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2020-11-27 Lotta Rubio Lind

**Quality assurance** 

2021-01-20 Lotta Rubio Lind (Approved)

# Notes; Quality assurance - LOT meeting 3

Date: November 27, 2020

### Participants:

Galson Sciences Ltd, Tim Hicks, Tamara Baldwin

**SSM**, Bo Strömberg, Henrik Öberg, Jinsong Liu, Michael Egan (participated the first part of the meeting)

**SKB**, Johannes Johansson, Magnus Kronberg, Magnus Westerlind, Lotta Rubio Lind

The meeting was held virtually using Skype.

SSM provided questions before the meeting which SKB gave written answers to and presented during the meeting. The questions and answers are all documented below. After each issue, the discussion that followed is summarized.

SKB was responsible for taking notes at the meeting. The notes were sent to SSM and SSM distributed the notes to Galson Sciences for a factual control before the notes were approved in SKB's document system.

It should be noted that SKB have added clarifying text after the meeting for issues 3, 8 and 18.

This meeting was the last in a sequence of three and the focus at the meeting was:

- (i) Management system & project management (2020-11-05)
- (ii) Retrieval, sampling, handling of samples & analysis (2020-11-13)
- (iii) Interpretation of results (2020-11-27)

# Quality management system

# 1. Selecting suppliers

What evaluation criteria do you used when selecting SKB's external supplier companies (e.g. KIMAB, Swerim, Clay Technology)? Do you have minimum requirements for respective company expertise and profile?

**SKB's reply/comment:** SKB prefers that the suppliers are ISO-certified. A supplier evaluation is performed before contracting a new supplier and is renewed regularly to assure that the company still meets SKB's expectations. In the evaluation the different criteria are described in 1056110 – Inköpsinstruktionen (Purchase instruction), and these criteria regard:

- Economy
- Quality assurance
- Environmental impact

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After delivery an evaluation of the new company used is done by the client according to a template provided by the procurement unit at SKB. For previously evaluated companies, additional evaluations should be made if called for by new experiences. These procedures are in place to ensure that past experiences are taken into account when considering a company for additional assignments.

Smaller companies and sole proprietorships do not necessarily have ISO-certifications and are then chosen for their unique expertise. The evaluation is then based on CVs, level of education, publications in the scientific literature, etc.

**Discussion:** SKB clarified that also for larger companies it can be specified which staff members at the supplying company that will perform the assignment. The competence of the individual is evaluated from CVs and descriptions provided by the companies. If the supplier, for some reason, has to change personnel for the assignment SKB has to approve the change. This is applicable for those assignments for which SKB requires specific personnel at the supplying company, and this procedure is stipulated the order.

#### 2. Collaboration with consultants

How long has SKB collaborated with KIMAB, Swerim, and Clay Technology?

**SKB's reply/comment:** The relationship between SKB and these companies has a long history. SKB (and its predecessor the KBS-project) has collaborated with the Swedish Corrosion Research Institute (Korrosionsforskningsinstitutet) and the Swedish Institute for Metals Research (Institutet för Metallforskning) at the Royal Institute of Technology (KTH) since 1977. Corrosion studies were performed from 1977, and creep studies started in 1984. These two institutes then merged and formed KIMAB, later Swerea KIMAB. A couple of years ago the company was divided into Rise KIMAB and Swerim.

Clay Technology was formed in 1988 and SKB has been using the company from the beginning. The company was formed by parts of SGAB (Sveriges Geologiska AB, a state-owned company formed in 1982) with which SKB collaborated before Clay Technology was formed.

**Discussion:** It was discussed how many companies there are in Sweden that perform these types of corrosion and material analyses relevant for experiments such as the LOT project. SKB knows of other companies in Sweden that can perform parts of the analyses but Rise KIMAB (Swerim) is the only one that can undertake the whole assignment. Using suppliers from abroad (which is done for other assignments) was considered, however, this alternative may also enhance the costs due to, for example, travel expenses. It may be noted that it was initially discussed to involve an academic research group abroad, but it was considered difficult due to the need for careful handling of samples during transportation. If using several suppliers, the effort in coordinating the suppliers will increase.

## 3. Report procedures

Do contracting companies provide internal company reports to SKB which are then further processed within SKB to SKB TR- or R-reports?

**SKB's reply/comment:** Yes, in some cases, depending on the degree of involvement of SKB's experts in the actual work done. When SKB experts are involved in the work they are normally involved at an early stage of experimental planning, analysis and reporting. In such cases the SKB expert is usually a co-author of the report. For other reports the only post-processing is editorial, i.e. typesetting and printing of the report.

**Discussion:** It was discussed how the LOT corrosion report TR-20-14 was written. Since it was a close cooperation from the beginning, Rise KIMAB did not make a separate report, rather it was written jointly from the start by the SKB and Rise KIMAB authors (see further the list of authors of TR-20-14). This has been the procedure for several SKB reports produced in collaboration between SKB and other companies, e.g. Rise KIMAB. Swerim provided a separate report for the TEM analyses, which were done at a late stage during the work (it was an additional separate order), which was included in the report TR-20-14 as an appendix. This section (as all other appendices) were subject to peer review with the rest of TR-20-14. The analytical measurements and results are to be considered as independent results which are entirely reported, and the conclusions are written mainly by SKB personnel with the support of the supplier.

It was discussed that for some assignments the supplier provides SKB with a draft report that SKB publishes in a SKB report series (TR-, R-, P-). When receiving the draft SKB goes through the draft (delivery control) and have a dialog with the author in order to clarify any ambiguities in the report and in some cases a new version with clarifications is requested before a formal review is performed of the report. The review of reports in the SKB report series are handled in the same way no matter if it is a TR-, R- or a P-report and with the same requirements on the review. The number of reviewers depends on the availability of reviewers with the right competence, sometimes it is necessary to combine several reviewers to fully cover the content/extent of the report. When choosing the reviewer/ers it is also important that the reviewer haven't been involved in the writing of the report. The reviewer/ers can be an internal staff member or an external consultant. After the review and before publication the report is adjusted to the SKB layout.

It was discussed if SKB have a written policy concerning openness of results. It is widely known at SKB (a SKB culture) that we should publish all of our results from scientific assignments in order to maintain trust from the authority and society, however, at the meeting SKB could not say if the openness-policy was clearly stated in the management system at present and would doublecheck this which has been done in the text below:

### Additional information after the meeting:

The SKB Policy refers to the four general principles that are expressed in SKB's Code of Conduct. One of the principles concerns *Openness*;

"Vår verksamhet drivs framåt av människor och samarbeten, och vi vet att transparens i ord och handling genererar förtroende och respekt. Vi är generösa med vad vi vet och delar gärna med oss av vår kunskap. Vi söker nya perspektiv, är innovativa och arbetar tillsammans med andra för att öka utvecklingstakten. När vi kommunicerar är vi proaktiva, transparenta, tydliga och relevanta."

"Our activities are driven by people and cooperation, and we know that transparency in words and deeds generate trust and respect. We are generous with what we know and willingly to share our knowledge. We seek new perspectives, are innovative and work together with others to increase the pace of development. When we communicate, we are proactive, transparent, clear and relevant" (unofficial translation).

Through the mandatory review of the RD&D programme reports, SSM oversees the development of management and disposal systems in the pre-licensing process. The review process includes opportunities for broad public participation in the development of the Swedish system for managing spent fuel and radioactive waste. The RD&D-programme can be found in the Management System for SKB in the part where the vision, goals and strategy of the company is described. In the latest SKB RD&D program published in 2019 it is stated in section *5.1.4 Review, openness and transparency* that:

"SKB's research is conducted on the basis of the requirement that research results should be correct, traceable, reproducible and relevant for SKB's mission. To achieve this, SKB has developed and applies procedures for quality assurance of the execution of research projects and tasks. There are also special procedures for quality assurance of safety assessments, which include approval of research results, data and models for use in the assessments, see also Section 5.4.

The fundamental principle is that SKB's research results will be published in the open literature to facilitate external review. Research results have, since the research programme was initiated in the 1970s, been published and will continue to be published in SKB's report series, which are available on SKB's website. Before they are published, the reports have undergone internal and/or external review in accordance with established procedures. Quality-assured data from SKB's site investigations, technology development and research are saved in databases and are available for the authorities in their review.

SKB also strives to publish relevant results in scientific journals and encourages its own personnel, as well as research institutions and consultants that SKB is collaborating with, to publish results. An independent review of the results is carried out through the peer review that takes place before publication. SKB's research results are also presented and discussed at scientific conferences and are published in the conference proceedings.

SKB is also working to disseminate research results outside the scientific community. For example by publication in popular scientific journals, publication on SKB's website and in SKB's paper Lagerbladet, themed evenings (mainly in Östhammar) and information at schools and universities."

### 4. External data deliveries

How are data deliveries from contractors controlled? That is, are data post-processed for recording in SICADA/use in reports?

**SKB's reply/comment:** Data is sent (by e-mail or via ftp) by the contractor and stored in SICADA, and a resource responsible for the work approves/releases the data in SICADA. With respect to post-processing, it depends on the data type. For example, temperature is stored as raw data, while other data are stored both as a raw data measurement files and calculated results in a template. For reports, calculated results are generally plotted or presented in tables. A general principle is that no data should be omitted. If it is obvious that e.g. a sensor is malfunctioning (for example showing unrealistic or unphysical values), it shall be documented that data from this sensor is removed and why.

