most of the floor of the cavity is a fine-grained sediment, made up by the insoluble parts of the anhydrite. In the northern part, the cave ends in a large breakdown room, the Chaosdom. Here, a connection to a sinkhole on the surface is evident, as numerous surface materials such as leaves, branches, and even artificial building material slowly migrate into the cave. The distance from the top parts of the Chaosdom to the surface, however, is still 44 m. The southern part of the cave is its lowest section and the last room Kuppeldom is completely filled with fine-grained sediments. While the cave extends vertically between 174-221 m N.N., the local base level Grenzbach is higher (266 m N.N.).

The cave interior is characterised by debris of gypsum material. This material is a result from the anhydrite-to-gypsum conversion in the cave rooms: The high humidity dissolves anhydrite from cave roof and walls, but gypsum is (almost) immediately precipitated. The increase in volume during the gypsum precipitation results in loose layers, which then easily detach under the gravitational acceleration imposed on them.

With the schematical cross section (Fig. 4), we describe the current hypothesis for the evolution of the cave: Aggressive water from the surface sinks along the permeable fractured limestone aquifer (in this case the Zechsteinkalk) and reaches calcium saturation with respect to the limestone. As the water table (before mining started) has been way above the cave location, the cavities evolved under deep phreatic conditions. Here, along the upper interface of the Zechsteinkalk, the water is in contact with Werraanhydrit, thus the water can still dissolve anhydrite along the interface. The widespread dissolution of the Werraanhydrit can be traced in the mine galleries along the limestone-anhydrite interface, as here the insoluble residuals from the Werraanhydrit accumulate as fine-grained sediments. However, the evolution of larger cupola-shaped rooms cannot be explained so easily, as they are not so widespread. Here, additional conditions need to be met, such as structural controls (e.g., faults) or preferential flow in the Zechstein layer causing an inhomogeneous distribution of water along the interface. Note that the cave follows the steep interface Zechsteinkalk-Werraanhydrit.

**Barbarossa-Cave**

The Barbarossahöhle is another mine cave of the typus Schlotte, also located in the Werraanhydrit (z1/zW) just above the Zechsteinkalk (z1/zW) layer (Fig. 4). The Barbarossa-Cave is located south of the Kyffhäuser Hills in the Permian Zechstein rocks, with the southern Kyffhäuser fault (SKF, for location see Fig. 2) nearby.

This large cave has been found in 1865 in search for new locations to mine the Kupferschiefer at the base of the Permian Zechstein rocks (e.g., Brust 2005; Kupetz 2005; Adler & Mertmann 2012; Kupetz 2014). Soon after the discovery, the mining operations were abandoned and the cave opened as a show cave.

The cave is around 800 m long, mainly horizontal and is characterised by a succession of larger rooms, often with wide cupola-shaped roofs. The groundwater table is exposed at various places, and in contact with the local base level. The spring Pfannspring about 700 m north of the cave provides surface water, which sinks after a short distance and reappears in the Barbarossa-Cave. Flow velocities, however, are very small (Adler 2013).

High humidity causes conversion of the anhydrite to gypsum along the roof areas of the chambers, the volume increase causes the layers of converted gypsum to shear off and fall down, once they become too heavy. This process is responsible for room enlargement. Parts of the large chambers are already in a state of breakdown, e.g., the Olymp, which reaches close to the surface, were the collapse sinkhole Teufelsgrube is located.

The Barbarossa-Cave does not follow the inclined bed of the Zechsteinkalk (as in the case of the Elisabethschächter Schlotte), instead it develops horizontally towards the local base level, the Wipper valley. The cave has formed under shallow phreatic conditions, but still its...
Fig. 4: Cave maps (left) and schematical cross sections (right) of three caves. The cave maps are shown in red, superimposed on the topography (color coded). For legend of geological cross sections see Fig. 3. Cave cross sections are shown in white, hypothetic water table as dashed black line, and faults as solid black lines. Top: Elisabethschächter Schloette, Middle: Barbarossa-Cave (dashed circle, location of collapse sinkhole), Bottom: Numburg-Cave (solid black outline-mapped parts, dotted black outline-hypothetical part). For legend see Fig. 3.
origin is hypogenic water from the Zechsteinkalk layer below the Werraanhydrit.

**Numburg-Cave**

The Numburghöhle is located east of the Kyffhäuser Hills along the southern shore of the Kelbra Reservoir (see Fig. 2 for location). The cave, which is developed in the Werraanhydrit (z1/zW), has been mapped to 1750 m length, and is mainly developed below the (current) water table (Völker and Völker 1991, 2014). The entrance of the cave has long been known as a cave spring, with brackish water emerging from a cave room, ending in a syphon. Starting in 1987, the extension of the Kupferschiefer-mining (Thomas-Münzer-Schacht, Bernhard-Koenen-Schacht) towards the west with its large-scale pumping was hampered by a substantial increase of leakage into the mining galleries. The groundwater withdrawal tapped the region around the Kelbra Reservoir, and the source of the Numburg-Cave became a sink, sucking around 10 m$^3$ of water per minute from the reservoir. The water level in the cave started to drop, and in August 1988 the phreatic parts of the cave became accessible, revealing a large cave (Fig. 4). As the inflow of water into the mining galleries become too large to be managed, the western part of the mine was sealed off by closing galleries with sealing dams and the water level in the Numburg-Cave and its surroundings started to rise again. In April 1989, the cave become flooded again up to its initial height.

The Numburg-Cave is a succession of large rooms, and it ends abruptly along the Kelbra fault. Here, in the last cave room, sandstone from the Carbon Period is exposed along a fault zone, and from this rock, water infiltrates the cave from several fissures in the sandstone. In the cave, brackish water emerges through two shafts, which have been explored to around 22 m depth. This brackish water ascends along the Kelbra fault zone from the Zechstein deposits in the north, which are located in greater depth, with the salt still present. Thus, the Numburg-Cave has at least two sources of water able to dissolve the Werraanhydrit, the aggressive water from the carboniferous sandstone, and the brackish water rising along the fault, the latter increasing the solubility of gypsum/anhydrite because of the presence of the dissolved salt.