**Discussion:** It was discussed how SKB handles identified errors in data already delivered and approved in the SICADA database. SKB replied that the supplier (internal or external supplier) then corrects the data sheet/table and send an updated version to the SICADA operators. The older version of data is then locked for use and only the updated data can be used/delivered for further use in, for example, modelling and reports. The SICADA operator also makes a SICADA bugreport in order to document the change.

It was also discussed how a contractor can access data from SICADA, this is done according to an instruction, using a template that is sent to the SICADA administration after which they provide approved data. If larger errors, errors that can affect model results and conclusions, are identified at a later stage, those who have received that data are informed about the changes made by the SICADA administration.

# Interpretation of LOT corrosion analysis and test conditions

### 5. Corrosion - function of time

What can be said about the extent of copper corrosion in the LOT test as a function of time considering all available data? Are there clear causes for variability apart from the apparent dependence on temperature? Could there be non-quantifiable causes for variation related to the design of the experiments and, if so, which are expected to be the most significant?

**SKB's reply/comment:** Corrosion on the warmest parts of the pipes was estimated to be of the order of  $10 \mu m$ , with a few  $\mu m$  deeper corrosion on the warmest parts of A3 (20 years) than for A2 (6 years). This is probably because  $O_2$ -induced corrosion proceeds beyond the depletion of  $O_2$  itself, since the intermediate oxidant Cu(II) prevails on much longer time scales (King et al. 2010, King and Kolář, 2019). However, it can not be excluded that the available amount of  $O_2$  differed between the test parcels. On the colder parts of the pipes, there was no significant difference between the A2 and A3 parcels.

For the coupons, gravimetric data is available from several test parcels and corrosion could be quantified with standardised methods. Two observations may be noted regarding variability;

- 1) The corrosion depths varied from 0.6 to 4.7 μm and were highest in the 1-year parcels (A0 and S1), intermediate in the 6-year parcel (A2) and lowest in the 20-year parcels (S2 and A3). The corrosion was thus inversely proportional with time, and correlation of measured corrosion depths (mass-loss) with temperature was very weak, indicating that there were other factors than time and temperature that controlled the corrosion in these experiments.
- 2) In the test parcel ABM 5 (another Äspö bentonite experiment, very similar to LOT), the variation in corrosion depth by mass-loss was 2.3 to 5 μm (Gordon et al. 2018), despite the fact that all specimens had the same temperature (ca 80 °C) and were exposed for the same time (5 years). It is not known exactly what causes this variability in LOT and ABM (within and between test parcels), but the most reasonable explanation is considered to be heterogeneous distribution or transport of O<sub>2</sub> and/or heterogeneous resaturation of the bentonite clay. When comparing the LOT and ABM test parcels it should be noted that the water supply systems were different. Concerning the different LOT parcels the natural flow from the boreholes may have contributed to spatial variations in the saturation process. Variations in the initial surface conditions of the copper specimens, especially for the pipes, can not be disregarded, e.g. regarding roughness and/or the initial oxide layer.

**Discussion:** It was discussed how the temperature gradient along the copper pipe and the gap between the bentonite and the pipe affects the results. Considering the gradient, the corrosion rates on the warmer parts would in the LOT case be overestimated and underestimated for the cooler parts. The coupons were imbedded from start (no gap) and there was no artificial temperature gradient and these effects can thus not be directly compared.

The general decrease of integrated corrosion rates between A2 and A3 (and S2) was discussed with reference to the report Johansson et al. (2014).

If there are other oxygen consuming processes in the experiment, such as microbial activity, was discussed. The conditions in the LOT experiment were not favourable for the microbes due to initial unsaturated conditions (low water content) and later high temperature and density so the oxygen consumption from bacteria should be considered low. This is further discussed in Issue 8.

### 6. Corrosion - highest temperatures

SKB has not measured the extent of corrosion on the part of the copper tube exposed to the highest temperatures. Why? What could be expected if these tube sections were analysed?

**SKB's reply/comment:** The corrosion of the warmest parts of the copper pipes has been measured based on the estimated concentration of copper in the bentonite (e.g. in block 9 and 11 for A3, see Table 3-19 in Johansson et al. 2020). Examination of the pipe surfaces was done for the pipes in blocks 21-23, since these were carefully handled during extraction of the coupons contained within the same bentonite blocks, in order to avoid damaging (scratching) the surfaces.

It may be noted that the temperature at the pipe surfaces examined metallographically was 70-80 °C for A3, which is not far from the peak temperature of 95 °C for any copper canister in the KBS-3 repository. On the other hand, the maximum temperature in A3 was 120 °C, which is higher than what is relevant for the canister in the repository. Initially, the canister surface in the repository will be ca 50 °C and it will take about a decade to reach the maximum temperature (95 °C) after which the temperature decreases slowly.

It is possible that the warmest part of the A3 pipe could be rougher or have slightly deeper pits than the examined part (depending on how the extent of these pits is influenced by corrosion). However, it should be noted that the deepest pits found on the pipe in A3 were only 25  $\mu$ m, while the expected range of pit-depths during initially oxidising conditions in the KBS-3 repository would be up to a few hundred  $\mu$ m, and with a low probability of pits even up to ca 1 mm (Briggs et al. 2020). Pipe material from LOT S2 and A3 has been saved for the possibility of further metallographic examination. It is, however, unlikely that the confirmation of deeper pits on the warmest parts of the pipe in A3 would change any conclusions regarding localised corrosion under the initially oxidising conditions of the repository.

### 7. Copper in bentonite - XRF

Do the bentonite copper content measurements by XRF provide an accurate measure of the local extent of corrosion on the copper tube? Would consideration of corrosion products left on the copper tube significantly affect the results?

**SKB's reply/comment:** The estimate from the XRF analysis of clay near the coupons showed that the mass of copper found in the bentonite clay was in the range 0.011-0.018 g. This is not far from the mass-losses of the coupons which were in the range 0.009-0.024 g, and which correspond to corrosion depths of 0.6-1.3 µm. The "error" in this comparison would be the adherent corrosion products which are included in the gravimetric analysis but not in the Cu in clay analysis. However, the oxide films on the coupons were generally very thin, typically around 1 um, at some positions less, and locally sometimes up to a few µm (see e.g. Fig 4-2 and Fig B-38 in Johansson et al. 2020). There was no apparent difference in oxide films on the coupons A3/K (50 °C) and S2/N (30 °C), as seen in the micrographs in Appendix B. It may be further noted that there was no obvious difference in the thickness of adherent corrosion products on the examined pipe samples (Appendices F and G in Johansson et al. 2020). Some micrographs show adherent layers of bentonite (high Si), while corrosion products were typically around 1 µm thick, sometimes less and locally sometimes more corrosion products in or near pits or surface defects. This was despite the fact that the temperature was higher in A3 (70-80 °C) than in S2 (50-60 °C), and despite the fact that both lighter and darker areas were examined for each test parcel (Figure G1-G5 in TR-20-14).

Laboratory studies in the Canadian program have shown that the estimate of corrosion from copper content in the clay, sometimes need to be multiplied by a factor to match gravimetric results. This factor has been estimated to range from 1 to 19 in the different experiments made

(Litke et al. 1992, King et al. 1997). While the experiments in Litke et al. (1992) used a 50:50 sand-bentonite mixture and had an initial  $O_2$  inventory of ca 30 mol/m<sup>2</sup> copper area, the experiments in King et al. (1997) used compacted bentonite without sand and had less than 1 mol/m<sup>2</sup> copper, which is closer to the initial conditions in LOT. The factors derived in the study by King et al. (1997) varied in most cases within the range 1-1.7. The fact that many of these values were close to 1 is in agreement with the general picture from LOT in which the adherent oxide film was mostly very thin.

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**Discussion:** It was discussed that the available amount of  $O_2$  is important for the thickness of the oxide film. SKB has discussed the Canadian experiments with one of the authors and it was noted that in the study by Litke et al. (1992), the copper surfaces in many cases had an appearance with thick deposits of blue-green corrosion products, while the copper surfaces in King et al. (1997) appeared more similar to the copper surfaces in LOT.

It was also discussed that the steep copper profile in the bentonite clay is indicative of Cu(II) being adsorbed. This is discussed in the Canadian studies cited and further references therein. SSM brought up a discussion about the form of copper in the clay and mentioned that ion exchange between copper ions and Na and Ca in the bentonite is known to occur. The possibility of complexation was also mentioned by SSM.

### 8. Oxygen consumption - microbes

It has been claimed in the LOT context that the initial oxygen content available during the LOT-experiments was rapidly consumed by microbial processes before the oxygen had a chance to react with the copper. What is SKB's position regarding this claim? What knowledge can be gained from SKB's other experiments regarding microbial oxygen consumption, pyrite oxidation and other reactions in bentonite clay?

**SKB's reply/comment:** The bentonite clay in LOT was initially cold and unsaturated, conditions under which microbial O<sub>2</sub> consumption is very inefficient (Birgersson and Goudarzi 2018, Giroud et al. 2018). When the heaters were turned on the temperature of the copper surface increased rapidly from rock temperature to ca 40 °C and then continued to increase. As indicated by laboratory studies, copper corrosion under aerobic conditions and temperatures 50-80 °C can occur rather rapidly (Litke et al. 1992, King et al. 1997). This supports the view that O<sub>2</sub> may have been consumed by corrosion before the microbial consumption started. Oxidation of pyrite is not significant at rock temperature but is known to occur at 55 °C. The occurrence of pyrite oxidation would however be difficult to confirm, which is further discussed in the reply to Q9.

**Discussion:** The activation energy for the oxidation of pyrite was discussed and that there is a temperature effect on pyrite oxidation. Different corrosion experiments were also discussed, and whether any of the experiments have measured the corrosion that occurs at a very early stage. SKB mentioned that the Canadian studies cited in the response to Q7 measured corrosion already after one month, and depending on the conditions they sometimes measured rather high corrosion rates on this short timescale, as compared with the rates measured after several months or years in the same experiments. This shows that the initial corrosion may occur much more rapid than the integrated corrosion rates measured for longer exposures.

SKB has started two new experimental studies in order to improve the detailed description of corrosion during the very early stage. One of the experiments is a small scale laboratory test conducted at a lab in the UK. The aim of this experiment is to measure of corrosion at different stages during the depletion of  $O_2$ . The second is a medium scale test conducted at the Äspö HRL and aims to follow the development of the gas-phase composition in a setup with copper and

unsaturated bentonite clay. Both experiments will focus on unsaturated conditions. The UK experiment is expected to finish during 2022. The first part of the Äspö experiment will be reported during 2021.

It was discussed if other studies made, for instance on corrosion on copper roofs, could give more information on the early corrosion. SKB pointed out that such studies differ from repository conditions in several ways, for example regarding the availability of O<sub>2</sub>, the chemical composition of rain water due to atmospheric pollutants, and temperature. It was further noted that in studies of such objects as for example copper roofs, corrosion is seldomly measured on shorter timescales.

### Additional information after the meeting:

It may be noted that the first sentence in SKBs reply to Q8 was a bit unprecise. Neither of the cited studies (Birgersson and Goudarzi 2018, Giroud et al. 2018) concerned microbial effects in the clay so it should rather be said that any type of O<sub>2</sub> consumption was inefficient under the unsaturated conditions of the experiments. Concerning LOT, microbial activity is not expected to have occurred, initially due to the unsaturated conditions (see for example SKB TR-20-08), and later due to the high clay density and high temperature.

Regarding the studies of the composition of the gas-phase cited in the reply to Q8, it may be appropriate to clarify what was actually observed. The study by Giroud et al. (2018) reported on experiments where open glass vials were filled with granulated bentonite and placed into stainless-steel bottles, subsequently sealed and airtight. All bottles were kept at laboratory temperature of ca 23 °C. The partial pressure of O2 decreased by a few percent over a time period of 15 days, however, the same pressure drop was seen also for inert gases like N2 and Ar, and the results were therefore interpreted as sorption rather than a reaction. Similarly, Birgersson and Goudarzi (2018) reported on a test that was conducted in isothermal (room temperature or 50 °C), isolated conditions, and involved only bentonite pellets. The test showed no noticeable O2 consumption at room temperature and very little at 50 °C over a period of 1 year. These studies show that, although inorganic reactions may consume O2 trapped in bentonite clay under some conditions, the reactions are slow at low temperatures, at least for an unsaturated conditions.

## 9. Oxygen consumption - other processes

If all the oxygen was not consumed in the experiment, could a certain fraction of the available oxygen have been consumed by processes other than copper corrosion? If so what is an approximate estimate of this fraction? Is there a formation of sulphate which could indicate pyrite oxidation?

**SKB's reply/comment:** Formation of sulphate from pyrite oxidation is likely to occur, however a bit difficult to verify, as the bentonite originally also includes gypsum (Ca-sulphate) and the sulphate moves in the bentonite, typically towards the central heater. Possibly the total sulphate content could be measured. Another strategy could be to look for disappearance of pyrite in XRD, however this is not trivial as the pyrite content is low already at the start. In Callovo-oxfordian (COX) clay, which is a clay stone rich in pyrite, signs of pyrite oxidation in the field experiment ABM1 was observed based on increasing Fe(III)/Fe(II) ratio, however this was not studied in detail as it was not the purpose of the study (Svensson and Hansen, 2013).

### 10. Availability of oxygen

Could there have been a slow leakage of oxygen from the tunnel through seals, fractures in the rock, the EDZ, through holes next to cables and pipes etc.? (Onset of reducing conditions does not *per se* exclude in-leakage if the prevailing redox reactions consuming oxygen are reasonably well buffered).

**SKB's reply/comment:** The weld between the bottom plate and pipe was pressure tested with He gas before the experiments were installed (Sandén and Nilsson, 2020). It seems unlikely that significant leakage could have occurred through concrete and clay, however, considering the large number of cables going through the concrete and around the test parcels it can not be completely excluded that some leakage could have occurred.

**Discussion:** The importance of cables was discussed for the different experiments in order to better understand the potential for leaking of oxygen for the different experiments. The experiments FEBEX (Nagra) and Prototype (SKB have a fairly large number of cables going through the clay system and have shown that leakage of O2 may be difficult to avoid on the scale of the experiments. The LOT experiment has fewer cables installed but SKB cannot definitely rule out that leakage of oxygen took place. SSM pointed out that a slow in-leakage of O2 would generally be difficult to detect if there are processes (like for example corrosion) that consumes O2 and keeps the concentration within the experiment low. In a situation in which the consumption rate of O2 is more rapid than the supply by mass transfer, a scenario which cannot be outruled for the case of LOT, the O2 concentration could in principle not only be low but be close to zero. Such slow mass transfer could still have some relevance on time-scales of several years.

### 11. Added groundwater - influence on oxygen

Could groundwater added to the LOT experiments have consumed oxygen through its reducing capacity? Is there knowledge of the reducing capacity of this groundwater (e.g. Fe(II) content)? If so how could addition of groundwater have affected the oxygen content available for copper corrosion?

**SKB's reply/comment:** The Fe(II) concentration of the groundwater that is supplied to LOT is low (around 70 ppb). Since the exchange of groundwater with the LOT-packages is limited the Fe(II) can be assumed to have negligible effect on the oxygen consumption.

**Discussion:** SSM mentioned that one way to determine if Fe(II) have been oxidised during the experiments would be to look for ferric precipitates, since Fe(III) would be very insoluble under the prevailing conditions. Other reducing agents mentioned and deemed to be less important were dissolved manganese and hydrogen sulphide. It was concluded that Fe(II) should control the redox potential of the groundwater.

The water filling procedure was shortly discussed and it was clarified that no pumps were used, see issue 12 below.

# 12. Water filling procedure

Were all slots and gaps completely filled with groundwater at the onset of the experiments (i.e. when the heaters were switched on)? For how long were the slots and gaps open? What was the pathway for water to enter the gaps (i.e. from titanium tubes and rock)? What is known about the groundwater inflow rates in the "deposition holes"? Is it possible to identify any water conducting feature in spite of the drilling near the holes?

**SKB's reply/comment:** The (natural) water inflow rates were measured at different depths in the bore holes for the LOT test parcels. The general conclusions from the pilot hole characterization program were that the water inflow was low, and that the water inlet points were few in all holes (therefore it was decided to use groundwater from the nearby hole, see below). The average water inflow to the borehole where test parcel S2 should be installed was ca 1 ml/min, which is nearly two orders of magnitude higher than for the other boreholes (Table 2-1 in Sandén and Nilsson 2020).

Before the start of the heaters and the water filling procedure, the Ti-tubes were open to the Äspö tunnel for ca 4 months in the S2 and A3 test parcels (Table 3-1 in Sandén and Nilsson 2020).

After installation of the test parcels in the holes, there was a remaining air-filled gap between bentonite blocks and rock surface with a width of 10 mm, and another air-filled gap between bentonite blocks and the central copper tube with a width of ca 1 mm. These gaps were slowly water filled (on February 2, 2000) in parallel with the onset of heating. The filling was made with groundwater from the adjacent borehole (HG0038B01) by use of the fixed installed Ti-tube and bottom filter placed in the sand below the parcel and close to the rock wall. The valve connected to the bottom filter was closed at the time when the groundwater had reached the uppermost filter (filter in block 32), which until then was open to the Äspö tunnel. This filter was then instead connected to the groundwater supply. This inflow point is assessed to simulate a point inflow from a water bearing fracture in the rock. The test parcels have had access to pressurised water from this point inflow during the entire test duration.

**Discussion:** It was discussed what determines the time to reach full saturation in the LOT experiments and that the high temperature at the pipe surface is an important factor. It was also discussed that the period before contact is established between the bentonite and the copper pipe surface could be important in the description of the development of corrosion in LOT.

In connection to Q10 it was discussed whether it is possible that leakage of air could have occurred through factures connected to the tunnel floor. SKB can not definitely exclude the possibility.