**Bad Frankenhausen**

We now move on from the known caves and cavities in the soluble Permian rocks of the Southern Harz and Kyffhäuser region to the city of Bad Frankenhausen, which is located right on the Southern Kyffhäuser Fault (SKF). In Bad Frankenhausen, the church Oberkirche has been built in 1382, but around 1640 the tower started tilting towards the east. Nowadays, the leaning church tower has an inclination of 4.9° (Fig. 5). The reason for this inclination can be found in the sub-surface: As already mentioned, the Southern Kyffhäuser Fault runs just south of the church, and here the soluble Permian Zechstein rocks are offset vertically by about 200 m. Several sinkholes, clustered around the Southern Kyffhäuser Fault, can be found, with the largest ones the Quellgrund, an old sinkhole in Bad Frankenhausen known since medieval times (998, and currently the spa gardens), and the Abtissinnengrube, an elongated sinkhole 160 by 120 m in size, known since the 16th century. On the western site of the latter sinkhole, in 2009 a new sinkhole appeared, 20 m in diameter and 13 m deep, indicating the active dissolution processes (e.g., Brust 2010).

Since 1997/1999, the area around the church tower has been surveyed by precise levelling (Scheffler & Martienßen 2013). Two points from the 8 node levelling network show significant horizontal movements of around 74-90 mm and vertical subsidences of around 100-160 mm within the period 1997/1999-2012 (Fig. 6).
deformation is, however, not uniform in time. Since 2014, Kersten et al. (2017) conduct both geodetic (120 points) and gravimetric (13 points) measurements in Bad Frankenhausen. Annual local subsidence values measured during this campaign amount to 4-10 mm, however, is partly related to recent construction work, thus separation of natural and anthropogenic effects is still difficult.

In 2013 and 2014, several boreholes have been lowered in vicinity of the church tower by the TLUG (Thüringer Landesanstalt für Umwelt und Geologie) and in parts by the LIAG (Leibniz-Institute of Applied Geophysics) to investigate the sub-surface down to 450 m depth (for locations of the boreholes see Fig. 6). In Fig. 7, simplified borehole lithologies are shown: Four boreholes have been drilled close to the church. While the main borehole Ky1/2014, (TLUG. LIAG), with ca. 457 m depth, crossed the Southern Kyffhäuser Fault, the two supplementary boreholes Ky2/2013, (TLUG) and Ky3/2013, (TLUG) only reached around 100 m in depth. The fourth borehole Ky1/1993, (TLUG), also around 100 m in depth, was drilled earlier. All four lithologies reveal around 3 m anthropogenic infill, then around 3-7 m Quaternary deposits. Below, the Permian Zechstein sequences start, with the Stassfurt-Anhydrite (z2/zS) from 7-74 m, the Stassfurt-Carbonate (Stinkschiefer) between 74-80 m depth. The full Werra-Anhydrite (z1/zW) is only present in Ky1/2014, reaching from ca. 80-160 m, followed by the Werra-Carbonate (Zechsteinkalk) between ca. 160-163 m. The Permian Zechstein sequences are completed by the lowest marine deposit, the Kupferschiefer between ca. 163-165 m, followed by the final conglomerate layer. Below, between ca. 167-348 m, the Carboniferous Kyffhäuser rocks are present (mainly sandstones). In 348 m depth, the Southern Kyffhäuser Fault is traversed, dipping with 50-70°. Below, both Stassfurt-(z2/zS) and Werra-(z1/zW) deposits are found again, but this time with salt still present.

Two older boreholes in vicinity of the sinkhole Quellgrund have been lowered in 1857 and 1866 to provide brine water for medical purposes (e.g., Schriel & Bülow 1926a). The borehole Saline1 revealed a similar lithology as the main borehole Ky1/2014, traversing the Stassfurt and Werra rocks, then the Carboniferous Kyffhäuser rocks, crossing the Southern Kyffhäuser Fault, and finally the salt in the Stassfurt sequence below. The bore-

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**Fig. 6: Simplified street map and topography of Bad Frankenhausen. Leaning church tower (red square), boreholes (blue stars with name of borehole), and Southern Kyffhäuser Fault (SKF, black dashed line). The Quellgrund is an ancient collapse sinkhole. Measured surface deformation is shown as red contours (vertical component, in mm) and as black lines with yellow arrow heads (horizontal components, longest arrow 90~mm).**

**Fig. 7: Borehole lithologies, for legend see Fig. 3. For the more recent boreholes Ky*, cavities (white) and cavity infill (brown) from the cored samples is shown in the grey inlays in the right part of the lithologies.**
hole Saline2, south of the Southern Kyffhäuser Fault, reveals a completely different lithology: Thick Tertiary and Triassic sequences till 217 m depth, then a thick Permian sequence (mainly Staßfurt anhydrite) down to the final depth of 423 m.

In the recent boreholes, voids and cavities are also logged, which allows us to discuss the degree of karstification in Bad Frankenhausen (Fig. 7). Voids logged in the four boreholes are frequent, ranging from heights of 20-50 cm up to the meter scale, with the largest ones extending to more than 4 m vertically. Often, these voids are accompanied by collapse areas with infill from both brecciated remnants of gypsum and anhydrite, and residual marls from the layered anhydrite. Most of the voids mapped are located in the Staßfurt-Anhydrit (z2/zS), with one void (1.45 m height) identified in the underlying Stinkschiefer (Staßfurt-Carbonate). The size of some of the voids indicates larger cavities, which might have similar dimensions to the mapped Schlotten. A georadar cross-borehole survey between boreholes Ky1/2014 and Ky1/1993 confirms these large-scale voids (Miensopust et al. 2015; Wadas et al. 2017). Thus we speculate that below the northern part of Bad Frankenhausen, larger sub-surface voids can be present and their partial breakdown can be responsible for the observed surface deformations.

FROM KARST TO CHEMISTRY …

We have described the widespread karstification of the entire region, both along the outcropping Permian Zechstein sequences and in the sub-surface. A special type of cavities, the Schlotten, entrance-less voids often discovered by mining operations, characterise large-scale hypogenic karstification of the anhydrite layers, with aggressive water arriving from the carbonate layers below. In the next section, we discuss the chemical evolution of a solution on its way through the limestone into the anhydrite above, with the potential to enlarge fractures to the meter-size and thus create large sub-surface entrance-less voids.

CHEMISTRY

In this section, we discuss the chemical properties of the soluble rocks involved (limestone, anhydrite, gypsum, salt) and the time scales of dissolution for these rocks types. We also consider the conversion of anhydrite to gypsum.