SSM made the point that there could be residual oxygen in bentonite and sand even after saturation that could diffuse to the copper surface.

SSM mentioned that the plugs in deposition tunnels may be a source of oxygen leakage during construction of the repository and this has been evaluated in one of the complementary investigations made by SKB after SR-Site

### 13. Water inflows in the LOT boreholes

Are there known differences between the LOT holes regarding water inflow and saturation? What is the time scale of full resaturation and is there an appreciable variability between the holes?

**SKB's reply/comment:** The natural flow was described in the reply to Q12. Reaching full saturation and swelling pressure of the bentonite clay generally took years. For blocks 8 and 14 in LOT A3, near the hottest part of the copper pipe, the process was particularly slow, taking 4-6 years to reach the final pressure (Sandén and Nilsson 2020). This can be compared with LOT A2, in which the saturation in blocks 8 and 14 took less than two years (Karnland et al. 2009).

**Discussion:** It was discussed if the slow resaturation of the bentonite near the warmest part of the copper pipe can result in the higher extent of corrosion there compared with the colder regions. The high temperature does not only influence the rate of  $O_2$  consumption by corrosion, but may also keep the initial gap between copper and bentonite clay open for a longer time due to dryer conditions and thus slower resaturation, as compared with colder areas. It was concluded that it is plausible that both these effects of higher temperature have contributed to the distribution of corrosion over the copper surfaces in LOT.

# SKB's position with respect to copper corrosion and alternative interpretations

### 14. Corrosion kinetics

At what point does the oxic corrosion of copper become diffusion controlled due to the formation of corrosion products? Is diffusion control of relevance (or mixed kinetics) as a rate limit for the initial phase of LOT with oxic corrosion?

SKB's reply/comment: Diffusion control could mean either anodic diffusion control (i.e. the diffusion of either copper cations across a passive Cu<sub>2</sub>O film or of dissolved Cu(I) diffusing in the buffer pore water) or cathodic diffusion control (i.e. diffusion of O<sub>2</sub> either through the buffer or across a precipitated corrosion product layer). Different experiments have indicated different types of diffusion control. For example, measurements of the depletion of O<sub>2</sub> with time in King et al. (1997) clearly indicated diffusion-limited O<sub>2</sub> consumption, which suggests that the cathodic reaction is diffusion controlled. In that study, the limiting step was probably diffusion through the buffer, rather than through a thin corrosion product film, as the buffer was compacted in situ (i.e. there were no air-filled gaps). On the other hand, anodic transport control may have been possible in Litke et al. (1992) because of the high amounts of O<sub>2</sub> in the tests. While the experiments in King et al. (1997) may be applicable to the coupons in LOT, the situation for the copper pipes in LOT is more complicated due to the fact that air-filled gaps may have been present initially and when these had been depleted of O<sub>2</sub>, the oxidant, O<sub>2</sub> or Cu(II), was only available by diffusion through the bentonite clay.

**Discussion:** The extent of different initial phases were discussed. The corrosion of the pipe surfaces may initially have been under kinetic control, while a similar initial phase for the coupons may have been much shorter (diffusion control is likely to have been rate limiting for a comparatively longer duration), since these were embedded in bentonite clay from the start. This is probably an important difference for the comparison of corrosion between pipes and coupons, although temperature variations were also important.

### 15. Corrosion - temperature dependence

What is the temperature dependence of copper corrosion rate for the relatively unexposed copper surfaces put in the LOT experiment (activation energy)?

**SKB's reply/comment:** For the copper coupons, the correlation of corrosion depth (or rate) with temperature was very weak. This may reflect that the corrosion was controlled by diffusion through the clay. However, it must also be noted that the temperature did not differ as much between the coupons as for the pipes. For the copper pipes, there was a clear temperature effect on the measured corrosion depths (and thus on the integrated corrosion rates) as shown in Figure 4-1 in Johansson et al. (2020). Since the integrated corrosion rates are determined for a very long period of time (20 years) during which the corrosion process goes through different mechanistic phases (O<sub>2</sub>, Cu(II), sulphide) with different kinetics, it is unclear what a detailed kinetic analysis of activation energies based on integrated rates would mean.

**Discussion:** It was discussed if there are publications on reaction rates for oxygen and copper. SKB replied that there are such papers in the general scientific literature, see for example King et al. 1995 and references therein. SKB mentioned that the reports cited in response to Q7 contain kinetic data for  $O_2$  induced corrosion of copper at a copper/bentonite interface.

### 16. Corrosion in sand

What are the differences in between corrosion of copper in sand and in bentonite (of relevance for interpretation of corrosion on the tube bottom plate)?

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**SKB's reply/comment:** This has not been investigated, since sand is not a material that will be used in the KBS-3 repository, it was only used in the specific LOT experiments. The verification of solid Cu(II) corrosion products adherent to the bottom plate, but not on other copper surfaces in LOT, may be due to ion exchange between corrosion products and bentonite clay, which does not occur with sand.

**Discussion:** It was discussed how the contact with bentonite clay allows ion-exchange between corrosion products and the clay, which is not possible with sand. This could be an explanation for the accumulation of Cu(II) corrosion products that was observed on the bottom plates but not on any surfaces in contact with bentonite clay. SSM noted that evaporation at the warm copper surface may be important for the formation of chloride containing corrosion products. SKB pointed out that one circumstance that may be of importance is that one of the Ti-tubes ran out in the sand just below the bottom plate. It is not known if Cu(II) corrosion products were formed under the bottom plates of earlier test parcels, e.g. A2, but in general the appearance of the pipe in A2 resembled the pipes in S2 and A3. In A2, small amounts of the solid Cu(II) phase paratacamite were found on one of the coupons. This has not been found on the coupons in S2 or A3. SSM also made the point that there is a great deal of uncertainty about the saturation process but the sand beneath the bottom plates would have saturated quickly. SKB agreed in general but notes that this depends on how fast water reached the sand in the deep bottom hole.

### 17. Corrosion kinetics, oxic-anoxic conditions

It has been claimed in the LOT context that corrosion of copper will not slowdown in response to a shift between oxic and anoxic conditions. What is SKB's position on that?

**SKB's reply/comment:** Experimentally, it has not been possible to distinguish the time dependence of the corrosion rate in the presence of O<sub>2</sub> versus the presence only of Cu(II), although the latter phase is expected to be longer (King et al. 2010). However, when O<sub>2</sub> has been depleted and the resulting Cu(II) intermediate has reacted, the only oxidant available in the deep ground water environment is sulphide, for which transport limitations will control the long-term corrosion rate. The integrated corrosion rates that can be calculated from corrosion depths obtained from copper coupons and pipes in the LOT series, clearly indicate that the integrated rate is decreasing with time. This is discussed in sections 4.2.1 and 4.2.2 in Johansson et al. (2020). As regards the possibility of corrosion of copper in pure, O<sub>2</sub>-free water, it is SKB's clear position that this process occurs to an extent that is compatible with established thermodynamic data, which is completely negligible in the LOT context, see further Hedin et al. 2018 and SKB 2019, Chapter 4.

### 18. Corrosion products, oxic-anoxic conditions

It has been claimed in the LOT context that the formation of the main detected corrosion product cuprite ( $Cu_2O$ ) in LOT A2 and S2, and also other typical corrosion products in similar experiments (malachite  $Cu_2(CO_3)(OH)_2$ , para-atacamite  $Cu_2Cl(OH)_3$ ) may be related to anoxic conditions. What is SKB's position regarding this issue? What are SKB's key arguments? Has knowledge of the thermodynamic properties of these phases been utilized to address such issues?

**SKB's reply/comment:** In order to form corrosion products, in such amounts to precipitate solid phases, some other oxidising species than water is needed (the dissolved copper oxidised by water is negligible; Hedin et al. 2018, SKB 2019). For this evaluation thermodynamic data is used to investigate what phases that can form under different conditions. For example, the copper oxide corrosion product Cu<sub>2</sub>O is formed under conditions where the redox potential Eh > 0 V SHE (at 25 °C, for pH up to 8, and a total copper concentration in solution of 10<sup>-6</sup> M). To form Cu(II) solid phases even higher potentials are required. This can be seen from Pourbaix diagrams (Puigdomenech and Taxén 2000), but can also be evaluated with more detailed speciation

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calculations, that has been done e.g. for corrosion in high chloride concentrations (Lilja et al. 2021), still using the same type of thermodynamic data. The claims that the solid corrosion products are related to anoxic conditions, may depend on the used definition of "anoxic conditions". For the LOT experiment (as well as in the bentonite buffer in the repository) the redox potential would be determined most probably by iron compounds and be in the range of that in groundwater (around -0.2 V; SKB 2011), when all O<sub>2</sub> is consumed. Such potentials are not compatible with formation of either copper oxides or paratacamite (or malachite). The observation of oxides and paratacamite in LOT is though consistent with thermodynamic data for the initial conditions with O<sub>2</sub> (high Eh). (Malachite would be formed with a water composition with slightly lower chloride concentration and higher carbonate concentration.) Even if the development of reducing conditions does not support formation of e.g. paratacamite, this does not exclude that compounds already formed may remain kinetically stable also during reducing conditions.