REATIONS

The chemical reactions in the aqueous systems of gypsum (CaSO₄•2H₂O), anhydrite (CaSO₄), and limestone (CaCO₃) can be described as (e.g., Duan & Li 2008; Li & Duan 2011):

\[
\begin{align*}
H₂O & \leftrightarrow H⁺ + OH⁻ \\
H₂O + CO₂ & \leftrightarrow H₂CO₃ \\
H₂CO₃ & \leftrightarrow H⁺ + HCO₃⁻ \\
HCO₃⁻ & \leftrightarrow H⁺ + CO₃²⁻ \\
CaSO₄•2H₂O & \leftrightarrow Ca^{²⁺} + SO₄²⁻ + 2H₂O \\
CaSO₄ + 2H₂O & \leftrightarrow Ca^{²⁺} + SO₄²⁻ + 2H₂O \\
CaCO₃ + 2H₂O & \leftrightarrow Ca^{²⁺} + CO₃²⁻ + H₂O \\
CaCO₃ + H⁺ & \leftrightarrow Ca^{²⁺} + HCO₃⁻ \\
CaCO₃ + H₂CO₃ & \leftrightarrow Ca^{²⁺} + 2HCO₃⁻ \\
\end{align*}
\]

The first reaction describes the dissociation of water, the second the solution of carbon dioxide (CO₂) in water, the next two ones the dissociation of carbonic acid, and the remaining equations the solution of gypsum, anhydrite and limestone, respectively.

EQUILIBRIA

A solution being in contact with a surface of soluble rock will dissolve the rock, until a chemical equilibrium between the dissolved species is attained. For gypsum, anhydrite, and limestone, we can characterise this equilibrium by tracking the calcium concentration, \(c_{eq}\) [mol/m³] and the respective calcium equilibrium concentration \(c_{eq}\) [mol/m³].

For gypsum and anhydrite, we can easily derive the calcium concentration from mass-balance considerations. For limestone, the derivation of an analytical relation for the calcium equilibrium concentration is a bit more tedious and results in an approximate solution (e.g., Dreybrodt 1988), which, however, compares well and to excellent accuracy with numerical solutions derived for the full electro-neutrality condition, which we verified by comparing our \(c_{eq}\) for limestone to results obtained from PHREEQC (Parkhurst & Appelo 2013).

The above mentioned chemical equations then result in calcium equilibrium concentrations \(c_{eq}\) for the different soluble rock types:
In (2), the mass-balance coefficients are:

\[ K_A(T,z) \text{ (Blount & Dickson 1973) the equilibrium constant for the dissolution of anhydrite, } K_G(T,z) \text{ (Blount \\
& Dickson 1973) the equilibrium constant for the dissolution of gypsum, } K_H(T,z) \text{ (Weiss 1974) the equilibrium constant for the dissolution of atmospheric carbon dioxide into water (Henry's law constant), } K_0(T,z) \text{ (Wissbrun et al. 1954) the equilibrium constant for the reaction of water and carbon dioxide to carbonic acid, } K_1(T,z) \text{ and } K_2(T,z) \text{ (Millero 1979; Mehrbach et al. 1973) the equilibrium constants for the dissociation of carbonic acid into bicarbonate, carbonate, and hydrogen, } K_C(T,z) \text{ (Mucci 1983) the equilibrium constant for dissolved calcite, } \gamma_{Ca^{2+}}, \gamma_{SO_4^{2-}} \text{ and } \gamma_{HCO_3^-} \text{ the activity coefficients for calcium, sulfate and bicarbonate, } p_{CO_2}^{\text{atm}} \text{ the carbon-dioxide partial pressure, } T [\text{oC}] \text{ the temperature of the solution, and } z [\text{m}] \text{ is depth, accounting for hydraulic pressure increase with depth.} \]

For limestone, (2) is valid for the open system, in which the solution is in contact with the atmosphere, and carbon dioxide will be replenished by further solution of CO\textsubscript{2} from the atmosphere. However, most fracture enlargement takes place under closed-system conditions, and here the CO\textsubscript{2} is consumed and thus decreases with dissolution. In this case, the carbon-dioxide partial pressure is (Dreybrodt 1988):

\[ p_{CO_2} = p_{CO_2}^{\text{atm}} - \frac{c_{eq}}{K_H \left(1 + \frac{1}{K_0}\right)} \]

with \( p_{CO_2}^{\text{atm}} \) the initial carbon-dioxide partial pressure obtained in the atmosphere and the soil.

In Fig. 8, the calcium equilibrium concentrations are shown for limestone, anhydrite, and gypsum, and for comparison also the sodium equilibrium concentration for the dissolution of rock salt (NaCl) is shown as a function of temperature and for different depth (representing hydraulic pressure) and in the case of limestone also for different carbon-dioxide pressure values.

While the equilibrium concentrations for limestone are between 0.1-3 mol/m\textsuperscript{3}, values for gypsum and anhydrite are about one order of magnitude higher, with values of around 15 mol/m\textsuperscript{3} for gypsum and 40 mol/m\textsuperscript{3} for anhydrite for temperatures below 40\textdegree C. For halite, the sodium concentration is even two orders of magnitude higher. Note that for temperatures below 50\textdegree C, the solubility of gypsum is lower than the one for anhydrite, an important observation, which we will discuss in more detail below. Also note that for limestone and anhydrite, the equilibrium concentration is a retro-grade function

![Fig. 8: Calcium equilibrium concentration as a function of temperature for limestone, anhydrite, and gypsum. Additionally, the sodium equilibrium concentration for halite is shown. Curves are shown for three different hydrostatic pressures (with hydraulic head as variable), and in the case of limestone for three different carbon-dioxide pressure representations.](image-url)
of the temperature, while for gypsum we observe a pro-
grade dependence on temperature for temperatures be-
low 30°C.

The dependence on hydraulic pressure is always
below one order of magnitude, and higher pressures in-
crease the equilibrium concentrations.

For limestone, the dependence on carbon-dioxide
pressure is non-linear and very pronounced, and respon-
sible for the well-known and investigated dissolution
processes in that rock.

**FLUX RATES**

With the calcium equilibrium concentration discussed,
we now know, how much of the soluble rock surface can
be dissolved. If we, however, also want to know, how fast
the dissolution process occurs, we need to introduce the
flux rate $F$ [mol/m$^2$/s].