**Discussion:** Regarding the terminology, SSM asked if the groundwater redox potential of -0.2 V (SHE) should be described as anoxic, reducing or if both descriptions are plausible. It is SKB's understanding that the terminology is not strictly defined and that both the terms anoxic and reducing could be used to describe the groundwater environment.

The thermodynamic and/or kinetic stability of different corrosion products formed in the presence of O<sub>2</sub>, such as Cu<sub>2</sub>O, CuOH, Cu(OH)<sub>2</sub> and Cu<sub>2</sub>Cl(OH)<sub>3</sub>), was discussed, for conditions under initial depletion of O<sub>2</sub>, as well as for the long-term sulphidic repository environment. SSM has also pointed out, to which SKB agrees, that it is possible that the prolonged presence of solid phases normally associated with oxidising conditions (such as paratachamite) could be due to slow dissolution kinetics in combination with slow mass transfer, rather than kinetic stability for the conditions and time scale of these experiments. In this context there was also a discussion about thermodynamic stability of cuprite together with copper both in the presence and absence of other redox affecting species such as hydrogen gas and dissolved hydrogen sulphide.

SSM asked whether SKB could provide a reference for thermodynamic data on paratacamite, a polymorph of Cu<sub>2</sub>Cl(OH)<sub>3</sub>, which has been identified in field tests such as LOT. SKB replied that such data were applied for thermodynamic calculations underlying the safety assessment SR-Site (Puigdomenech and Taxén 2000).

### Additional information after the meeting:

Since the meeting, SKB has looked into the data used in Puigdomenech and Taxén (2000) and a few things needs to be clarified. First of all, the data applied in Puigdomenech and Taxén (2000) was for atacamite, which is a different polymorph of Cu<sub>2</sub>Cl(OH)<sub>3</sub>. It is noted that SKB has not always been strictly consistent or precise when referring to the polymorphs of Cu<sub>2</sub>Cl(OH)<sub>3</sub>, as is also the case for older literature. Further, naturally found paratacamite often contains some zinc. However, as discussed by Pollard et al. (1989) the  $\Delta G^{\circ}_{f}$  (Gibbs free energy of formation) values for the three polymorphs of Cu<sub>2</sub>Cl(OH)<sub>3</sub> are very close, with paratacamite being the most stable, closely followed by atacamite, and botallakite being the least stable of the three (there is a also other crystallographic variants, like clinoatacamite, Jambor et al. 1996). The difference in  $\Delta G^{\circ}_{f}$ between paratacamite and atacamite given by Pollard et al. (1989) was less than 7 kJ/mol and in the same paper, comparison was made with earlier studies showing the same trend in the relative thermodynamics of the polymorphs. The value for  $\Delta G^{\circ}_{f}$  assigned to atacamite in Puigdomenech and Taxén (2000), lies within the range of values for atacamite and paratacamite given in other studies (Woods and Garrells, 1986). The subtle differences between the  $\Delta G^{\circ}_{f}$  values reported for the polymorphs of the Cu<sub>2</sub>Cl(OH)<sub>3</sub> does not change the stability area to any noticeable extent. It has thus no implication for the interpretation of the observation of Cu<sub>2</sub>Cl(OH)<sub>3</sub> in LOT or the thermodynamic evaluation of corrosion products to be expected in the spent fuel repository.

The structure, thermodynamics, and chemical stability of the CuOH phase mentioned above, has been studied experimentally as well as by theoretical methods, see further Soroka et al. 2013 and Korzhavyi et al. 2012.

### 19. Corrosion products- Cu(II) phases

In King's review, a presence of Cu(II)-phases is suggested. According to established knowledge, ordinary copper corrosion in an oxic environment involves formation of a double-layer of Cu<sub>2</sub>O and CuO on top of the metal, with the monovalent oxide forming closest to the metal surface. To what extent does the presence of the buffer affect these reactions? What is the significance of an absence or presence of Cu(II)-phases for the interpretation of LOT? It has been claimed in the LOT context that a lack of Cu(II) suggests that corrosion has been anoxic. What is SKB's position on this issue?

**SKB's reply/comment:** As elaborated on in the answer to Q18, the oxidation state of corrosion products, more oxidising conditions (more positive Eh) supports the formation of Cu(II) compounds (e.g. CuO in fresh water or paratacamite, Cu<sub>2</sub>(OH)<sub>3</sub>Cl, in chloride-rich groundwater), while less oxidising conditions (less positive Eh) results in the formation of Cu(I) corrosion products (e.g. Cu<sub>2</sub>O or CuCl<sub>2</sub><sup>-</sup>). In aerobic systems, it is common to get a bilayer structure with a Cu(II) phase on top of Cu<sub>2</sub>O, and this was found in the copper-bentonite experiments in both Litke et al. (1992) and King et al. (1997). The bentonite tends to adsorb the Cu(II) and, hence, lower the tendency for Cu(II) precipitation at the copper surface. In Litke et al. (1992) the initial [O<sub>2</sub>] was high (30 mol/m<sup>2</sup> Cu) and the average corrosion depth was ca 40 μm, and precipitated Cu(II) was clearly visible as blue-green corrosion products. In King et al. (1997), the initial [O<sub>2</sub>] was lower (<1 mol/m<sup>2</sup> Cu) and the fraction of Cu(II) in the corrosion product was in the range 1-56%, while the steep copper profiles in the bentonite clay adjacent to the copper surfaces implied the presence of Cu(II) adsorbed in the clay. LOT seems to be more similar to King et al. (1997), i.e. no visible adherent Cu(II) products but steep Cu profiles in the adjacent bentonite clay. See also answer to Q16.

### 20. Hydrogen embrittlement

One of key issues raised by the Swedish Land and Environment Court is hydrogen embrittlement of copper. Are there any results of relevance to this issue for the LOT S2 and A3 phases?

**SKB's reply/comment:** Analysis of hydrogen levels in the copper pipes from LOT S2 and A3 (section 3.4.2 in Johansson et al. 2020), shows that hydrogen is only present in or near the surface of the material, which is also where corrosion products and bentonite deposits are present. Comparison of bulk levels of hydrogen of the S2 and A3 pipes with a reference pipe showed very low levels of hydrogen and no significant difference between the pipes. Having said this, it may be noted that the only process which could hypothetically lead to hydrogen uptake by copper under the conditions in LOT, is the long-term corrosion by sulphide (H<sub>2</sub>S/HS<sup>-</sup>), and as is evident from the results in Johansson et al (2020), the extent of sulphide corrosion in LOT was very limited.

**Discussion:** The form of hydrogen in the surface deposits was discussed. The role of hydroxide groups and water associated with bentonite deposits and corrosion products was discussed, as well as the amount of adsorbed water on particles and surface deposits. SKB mentioned that results similar to LOT have been obtained in a Round-Robin study of the performance of the Leco method for hydrogen measurements on uncorroded copper specimens (Granfors 2017).

### 21. FEBEX - LOT

Are there any key findings from the FEBEX experiments that are of relevance for the interpretation of the LOT results?

**SKB's reply/comment:** The results from FEBEX are regarded as part of the general scientific knowledge of copper corrosion under early repository conditions dominated by O<sub>2</sub>-induced corrosion (Wersin and Kober 2017). The results, e.g. regarding corrosion products and corrosion morphology, are discussed in Johansson et al. (2020) and will be integrated in the safety assessment PSAR for the spent fuel repository.

A few detailed results may be worth mentioning in this context. The mass-loss of the FEBEX coupon 4A2 corresponded to an average corrosion depth of 8.5  $\mu$ m, while the deepest pit observed on the FEBEX coupon 4A1 was ca 90  $\mu$ m. These numbers are both larger than the corresponding values for the coupons in LOT, which may be due to a combination of higher temperature, larger clay volume (more  $O_2$ ) and an extended oxic period in FEBEX as compared with LOT. It may be noted that FEBEX differed from LOT in several aspects, for example there was no sulphur on the coupons in FEBEX (while low levels of sulphur were found on all copper surfaces is LOT), there were clear signs of blue-green Cu(II) corrosion products on the coupons in FEBEX (not present generally in LOT S2 or A3), and FEBEX is thought to have had a longer oxic period initially (possibly due to leakage from the tunnel).

# Concluding remarks by SKB

In the context of the above questions from SSM, SKB would like to make the following summarizing points regarding the interpretation of copper corrosion in the LOT series:

### Purpose with the LOT experiment

- Since the LOT experiments were designed with the purpose to study the behavior of bentonite clay under repository-like conditions, the experiments can only give limited information about copper corrosion. In order to be clear about this, a number of limitations of LOT, if considered as a corrosion experiment, are discussed in Section 1.5 in Johansson et al. (2020).
- There is no information available concerning the initial development of redox conditions in the LOT experiments, and, consequently, the development of the corrosion process can not be described in detail.

#### Oxygen consumption

- There is a large uncertainty considering the amount of  $O_2$  (from air) that has been available for corrosion.
- It is considered likely that the copper pipe surfaces consumed a large fraction of the initially available O<sub>2</sub> when the heaters were turned on and during the following months during which the temperature of the pipes increased from 40 °C to 90 °C or higher.
- It is considered likely that the copper pipes consumed most of the O<sub>2</sub> in the unsaturated clay system faster than chemical processes in the clay (e.g. oxidation of pyrite) since the copper surface had a higher temperature than the outer parts of the bentonite blocks.
- Microbial consumption of O<sub>2</sub> is not regarded as efficient in unsaturated bentonite clay.