Calcium flux rates are known both from labora-
tory experiments and numerical modelling. For lime-
stone, flux rates have been measured experimentally
(e.g., Plummer *et al.* 1978; Svensson & Dreybrodt 1992;
Eisenlohr *et al.* 1999), as well as determined by numerical
experiments (e.g., Buhmann & Dreybrodt 1985a, 1985b;
Dreybrodt & Kaufmann 2007; Kaufmann *et al.* 2010).
For gypsum and anhydrite, flux rates have also been mea-
sured in the laboratory (e.g., James & Lupton 1978; Go-
bran & Miyamoto 1985; Lebedev & Lekhov 1990; Jeschke

All of the above mentioned flux rate experiments
can be described by a flux rates $F$ [mol/m$^2$/s] with a
piece-wise function of the calcium concentration $c$ with
respect to the calcium equilibrium concentration $c_{eq}$:

$$F = k_i \left( \frac{m_i - c}{c_{eq}} \right)^{n_i}, \quad i = 1, 3$$

with $k_i$ [mol/m$^2$/s] a rate coefficient, $m_i$ [-] a flux rate
regime coefficient, $c$ [mol/m$^3$] the actual calcium
concentration, $c_{eq}$ [mol/m$^3$] the calcium equilibrium
concentration, and $n_i$ [-] a power-law exponent (see Kaufmann *et al.* (2010) for more details, and Tab. 1 for parameter values). The counter $i$ marks different re-
gimes of the flux rate function. The rate coefficient de-

deps on both surface-controlled rates and diffusion-
controlled rates:

$$k_i = \frac{k_i^S k_i^D}{k_i^S + k_i^D}$$

with $k_i^S$ [mol/m$^2$/s] the surface-controlled rate coefficient,
and with $k_i^D = 2Dc_{eq}/d$ [mol/m$^2$/s] the diffusion-controlled
rate coefficient, $D$ [m$^2$/s] the diffusion coefficient, and $d$
[m] the film thickness. The film thickness describes the
thickness of the solution layer on the fracture wall, e.g.,
for a circular conduit it can be defined by the ratio of flu-
id volume in the conduit $V$ [m$^3$] to reactive surface area
of the conduit $A$ (m$^2$), $d=V/A$.

**Tab. 1: Parameter values for soluble rock chemistry.**

<table>
<thead>
<tr>
<th>Rock</th>
<th>atomic mass $m_{rock}$ [kg/mol]</th>
<th>density $\rho_{rock}$ [kg/m$^3$]</th>
<th>rate constant$^1$ $k_i$ [mol/m$^2$/s]</th>
<th>rate exponent$^1$ $n_i$ [-]</th>
<th>Switch$^1$ $c_{eq}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Limestone</td>
<td>0.100</td>
<td>2600</td>
<td>4x10$^7$</td>
<td>1.0</td>
<td>0.90</td>
</tr>
<tr>
<td>Anhydrite</td>
<td>0.136</td>
<td>2900</td>
<td>4x10$^5$</td>
<td>5.4</td>
<td></td>
</tr>
<tr>
<td>Gypsum</td>
<td>0.172</td>
<td>2200</td>
<td>1.1x10$^4$</td>
<td>4.5</td>
<td>0.95</td>
</tr>
</tbody>
</table>

$^1$ Buhmann *et al.* (1985a,b)

$^2$ Jeschke (2002)

$^3$ Jeschke *et al.* (2001)
In Fig. 9, flux rates following (4) are shown for anhydrite, gypsum, and limestone. For both limestone and gypsum, the flux rate function is linear for strong undersaturation with respect to calcium, but becomes non-linear close to the calcium equilibrium concentration. For anhydrite, the entire flux rate below saturation is non-linear. The non-linearity, which results in much lower flux rates close to saturation, is often a result of impurities in the rock (e.g., shales, marls), which accumulate on the rock surface, while the soluble parts of the rock are dissolved and transported away in solution (Svensson & Dreybrodt 1992; Eisenlohr et al. 1999; Buhmann & Dreybrodt 1985a,b).

Beyond the saturation point, precipitation of the mineral will start. Here, experimental data are rare, and only precipitation with calcite and limestone is based on experimental data (Buhmann & Dreybrodt 1985b; Dreybrodt & Buhmann 1991). For precipitation rates of anhydrite and gypsum, we simply assume a linear relation (see Kaufmann et al. 2017 for more details). This assumption rests on the fact, that during dissolution impurities accumulate on the interface solution-rock, and are responsible for the non-linear rates close to equilibrium, while during precipitation the pure mineral species will deposit, which we assume can be described by a linear rate law.

**TIME SCALES**

We now know what the amount of dissolved material is \(c_{eq}\) and how fast is will be dissolved \(F\). We combine these two informations to describe the temporal evolution of the calcium concentration \(c\) in the water film with a differential equation:

\[
\frac{\partial c}{\partial t} = \frac{A_s}{V} F(c, d),
\]

with \(A_s [\text{m}^2]\) the reactive surface area, and \(V [\text{m}^3]\) the fluid volume.

The ratio \(A_s/V\) is known as specific surface, and we can express this in terms of film thickness \(d\) by defining a circular conduit with surface area \(A_s = 2\pi r l\) and volume \(V = \pi r^2 l\), with \(r [\text{m}]\) the conduit radius \((d=2r)\) and \(l [\text{m}]\) the conduit length. We thus find \(A_s/V \sim 1/r\). Similar relations hold for rectangular conduits. Note that (6) describes a stagnant film, transport and advection will be considered in the next section.

With both \(c_{eq}\) and \(F\) known for anhydrite, gypsum, and limestone, the temporal evolution of the modelled calcium concentration \(c\) is shown in Fig. 10. All solutions start with a calcium concentration of zero, \(c_{in} = 0\). For the low film thickness value of \(d=0.001 \text{ m}\) (solid lines), a water film on limestone reaches equilibrium after about 10 h, while the same solution on a gypsum surface would be saturated after only 1 h. For anhydrite, the non-linear flux rate is responsible for reaching the saturation point even later, around 50 h in this case. Note the curve for halite, which we plotted for comparison. The sodium equilibrium concentration in this case is reached within minutes!

If we increase the film thickness by an order of magnitude \((d=0.010 \text{ m},\) dotted lines), the times needed to reach the equilibrium concentrations for the soluble
rock types shown increase by about one order of magnitude. The reason for the increase in time with increasing fracture width is the boundary layer, which for this larger fracture is more pronounced and the diffusion of dissolved species out of this boundary layer takes longer (e.g., Buhmann & Dreybrodt 1985a, 1985b). Thus, increasing the film thickness $d$ changes the rate coefficient (5).

**ANHYDRITE-GYPSUM CONVERSION**

We now finally discuss a water film in a fracture in anhydrite, with solution undersaturated with respect to calcium passing through it. As discussed above, the calcium concentration $c$ increases in the fracture, but now we allow gypsum to precipitate, once the calcium equilibrium concentration for gypsum, $c_{eq}^{\text{gypsum}} \approx 15 \text{ mol/m}^3$, is reached (Fig. 11). Note that the water film is still stagnant.

![Graph 1](image1.png)

**Fig. 10:** Calcium concentration as a function of time. Shown are concentrations for anhydrite, gypsum, and limestone approaching their respective calcium equilibrium concentrations (dashed lines), for a temperature of $T=10^\circ$C, a hydraulic head of $z=0$ m, a film thickness of $d=0.001$ m (solid lines) and a film thickness of $d=0.01$ m (dotted lines), and a carbon-dioxide pressure of $P_{\text{CO}_2}=0.05$ atm for calcite. Additionally, for halite the sodium concentration and the respective sodium equilibrium concentration is also shown (note that these quantities are scaled by a factor $1/200$).

![Graph 2](image2.png)

**Fig. 11:** Calcium concentration as a function of time in the case of anhydrite-gypsum conversion. Once the calcium concentration of the solution passes the calcium equilibrium concentration of gypsum, gypsum starts to precipitate. Also shown is the reduction in fracture width $d$ as a result of the gypsum precipitation for two initial fracture width (see legend).
We follow a strategy first described by Kontrec et al. (2002) and later extended by Serafeimidis and Anagnostou (2012). We solve the coupled reaction equations for anhydrite and gypsum concentration and fracture width:

\[
\frac{\partial c}{\partial t} = \frac{A_s}{V} F_{\text{anh}}(c,d) + \frac{A_s}{V} F_{\text{gyp}}(c,d), \\
\frac{\partial r}{\partial t} = \frac{m_{\text{anh}}}{\rho_{\text{anh}}} F_{\text{anh}}(c,d) + \frac{m_{\text{gyp}}}{\rho_{\text{gyp}}} F_{\text{gyp}}(c,d).
\]

In (7), \( r \) [m] is the fracture half-width, and \( m_{\text{anh, gyp}} \) [kg/mol] the atomic mass and \( \rho_{\text{anh, gyp}} \) [kg/m^3] the density of anhydrite or gypsum, with anhydrite the dissolving and gypsum precipitating mineral. Note that the terms for the gypsum flux rate, \( F_{\text{gyp}} \), are only active, when the calcium concentration \( c \) in the solution exceeds the calcium equilibrium concentration for gypsum, \( c_{\text{eq, gyp}} \), because at the start of the experiment no gypsum is present.

The concentration curve for anhydrite dissolution starts as before (green line), increasing from zero and heading towards the calcium equilibrium concentration of anhydrite, \( c_{\text{anh}} = 45\text{ mol/m}^3 \). However, soon after passing \( c_{\text{eq, anh}} = 15\text{ mol/m}^3 \) (blue dashed line), gypsum starts to precipitate and the concentration curve approaches a plateau at \( c_{\text{eq, gyp}} = 17-18\text{ mol/m}^3 \). With time, the concentration slowly drops again. The reason for this drop is that gypsum precipitates faster than anhydrite dissolves, and the fracture width therefore slowly reduces. This can be seen for the two black lines with different initial fracture widths \( d=0.001\text{ m} \), solid black line, \( d=0.0005\text{ m} \), dotted black line): Both fractures initially grow a bit in size (hardly visible in the plot), but once the calcium concentration exceeds the calcium equilibrium concentration for gypsum \( (c > c_{\text{eq, gyp}}) \), the fracture width decreases. This closure of the fracture is forced by the larger volume the gypsum occupies, when compared to anhydrite.

After about 100 h (roughly four days) for the fracture with \( d=0.0005\text{ m} \) initial width and about 1000 h (roughly a month) for the fracture with \( d=0.001\text{ m} \) initial width, the clogging accelerates and after around 1 year the fracture is clogged. Only fractures with larger initial width \( d>0.01\text{ m} \) remain open, as from this size on the amount of gypsum precipitating is no longer enough to fill the fracture completely.

FROM CHEMISTRY TO EVOLUTION ...

We have described the dissolution (and precipitation) of soluble rocks for the three important rock types limestone, gypsum, and anhydrite present in the Permian Zechstein sequences in the Southern Harz and the Kyffhäuser Hills. For our geological setup, it is important that meteoric water is transported deeper into the sub-surface through the well karstified carbonates, often saturated with respect to this carbonates, but still able to dissolve anhydrite, which has a much higher calcium equilibrium concentration. As we are most likely still in a flow regime with temperatures below 60°C, during the dissolution of anhydrite the calcium equilibrium concentration for gypsum is reached, and gypsum starts to precipitate. Depending on the initial fracture size distribution in the anhydrite, smaller fractures are quickly clogged and then seal off the anhydrite (as aquitard). Only a thin layer of anhydrite along the carbonate-anhydrite boundary is dissolved, producing the Schlottenlehmen, a residual clay material often found along this interface.

However, in areas characterised by larger initial fractures, e.g., faults, the precipitated gypsum cannot clog the entire fracture, and the anhydrite dissolution proceeds deeper into the anhydrite rocks, creating the initial voids needed for the evolution of the Schlotten. We speculate that the observed surface deformation with its visible landmark the tilted church tower of the Oberkirche in Bad Frankenhausen is also a result of solution processes within the Permian Zechstein rocks beneath the city.

With this knowledge of dissolution and precipitation in the soluble Permian Zechstein rocks, we proceed to the next section, discussing a three-dimensional model for the evolution of flow and transport in the system. We aim to describe a simplified system with hypogene water supply through limestone into anhydrite, and the creation of local cavities, the Schlotten.