### Evaluation of corrosion

Since the roughness of the copper surfaces in LOT were not characterized prior to the
experiments (one of the limitations of LOT if regarded as a corrosion experiment),
localized corrosion can only be pessimistically evaluated by assuming that the topography
observed is solely due to corrosion. However, even under that assumption, the observed
pits are consistent with the expected corrosion morphology under initially oxidizing
conditions in the repository.

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- Given the uncertainties due to the limitations of LOT as a corrosion experiment, the results
  are consistent with the conceptual corrosion model applied by SKB in the assessment of
  post-closure safety for the spent fuel repository.
- A group of researchers has suggested that copper corrodes in pure O<sub>2</sub>-free water, i.e. by reactions between copper and water molecules, to an extent that by far exceeds that predicted by established science and thermodynamic data (Hultquist et al. 2015). It is emphasized that this hypothesis can not be evaluated in a complex field test like LOT, in which air (O<sub>2</sub>) was initially available and since groundwater is not pure water. This hypothesis has instead been thoroughly evaluated in careful laboratory experiments, which have been published in the scientific literature, see further Hedin et al. 2017, 2018, Ottosson et al. 2017, and references therein.
- One of the observations made in LOT S2 and A3 was a slightly difference appearance and the formation of Cu(II) corrosion products under the bottom plates of the test parcels. This difference between the bottom plate and other copper surfaces in the test parcels is attributed to different reactivity at the copper-sand interface, as compared with the copper-bentonite interface. Since a copper-sand interface will not be present in a KBS-3 repository, these observations are not regarded as directly applicable to the assessment of copper corrosion in the spent fuel repository.
- The detailed evaluation of corrosion in SKBs safety assessment is based on numerous studies (laboratory experiments, models, and field tests) of different aspects of copper corrosion under repository conditions (radiation, O<sub>2</sub>, sulphide, microorganisms, groundwater etc) and is not dependent on a single experiment like LOT, especially since it was not designed for detailed studies of corrosion.

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Notes Author

2020-12-03 Lotta Rubio Lind

# Notes; Quality assurance - LOT meeting 2

Date: November 13<sup>th</sup>, 2020

Participants: Galson Sciences Ltd, Tim Hicks, Tamara Baldwin

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SKB, Johannes Johansson, Magnus Kronberg, Magnus Westerlind, Lotta Rubio

Lind

The meeting was held virtually using Skype.

SSM provided questions before the meeting which SKB gave written answers to and presented during the meeting. The questions and answers are all documented below. After each issue, the discussion that followed is summarized.

This meeting was the second in a sequence of three and the focus at the meeting was:

- (i) Management system & project management (2020-11-05)
- (ii) Retrieval, sampling, handling of samples & analysis (2020-11-13)
- (iii) Interpretation of results (2020-11-27)

# Retrieval, sampling, handling of samples and analysis

### 1. Pre-characterisation

TR-20-14 states up front that copper coupons were 'not intended or prepared for detailed corrosion analysis', but the 1998 LOT test plan (IPR-99-01) stated that the copper coupons and 'interesting' parts of the central copper tubes will be analysed to evaluate the mean corrosion rate, and to identify pitting corrosion and corrosion products. Some detailed corrosion analysis has been undertaken, although various caveats are noted in TR-20-14, such as lack of pre-characterisation of the surfaces of the copper coupons and tubes.

- a. Why was the pre-characterisation not done?
- b. How significant are the uncertainties associated with this lack of pre-characterisation?

That is, do these uncertainties significantly affect confidence in results?

# **SKB's reply/comment:**

- a. The importance of pre-characterisation at the microscopic level was probably not realised at the time of initiation of the experiments. In the 1990s, SKB's assessment of localised corrosion was mainly based on literature studies of pitting of copper pipes, archaeological artefacts etc.
- b. The uncertainty is difficult to quantify, but it should be noted that the reference coupons had pits of similar magnitude as the corrosion coupons, despite the fact that their average corrosion depth was only about 10% of the corrosion coupons. Similar results have been obtained by comparison of corroded and reference specimens in ABM 5 (TR-18-17), a test

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parcel from another experimental series similar to LOT. It may be further noted that even newly prepared and polished coupons, displayed pits of tens of µm (TR-18-14).

It is thus possible that the topography observed in LOT is due to initially occurring defects of mechanical origin that has later been affected by the corrosion process.

**Discussion:** It was discussed if it is possible to distinguish between pits formed by corrosion and mechanically induced pits. In the report TR-18-14 it is obvious what has caused the pits on the unexposed (reference) coupons. But for the LOT experiment it is difficult to say for certain due to the lack of pre-characterisation.

Probabilistic modelling of localized corrosion during initially aerobic repository conditions was discussed in the context of field tests like LOT and Febex.

Differences between Febex and LOT were discussed; e.g the FEBEX experiment was probably exposed to oxygen for a longer time period than LOT, (both due to the clay volume and possibly due to leakage via cables).

It was concluded that SKB assesses the measured pit depths on the LOT specimens conservatively by assuming that all pits are results of localised corrosion.

# 2. Milled or polished side

What is the significance of having a milled side and a polished side to the copper coupons?

**SKB's reply/comment:** The polished side was probably intended for the evaluation of localised corrosion. The milled side was examined with LOM since it was judged to be potentially more reactive than the polished side and because it more closely resembles the rough KBS-3 canister surface (P-12-22, P-13-50, P-17-11).

Examination of cross-sectioned coupons with SEM was done on the polished sides for one coupon from each test parcel. It may be noted that newly prepared and polished coupons (SiC, grit size P4000), also displayed pits of tens of µm (TR-18-14).

**Discussion:** It was discussed if there would be any advantage of polishing the copper canister. It may be noted that the main reason for localised corrosion would not be surface topography but rather passivation of the surface. Also, in the repository (as well as in field tests), localised corrosion during initially oxygenated conditions is a process limited by mass-balance.

### 3. Deviations from activity plan

Were any deviations from the activity plan for division and retrieval of the parcels necessary?

**SKB's reply/comment:** With respect to retrieval, there was the previously discussed issue with rock edges between boreholes which prompted additional drilling of core holes (meeting 1, question 8).

There were no deviations from the activity plan for division of the parcels.

**Discussion:** SKB handled the challenges with the seem drilling straightness during retrieval without affecting the copper samples, but it did increase the costs slightly and it took more time to retrieve the test parcels. The corrosion analysis was controlled by internal steering documents at Kimab and Swerim ,not by SKB's activity plans. There were no major deviations from the original plan for the corrosion analysis. The pickling method used in the gravimetric analysis had to be

tested and adjusted (as described in TR-20-14), however, that was not totally unexpected and is not regarded as a deviation in that sense.

### 4. Hand tools

Hand tools were used to extract the coupons so as not to damage or scratch them. What tools were used and was damage avoided?

**SKB's reply/comment:** First of all, the original design drawings for the LOT parcels were used to identify the positions of the coupons. Next, a metal detector was used to verify the position to an accuracy of about a few cm.

Finally, a rubber hammer and a wedge made of wood were used to remove the clay. When the edge of the wedge came close enough to the position of the coupons, the remaining clay typically divided in a way that part of the coupon was exposed.

**Discussion:** Two lessons learned from earlier retrievals of large-scale field tests were that it is difficult to remove compacted bentonite clay and that it is easy to scratch or damage small specimens inside the bentonite. SKB used these experiences and took several precautions when extracting the copper coupons. For example, the position of the copper coupons were determined from the original drawings from the assembly of the test parcels and the exact positions were verified using a metal detector. In this way, the extraction took longer time but it was successfully done, all eight coupons were retrieved without any damage.

### 5. Calibration

Were there any issues identified with regard to calibration and are calibration records available?

### SKB's reply/comment:

- Analytical balance (Rise KIMAB): Calibrated annually.
- XRD (at Rise KIMAB): Calibration of the instrument is done on regular basis by measuring Corundum NIST standard sample. The peak position identification and offset calculation is then done using EVA Diffraction program available by Bruker.
- SEM-EDS (at Rise KIMAB): Typically, the calibrations for magnification and energy positions in the EDS spectra are checked every year during maintenance. Normally, there is no need to correct them since they vary very little over time. For EDS you will know directly if the calibration is wrong, since all peak positions will be off. Either everything is wrong or nothing. An experienced operator will notice if anything is wrong.
- TEM (at Swerim): Magnification calibration, i.e. the scale bar for imaging and also for diffraction in the TEM, may not be more accurate than +/-5%. For diffraction, typically calibratation/normalization is done using a known phase before extracting the data, e.g. pure Cu.
- Regarding XRF for bentonite analysis, the instrument has an internal calibration named Omnian from Panalytical.
- SEM for bentonite had an internal calibration function.

Regarding test sensors there are also calibration protocols available.

**Discussion:** Overall the heating and temperature sensors have operated well. The heating system was updated with new components in September 2014 which resulted in a small drop in temperature, however, it does not affect test results in any significant way. This is reported in TR-20-11.