EVOLUTION

In this section, we describe the numerical tool SOCKS3D (SOluble roCKS), which is used to describe the evolution of flow and permeability in an aquifer, which is in parts composed of soluble rocks. Our aim is to describe a situation as depicted in Fig. 12: The hypothetical model describes a simplified cross section similar to the Bad Frankenhausen region, with a single sequence of Permian Zechstein rocks (limestone, anhydrite, sandstone), offset by a fault. On the left side of the fault, the soluble rocks are in greater depth, and here salt is present on top of the anhydrite.
**MATHEMATICAL FRAMEWORK**

We use the numerical program package SOCKS3D, which is a new fork of our widely used KARSTAQUIFER package (see Kaufmann et al. 2010; Hiller et al. 2011; Kaufmann et al., 2012; Hiller et al. 2014, for more details), including the description of multiple rock types. The KARSTAQUIFER package has been used to simulate the speleogenetic evolution of an aquifer comprised of soluble and/or insoluble rock. Here, we will provide only a short model description, the reader interested in more details is referred to the references listed.

The model is based on a two-dimensional digital topography ($nx \times ny$ grid points), and then is extended into depth with $nz - 1$ layers, resulting in $nx \times ny \times nz$ nodal points. On the discretisation level, rock matrix is identified by 3D parallelepipedal elements, rock fractures by 1D linear bar elements, both represented with finite elements.

The SOCKS3D packages solves the following set of equations in a three-dimensional modelling domain describing a fractured, porous aquifer in soluble rocks under steady-state or transient flow conditions and a realistic topography:

$$\frac{\partial h}{\partial t} = \nabla \cdot \vec{q}$$

$$\frac{\partial \vec{V}}{\partial t} = \frac{p}{A} F(c)$$

$$\frac{\partial d}{\partial t} = \frac{m_{\text{rock}}}{\rho_{\text{rock}}} F(c)$$

The first equation is a conservation equation for flow, with $h$ [m] the hydraulic head, $\vec{q}$ [m/s] the Darcy velocity, $S_s$ [1/m] the specific storage, $\vec{V}$ [1/m] the nabla operator, and $t$ [s] time. The second equation is the transport equation for the calcium concentration $c$ [mol/m$^3$], with $P = \pi d$ [m] and $A = \pi d^2/4$ [m$^2$] the perimeter and the cross section of a circular fracture (not to confuse with $A_s$, the specific surface area), $d$ [m] the fracture diameter, $F$ [mol/m$^2$/s] the calcium flux rate, and $c$ [mol/m$^3$] the calcium concentration. The third equation describes the reactive enlargement of a fracture with $m_{\text{rock}}$ [kg/mol] the atomic mass and $\rho_{\text{rock}}$ [kg/m$^3$] the density of the soluble rock. Darcy velocity is given by

$$\vec{q} = K \nabla h,$$

with $K$ [m/s] the hydraulic conductivity.

For fracture elements, the hydraulic conductivity is given as

$$K_f^{\text{lamin}}(t) = \frac{\rho g}{10 \pi \eta} d(t)^2, \text{laminar}$$

$$K_f^{\text{turb}}(t) = \frac{2g}{f q_i} d(t), \text{turbulent}$$

with $\rho$ [kg/m$^3$] the density of the fluid, $g$ [m/s$^2$] the gravitational acceleration, $\eta$ [Pa s] the viscosity of the fluid, $f$ [-] the friction factor, and $q_i$ [m/s] the Darcy flow rate component in flow direction from the previous iteration step. Note that flow in the presence of turbulence becomes non-linear. For matrix elements, the hydraulic conductivity is derived from the Kozeny-Carman-relation

$$K_m(t) = \frac{\rho g}{10 \pi \eta} \frac{d'(t)^2}{l^* \Phi(t)^2}$$

with $\Phi$ [-] the porosity of the rock matrix, scaled between 0 (no pores) and 1 (only pores), $l^*$ a tortuosity factor accounting for the complex geometry of the flow through the pores, and $d'$ defined as typical grain size.

**MODEL SETUP**

The three-dimensional model extends 100 m in north-south and 200 m in west-east direction, and 20 m vertically, using 51x101x11 nodal points (Fig.13). With this setup, we simulate a section of the Zechstein deposits of the southern Harz Mountains, simplified with one depositional sequence.

The model has a 2 m resolution, with around 160,000 fracture and 50,000 matrix elements. While the hydraulic properties of the matrix elements are fixed to a hydraulic conductivity of $K_m = 10^{-7}$ m/s and a specific storativity of $S_m = 5 \times 10^{-6}$ 1/m, hydraulic conductivities of...
the fracture elements change with time due to dissolution and precipitation of the mineral species. Specific storativity of the fracture is low, $S_f = 10^{-8}$ $1/m$.

The model consists of up to three different rock types, from top to bottom sandstone, anhydrite, and limestone. Surface temperatures are set to $T = 10^\circ C$.

As boundary conditions, we define a fixed head boundary condition along the left side ($h = 10$ m, blue dots), defining a regional base level of the aquifer system. As input, a fixed inflow condition along the bottom part of the model along the right side is defined, mimicking a regional groundwater-flow component into the limestone layer ($q = 20000$ m$^3$/yr, $T = 10^\circ C$, $p_{CO_2} = 0.02$ atm, $c_{aq} = 1.01$ mol/m$^3$, $c_{in} = 0.90$ mol/m$^3$, red dots). Thus the inflow occurs into the limestone layer, while the local base level is the gypsum/anhydrite formation along the right side. This conditions mimic hypogene conditions for the evolution of cavities.

The resulting initial groundwater-flow pattern can be seen in Fig. 13: The head contours reveal flow from right to left, with only small local perturbations resulting from the initially heterogeneous fracture conductivities visible in the distorted head contours, which result from our assignment of initial fracture diameter values using a log-normal distribution, assigning an average value $d_{av}$ [m] and a standard deviation $d_{\sigma}$/m].

MODEL RUNS

With the geometrical, hydrological, and chemical properties defined, we discuss a set of model runs for the evolving aquifer, which are all run for 20,000 years. Our strategy is as follows: We start with a simple one-material limestone aquifer below a sandstone domain to realise how the carbonates of the Zechstein sequences become karstified, both in time and geometrically. We then add a gypsum layer between the limestone and the sandstone to understand the impact of dissolution of different soluble rock types, resulting in different time scales of the evolution of secondary permeability in both soluble rock sequences. We then swap the gypsum layer with an anhydrite layer to describe the simplified setup of the Zechstein sequence in the southern Harz region. Our aim is to understand the clogging of the anhydrite sequence due to the precipitation of secondary gypsum. As we will see, we efficiently block the anhydrite, and therefore we finally need to introduce conditions, which can lead to

![Fig. 13: Setup of numerical evolution model. Top: Schematic cross section. Bottom: Topography (transparent), limestone (red), anhydrite (grey), sandstone (blue). Inflow boundary (right, red dots), and base level (left, blue dots). Groundwater-flow head contours (colour-coded). The block in the background marks the different rock layers (for colour code see Fig. 13 top).](image-url)
localised enlarged voids in the anhydrite, resembling the Schlotten-type cavities.

**LIMESTONE AQUIFER**
The first model is made up of a 6 m thick limestone layer and a 14 m thick sandstone rock above. With this model, we introduce a standard model of karst aquifer evolution with its typical time scale of evolution.

The sandstone has fairly low permeability, and recharge is only via a regional flow component from the right into the limestone layer at depth, mimicking the regional flow component.

The initial fracture diameter distribution for both the sandstone and the limestone is based on a log-normal distribution with $d_{ave}=0.1 \text{ mm}$ and $d_{0}=0.05 \text{ mm}$. With this large standard deviation, we expect several competing preferential flow paths, as the distribution of initial fracture diameter values becomes wide enough to provide less direct flow path options.

Water entering the limestone layer from the right side is slightly aggressive, with $c_{aq}=0.90 \text{ mol/m}^3$ it has already attained 90% of the equilibrium values of $c_{eq}=1.01 \text{ mol/m}^3$ for this temperature / carbon-dioxide pressure setting for limestone. The evolution of this model is shown in Fig. 14 for two snapshots in time.

5,000 years into the evolution, several clusters of enlarged fractures have started developing from the right side towards the base level at the left side. Enlarged fractures have already diameter values in the cm-range. The different clusters compete for flow, and the bottommost cluster has already attained an advantage, when compared to the other clusters, which can be seen from the size of the cluster and the head distribution in front of it.

![Fig. 14: Limestone aquifer: Two snapshots in time, with fracture diameter and hydraulic head colour-coded. The block in the background marks the different rock layers (for colour code see Fig. 13 top).](image)
20,000 years into the evolution, the bottommost cluster has succeeded in capturing the majority of flow, and a preferential flow path connecting the right and the left side is established, with fracture diameter values in the 10 cm-range. Hydraulic pressure drops significantly, and the further evolution is deferred due to the lower flow velocities in this stage. This situation, called catchment-controlled evolution in the karst aquifer literature, limits flow through the evolving aquifer, as inflow is controlled by the amount of water coming from a limited catchment.

LIMESTONE-GYPSUM AQUIFER

We proceed with the second model, which is made up of a 6 m thick limestone layer, a 6 m thick gypsum layer, and a 8 m thick sandstone rock above. With this second model, we introduce a model of karst aquifer evolution made up of two adjacent layers of different soluble rock. The different properties of the limestone and gypsum rocks will cause a completely different evolution, as we will see below.

The model is recharged as above, via the right side of the limestone part, with water close to the calcium saturation for limestone under the chosen climatic conditions (\(c_{\text{eq\ limestone}} = 0.90 \text{ mol/m}^3, c_{\text{eq\ limestone}} = 1.01 \text{ mol/m}^3\)). Thus the incoming water can dissolve limestone (in the non-linear regime of the limestone flux rate), but has much more potential to dissolve the overlying gypsum, which with the chosen surface temperature has a calcium equilibrium concentration of \(c_{\text{eq\ gypsum}} = 14.57 \text{ mol/m}^3\). The evolution of this model is shown in Fig. 15 for two snapshots in time.

After already 300 years of evolution, the enlargement of fractures in the gypsum aquifer is substantial,

![Fig. 15: Limestone-Gypsum aquifer: Two snapshots in time, with fracture diameter and hydraulic head colour-coded. The block in the background marks the different rock layers (for colour code see Fig. 13 top).](image-url)
with clusters enlarged to the 10 cm size. After 20,000 years of evolution, the entire gypsum sequence is basically dissolved.

Hardly visible from the current perspective view and therefore only described is the fact, that in the limestone sequence almost no evolution of secondary porosity occurs. The reason for this behaviour is that water injected into the limestone along the right size of the model domain finds an easier flow path through the gypsum sequence, which through its fast dissolution becomes much more permeable than the limestone and attracts flow. Thus, the dissolution kinetics (gypsum versus limestone) entirely controls the evolution of this combined system of two soluble rocks, with gypsum having a clear advantage due to its much higher solubility.

LIMESTONE-ANHYDRITE AQUIFER
We, however, know that in the southern Harz Mountains the limestone is capped by anhydrite. While both gypsum and anhydrite are made up of the same material, considering the evolution of a combined limestone-anhydrite system is completely different, as we will see below in the snapshot shown in Fig. 16.

In this model, we keep the hydraulic boundary conditions as above, but in the anhydrite layer we allow for gypsum precipitation, once the calcium concentration passed the calcium equilibrium concentration for gypsum, $c_{eq}^{gypsum} \approx 15.34 \text{ mol/m}^3$. As we have already discussed in section 4, this combined anhydrite dissolution-gypsum precipitation can clog small fractures on time scales below a year. Thus it is not surprising to find an evolved aquifer, which after 20,000 years of evolution time resembles more or less the situation of the single limestone aquifer: A couple of enlarged clusters of fractures in the limestone part of the model, with the bottommost cluster on its way to win the competition. However, no evolution of secondary porosity in the anhydrite sequence, as here the water entering from the limestone layer below starts dissolving the anhydrite, but quickly passes the gypsum equilibrium concentration, resulting in gypsum precipitation. As the precipitated gypsum occupies a larger volume as the dissolved anhydrite, the small fractures initially present in the anhydrite clog and the anhydrite layer becomes almost impermeable.

With this model we have shown, how effective the anhydrite-gypsum conversion is in reducing the permeability of an anhydrite layer, resulting in properties more characteristic for an aquitard for the anhydrite, which is a common observation in nature.