Calibration of scales at the laboratory at Äspö was discussed. SKB's water chemistry laboratory is an accredited laboratory and has routines for these types of calibrations which are also implemented in the Material research laboratory where the majority of SKB's internal bentonite analyses are done (the SEM is located at the Canister laboratory).

#### 6. O content

The tube copper had a higher O content than the Cu-OFP used in the copper disposal canisters. What is the significance of the difference in Cu grade in terms of possible corrosion rates and mechanisms?

**SKB's reply/comment:** The difference is probably of low significance but we found it appropriate to mention. However, no conclusions in the report TR-20-14 are based on this difference. The main difference is the content of P and O. Cu susceptible to H sickness typically has  $\sim 10^2$  wt-ppm O.

	O (wt-ppm)	P (wt-ppm)
Cu-OFP	<5	30-70
SS 5015-04	~30	150-400

**Discussion:** The difficulties of comparing different field experiments were discussed. SKB considers it difficult to compare coupons and pipes in the LOT experiment since they have to be examined with different method due to their size and degree of pre-characterisation.

SSM mentioned that preliminary results based on electrochemical potentiodynamic measurements suggest that pure copper is more prone to passivation as compared with OFP-copper. There are, however, still no definite results.

### 7. Reference materials

Regarding the reference materials:

- a. Was any pre-characterisation work done on the reference coupons and reference tube?
- b. Have estimates been made of the rate and type of any corrosion expected on the reference coupons and tube during dry storage? In Section C1.2, it is noted that there is cuprite on the reference coupons.
- c. Table 3-3 of TR-20-14 indicates that, during storage, the reference coupons corroded more than the test coupons. Was this expected, or is it possible that what is seen on the reference coupons are surface defects that could be present on manufacture?
- d. Would there be any benefit in looking further at the surface of newly prepared copper to understand its characteristics and surface defects on manufacture (as noted in Section 4.3)?

### SKB's reply/comment:

- a. No pre-characterisation was made (see reply to Q1).
- b. Photos of the reference coupons and pipe are shown in Fig A-11 and A-16 in TR-20-14 and it's evident that some oxidation has occurred during storage.
- c. The reference coupons were analysed gravimetrically and the total mass-loss corresponded to 0.07 and 0.16  $\mu m$  corrosion for the two coupons (Table 3-2, TR-20-14). The data in Table 3-3 are pits/defects; these are not necessarily related to corrosion as explained in reply to O1 and O2.
- d. In general, yes, to facilitate interpretation of future field tests. Especially older field tests in which the importance of microscopic pre-characterisation wasn't fully realized. Must be

done by preparing new coupons according to the same method as the corroded specimens, e.g. regarding cutting technique and polishing depth and/or fineness.

**Discussion:** It was discussed whether it would actually be useful to evaluate newly prepared surfaces, since it would require detailed documentation of the preparation of the specimens in order to be comparable.

SKB mentioned that the preparation of specimens of the MiniCan experiment is well documented, but for the LOT experiment SKB is not sure if it would be possible or even meaningful due to interpretation difficulties. The LOT coupons were prepared at Studsvik more than 20 years ago.

### 8. Microbes

Regarding microbes:

- a. IPR-99-01 stated that microbial populations in groundwater will be analysed before emplacement and at the end of the experiment and that bentonite samples will be examined for microbial populations. Was information on microbial populations obtained?
- b. How was information at the start of LOT recorded and stored, and was information about preparation of the bacteria samples managed separately to result in the information being lost?

### SKB's reply/comment:

- a. Microbes in Äspö groundwater have been studied in several projects and close to the LOT site is the Microbe site with measurements reported in *IPR-00-36 The microbe site*, *Drilling, instrumentation and characterization*. With respect to groundwater measurements, no specific measurements were made in LOT, so available data is from these nearby experiments. It should be noted that there are quite large local variations.
- b. Data were stored in SICADA from the start. For example, there is 47 SICADA activity Ids stored regarding the first two pilot parcels. Information regarding bacteria in the bentonite plugs is not lost; the work which has been done is described in *TR-00-22 Long term test of buffer material*. *Final report on the pilot parcels* and *TR-09-29 Long term test of buffer material at the Äspö Hard Rock Laboratory, LOT project. Final report on the A2 test parcel*. Bacteria in the plugs at the termination of the pilot parcels are reported in TR-00-22, chapter 7. Furthermore, Appendix 1 in TR-09-29 covers the bacteria work for A2. By mistake data concerning bacteria placed on the copper coupons at installation were not stored. That work was part of LOT and was not managed separately (however done by another contractor, Micans, but it should be noted that SKB is always responsible for the final documentation).

**Discussion:** SKB have been in contact with he supplier that prepared the coupons with bacteria before test installation. It is considered likely that the bacterial media contained sulfate reducing bacteria (SRB), since these are considered as the microorganism having the most potential influence on the corrosion process, however, by mistake the documentation of this information has been lost.

It was discussed if any studies of bacteria could have been done on the sand that is underneath the test parcels but since copper does not interact with sand in the KBS-3 system, it has not been studied at this stage.

It was discussed why there was an 8 m pilot hole drilled when the test parcels are 4 m. Hydraulic testing is reported in TD-99-25 Äspö Hard Rock Laboratory, Long term test of buffer material, Hydraulic tests in five boreholes in the G-tunnel of Äspö HRL, however, it has not been

documented why 8 m pilots were used, possibly to get a better picture of the area close to the experiments.

### 9. Saline water

The formation water supplied to the boreholes was observed to become more saline and alkaline over the duration of the test. Is there an explanation for these changes and could they influence copper corrosion mechanisms or bentonite behaviour in any significant way?

**SKB's reply/comment:** SKB has not done an in-depth analysis on the water in the area closest to LOT, however, 'upconing', when more saline, deeper, water rises towards the tunnel system is frequently observed at Äspö and is most likely the explanation also in this case.

With respect to affecting the bentonite the changes will have only marginal effect on the bentonite properties, see for example TR-06-30 Mineralogy and sealing properties of various bentonites and smectite-rich clay materials

There is very little Cl on the copper surfaces, typically <1 at-% and a few observations of up to  $\sim3$  at-% locally. This is similar to (or less than) what was found in LOT A2 (TR-09-29) and Prototype Repository (P-12-22).

**Discussion:** It was discussed why there is so little Cl at the copper surface. SKB states that the salinity is difficult to compare with previous test parcels since they were not analysed in the same way or to the same extent as LOT S2 and A3.

### 10. Monitoring corrosion potential

Why was monitoring of the corrosion potential of the copper or redox potential not attempted during LOT?

**SKB's reply/comment:** In general, the main focus of the LOT series was not copper corrosion. However, attempts were made to measure corrosion rates electrochemically by an electrode system installed in the test parcel LOT A2. Rate measurements were made but there was no recording of  $E_{corr}$  (Rosborg and Pan, Electrochimica Acta 53 (2008) 7556–7564).

E<sub>h</sub> was regarded as practically unfeasible to measure in compacted bentonite in the 1990s, especially in a field test where it is difficult to install and maintain a reference electrode on long time-scales.

### 11. SEM cross-section analysis

Tube samples were chosen for SEM cross-section analysis based on visual appearance (Section 2.3). What criteria were used to judge the area to select?

**SKB's reply/comment:** It was decided to sample "type-areas" based o their visual appearance, for example light (Cu coloured), dark/black, and with grey deposits (could be bentonite and/or gypsum). This is shown in figures G1-G5 in TR-20-14. As exemplified for area 1 for the pipe sample from A3, several cross-sections were examined for each area (Fig G6 in TR-20-14).



**Discussion:** SKB believes that sampling "type-areas" was an appropriate way to choose samples for the analysis. It should be noted that the darkest parts are not necessarily the most corroded, since corrosion products may in some cases adhere to the bentonite surface in contact with the pipe.

SKB has examined parts that were exposed to temperatures relevant for the repository. The pipe sections in blocks 21-23 were also selected for practical reasons, that is, they were in the same bentonite blocks as the coupons. Other parts of the pipes have been stored at Äspö HRL and could still be analysed.

It was discussed whether the thermal gradient in the experiment can affect the corrosion depth (and rates). Due to this gradient the oxygen will be transported from the cooler part and it can cause a higher corrosion rate at the warmest part of the pipe. This could mean that the corrosion rate is overestimated for the warmer part, while underestimated for the cooler part, as compared with the situation without an axial thermal gradient along the pipe. It was noted that such an axial thermal gradient would not be present in the repository, where the temperature increase is caused by spent fuel residual heat rather than an electric heater.

### 12. Mass loss measurements

What is the accuracy of the mass loss measurements?

**SKB's reply/comment:** The accuracy of the gravimetric analysis done at Rise KIMAB was +/-0.00009 g.

**Discussion:** The pickling process was discussed. It is a standardised methodology using a reference copper sample.

### 13. XRD results

Regarding the XRD results, Figure C-2 shows a peak at position 42 for coupon A3/K. The peak is attributed to  $Cu_2S$ , but isn't it  $Cu_2O$ ? Also, on a close look, the peak at position 30 seems to coincide with a small  $Cu_2O$  peak rather than bentonite clay. Any correspondence with components of bentonite clay does seem very weak.