LIMESTONE-ANHYDRITE AQUIFER WITH SCHLOTTEN-TYPE CAVITIES
Now we have to come back to our initial question, how do Schlotten-like cavities evolve in a combined limestone-anhydrite sequence as observed in the southern Harz Mountains. As we are able to clog the anhydrite recharged with an aggressive water source from below (hypogene setting), we need some conditions to allow for the evolution of secondary porosity locally. We have several options available, which can be discussed with our knowledge from section 4. Two options applicable to the southern Harz Mountains and the Kyffhäuser setting will be discussed:
- **Chemical control:** Coming back to Fig. 8, we have based our discussion of the effective clogging of the anhydrite layer on the different calcium equilibrium concentrations of gypsum and anhydrite, with around $c_{eq}^{gypsum} = 15 \text{ mol/m}^3$ and $c_{eq}^{anhydrite} = 45 \text{ mol/m}^3$ for a temperature of 10°C. This large difference in $c_{eq}$ and the volume increase between the dissolved anhydrite and

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**Fig. 16:** Limestone-Anhydrite aquifer: One snapshot in time, with fracture diameter and hydraulic head colour-coded. The block in the background marks the different rock layers (for colour code see Fig. 13 top).
the gypsum precipitate causes the clogging of the layer. However, when we increase the temperature of the solution to 30-40°C, this difference becomes very small, and the gypsum precipitation less efficient. We argue that ascending thermal water through a fault zone (as the southern Kyffhäuser Fault in Bad Frankenhausen) can cause this reduction in gypsum precipitation, thus leaving voids from enlarged anhydrite fissures open.

- **Structural control:** Another possible mechanism to create localised voids in the anhydrite can be derived from an observation in Fig. 11: The time scale for clogging an anhydrite fracture with precipitating gypsum depends on the initial width of the anhydrite fracture, larger initial fractures will clog later. This effect can be substantial, as small increases in initial fracture width already result in orders-of-magnitude delay in clogging the fracture. We argue that a fault through the Zechstein sequence, providing a larger initial permeability of the rock sequences, can allow fractures in the anhydrite to remain open and to enlarge to larger voids.

We will show that both hypotheses result in localised enlarged voids in the anhydrite layer in our simplified model. The model shown in Fig. 17 is based on the limestone-anhydrite-sandstone setup of Fig. 16, but with two vertical fault zones (shown as vertical semi-transparent planes).

The fault on the left side close to the base level has an initial fracture diameter of 0.001 m, with is one order of magnitude larger than the average initial fracture diameter of the other fractures (assigned with log-normal distribution). The fault on the right side of the model domain has an initial fracture diameter of 0.005 m, thus it is hydraulically more effective, when compared to the

Fig. 17: Limestone-Anhydrite aquifer with Schlotten-type cavities: Two snapshot in times, with fracture diameter and hydraulic head colour-coded. The two faults are shown as vertical semi-transparent planes. In the bottom figure, only fractures enlarged to the 50 cm size and above are shown. The block in the background marks the different rock layers (for colour code see Fig. 13 top).
other fault. While the fault on the right side is used as structural-control element, along the fault on the left side we inject water from below with a temperature of 40°C, $p_{CO_2}=0.02$ atm, and a calcium concentration of $c_{eq}=1.01 \text{ mol/m}^3$. Thus, this thermal water component is saturated with calcium with respect to the limestone chemical-control element. Pointing back to our Fig. 12, the left fault mimics groundwater ascending through the southern Kyffhäuser Fault in Bad Frankenhausen as the origin of the enlarged voids observed in the boreholes, while the right fault depicts a situation similar to the Elisabethschächter Schlotte with its relation to the collapse sinkhole on the surface.

10,000 years into the evolution, we observe a preferential enlargement of fractures in the limestone part of the rock sequence, but the pattern is different when compared to our other runs. The reason is the right fault, which attracts flow because of its larger initial fracture diameter values. The fault acts as local base level, from which new preferential pathways in the limestone start to evolve towards the base level on the left side. In the anhydrite part of the two faults, several anhydrite fractures remain open and enlarge width time to the cm-to-meter range.

In the second snapshot taken 20,000 years into the evolution, we just visualize fractures enlarged to at least 50 cm to clarify the substantial local growth in the anhydrite layer.

In the anhydrite part of the right fault, several anhydrite fractures remain open and become enlarged to the meter size, generating a local system of interconnected enlarged voids, similar to the Elisabethschächter Schlotte!

The evolution around the left fault, in which we inject thermal water, is different. Here, several isolated clusters of enlarged voids grow on an downstream of the fault, which quickly reach sizes in the 10 cm to meter range. Such voids are possibly observed in the boreholes lowered around the church Oberkirche in Bad Frankenhausen! We note, however, that here also water from the neighboring sandstone can provide additional aggressive solution, as in the case of the Numburg Cave.

CONCLUSIONS

In this final section, we review our models and summarize our key results in view of the questions asked in the introduction. We have asked three key questions in the introduction, which we now can answer:

1. Can we explain the evolution of the Schlotten-type cavities with a large-scale karst evolution model, accounting for dissolution in a combined carbonate-sulfate setting? Yes, we can. Our numerical modelling results suggest that in a setting where water seeps into the Permian Zechstein sequences through the carbonate layers (Zechsteinkalk, Stink-schiefer), it dissolves the anhydrite from below. However, the dissolution of the anhydrite is mainly occurring along the interface carbonate-anhydrite. Fissures in the anhydrite are usually clogged by gypsum precipitation. Only along prominent fault zones, flow is diverted and focussed into these hydraulically more permeable zones and anhydrite dissolution outpaces the gypsum precipitation. Thus, along these prominent fault zones, Schlotten-type cavities can form.

2. Can we explain the surface deformation observed in Bad Frankenhausen with numerical models of large-scale dissolution in the underlying sulfate layers? Yes, we can, at least indirectly through the evolution of large isolated cavities in near-surface layers along prominent fault zones, which develop in the anhydrite and due to their sizes (often in the meter range, as our models suggest), these voids are prone to collapse.

3. Can we estimate the role of aggressive mineral-rich brine water arriving from below through fault systems (hypogene setting)? We can also speculate that rising thermal water (as in our setup of one of the prominent faults) changes the chemical equilibrium conditions and thus enhances the dissolution of anhydrite, resulting in even faster development of large sub-surface voids.
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