**SKB's reply/comment:** It is a mistake in the report, the peak at 42 corresponds to Cu<sub>2</sub>O. Cu<sub>2</sub>S peaks should be at 37, 46, and 48. Peaks are weak but present on several samples.

SKB agrees that the peak at 30 seems more like Cu<sub>2</sub>O than bentonite, although we don't see that this is commented on in the report. Other peaks from bentonite are present in the diffraction patterns for some coupons, e.g. the peak at 22 for coupon A3/K and A3/I.

**Discussion:** The precision of the methods used was discussed broadly.

SKB clarified that the peak at 22 corresponds to bentonite. The bentonite is not evenly distributed over the coupon surface and the samples have to be washed with deionized water before the analysis otherwise the bentonite layer would be too thick.

### 14. C contamination EDS

What is the source of the C contamination noted in the EDS results (e.g. Figure C-46)?

**SKB's reply/comment:** The contamination is typically a few monolayers on the surface, covering the whole sample surface, but sometimes more. In SEM and TEM, the electron beam attracts adsorbed carbon and hydro carbons, so that carbon migrates to the beam and is there cracked and builds up a layer of carbon in the area which the beam scans. During an EDS analysis, the beam scans the same area for a long time, so a lot of carbon can build up. In SEM, the vacuum is lower, so even more carbon can end up on the sample from parts inside the chamber.

It may be noted that EDS data both including and excluding C is presented in Appendix C. There is no major impact on the evaluation of the levels of other elements due to C (See e.g. Fig C-46).

#### 15. Zinc contamination

Has any reason been found for the detection of zinc in the reference coupons?

**SKB's reply/comment:** The source of Zn on the reference coupons is unclear, we did not identify a source of zinc, and nothing in the handling of the samples at Rise KIMAB should have contaminated the samples. Possibly, there was an unknown source for Zn in the laboratory at Clay Technology where the reference specimens were stored for over 20 years.

# 16. Bentonite composition or fingerprint

For the EDS analysis (Section 3.3), would it be possible to identify a bentonite composition or fingerprint that would enable the bentonite component to be removed from the EDS results to give clearer focus on the corrosion product composition?

**SKB's reply/comment:** Rise KIMAB has replied; the problem is that the quantification is not so accurate when there are several phases in the same position/pixel. Also, the bentonite doesn't seem to be homogeneously distributed. It contains a lot of small nanoparticles, so the composition is not the same in different positions. There is sometimes water present. What you get is only a mean composition, which may vary spatially. Therefore, you don't know what composition to deconvolute and the EDS results should mainly be used in a qualitative way.

### 17. EDS analysis Cu<sub>2</sub>S

The EDS analysis (Section 3.3.1 and Appendix C) does appear to consistently indicate a  $Cu_2S$  phase at the surfaces of the coupons that were at cooler temperatures (block 30) but not at the surfaces of the coupons that were warmer (block 22). Is there any explanation for this? Is it statistically significant? The results for tube sample S2 (Figure 3-20) appear to contradict this so perhaps it is not significant.

**SKB's reply/comment:** There is some support of this observation in the EDS analysis made for larger areas (mm<sup>2</sup>) of the coupons (Table 3-1). The same trend is not seen in the corresponding data for the pipes (Table 3-5), however, the data set is small.

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Table 3-1. EDS data of copper coupons at 100× magnification (at%).

Coupon	Block	0	Si	s	Cu	Ca	Fe
A3/I	22	50.8	12.1	2.0	22.6	3.1	0.4
A3/J	22	68.7	3.2	0.7	23.7	1.6	0.2
A3/K	30	28.7	5.7	7.8	53.9	0.3	0.2
A3/L	30	34.9	8.7	8.7	42.4	0.4	0.3
S2/M	22	40.2	10.5	2.2	37.3	0.5	0.3
S2/N	22	37.3	10.3	1.5	44.3	1.4	0.3
S2/O	30	38.7	9.3	5.9	38.5	0.5	0.3
S2/P	30	40.9	11.5	7.3	33.3	0.4	0.4

Table 3-5. Selected EDS data of copper pipe samples at 100× magnification (at%).

Sample area	0	Si	S	Са	Fe	Cu
A3 pipe "white"	46.2	10.5	12.3	21.1	1.1	3.5
A3 pipe "dark"	29.1	16.9	4.2	1.2	1.1	38.2
S2 pipe "Cu"	16.4	7.9	1.3	2.1	0.7	61.6
S2 pipe "dark"	23.4	16.0	3.7	1.2	1.4	41.8

**Discussion:** It was concluded during the discussion that the small number of observations makes it difficult to draw any conclusions. It was mentioned that sulphide comes with the groundwater, which may not have been evenly distributed in the test parcels.

### 18. Diffraction analysis

The diffraction analysis (Section C1.4.3) appears to provide a means of identifying the composition of corrosion products, but the discussion is complex and the d values to compare with 1/d do not seem to be provided. Will a more detailed explanation and results be published?

**SKB's reply/comment:** Swerim has replied; It is not possible to unambiguously determine which phase(s) exist in the samples as there are so many small particles close to each other and each diffraction pattern obtained usually contains information from several different phases. When these phases have similar d-values and in many cases almost the same, it is not possible to distinguish which phase it is with our TEM equipment. This is further affected by the fact that it is not possible to get a better accuracy than about +/- 5% on all values. The accuracy does not depend on the calibration of the instrument but on a number of factors during measurement and postprocessing of data.

**Discussion:** SSM and SKB agree that TEM is an interesting method for trying to distinguish between different corrosion products. SKB will continue exploring this method.

### 19. Figure D-7 Cu<sub>2</sub>S

In Figure D-7, it is not convincing that Cu<sub>2</sub>S is being indicated as present on coupon S2/P. Is the figure showing the reflections discussed in Section C1.2 rather than a signal for Cu<sub>2</sub>S?

**SKB's reply/comment:** Yes, but on peak 49 there is a small "shoulder", which is Cu<sub>2</sub>S, according to Swerim. It is not very clear but it is an indication. SKB noted that the stoichiometry of Cu<sub>2</sub>S is

also in agreement with the Cu:S ratio obtained from TEM-EDS, which supports the interpretation that Cu2S was formed.

## 20. The purpose of the H measurements

What is the purpose of the H measurements? Is it to provide evidence to support arguments that hydrogen embrittlement doesn't occur?

**SKB's reply/comment:** The purpose of the H measurements was to investigate whether there was any uptake of H in the copper material, which could potentially lead to embrittlement effects.

**Discussion:** It was discussed whether H-loading could occur in the experiment. During sulphide corrosion it would in principle be possible, but in this experiment the extent of sulphide corrosion was very low (which is expected due to the low sulphide concentration in the groundwater and the slow inflow of groundwater through the bentonite).

## 21. Timescale for oxygen consumption

Could the measured amount of corrosion and the expected rate of corrosion give an indication of the timescale for oxygen consumption in LOT?

**SKB's reply/comment:** The rate of corrosion has probably changed significantly during the initial period of exposure. Integrated corrosion rates are discussed in TR-20-14 for both coupons and pipes, however these will underestimate the initial corrosion rates and overestimate the long-term rates. Comparison with data from LOT A2 shows a clear decrease in the integrated corrosion rates for all types of copper surfaces.

It may be noted that laboratory experiments with copper in bentonite under aerated conditions reported gravimetrically determined corrosion rates of  $100\text{-}200~\mu\text{m/y}$  for the first month at  $50\text{-}80^{\circ}\text{C}$  (Litke C et al., 1992. A mechanistic study of the uniform corrosion of copper in compacted clay and soil. AECL-10397, AECL, Canada.) This implies that the depletion of  $O_2$  could be rapid. However, this doesn't mean that the environment becomes reducing, since Cu(II) may be present for longer periods (TR-10-67, TR-18-08).

**Discussion:** This issue was only discussed briefly and was planned to be discussed further at the third meeting (2020-11-27).

### 22. Coupon S2/P

Are there any further views on why coupon S2/P was less corroded than other coupons?

**SKB's reply/comment:** Not really, but it may be noted that the corrosion of gravimetric specimens in earlier LOT test parcels and the similar ABM 5 experiment showed variation both within and between test parcels, which is also discussed in section 4.2.1 in TR-20-14.

**Discussion:** Why this variation occurs was briefly discussed. It is unclear if variation in transport conditions or chemical variations can have an effect.

# 23. Correction factor (the issue was added at the meeting)

**Discussion:** The correction factor mentioned by Peter Szakalos was discussed. This means that corrosion estimated from measurements of copper in the clay needs to be multiplied with a factor in order to capture the total corrosion depth, i.e. including adherent corrosion products- SKB has measured copper in the bentonite adjacent to coupons and adherent corrosion compounds at the surface of the coupon samples. This was then compared with the mass loss of copper from the

sample. This method performs quite well, since the adherent layer of corrosion products was generally very thin (ca 1  $\mu$ m). SKB did not apply a correction factor. It was also noted that the estimate of corrosion from copper in the clay is not a standard method.

This issue will be further discussed at the third meeting.