Quantitative imaging of the 3-D distribution of cation adsorption sites in undisturbed soil

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Abstract. Several studies have shown that the distribution of cation adsorption sites (CAS) is patchy at a millimeter to
 centimeter scale. Often, larger concentrations of CAS in biopores or aggregate coatings have been reported in the literature. This heterogeneity has implications on the accessibility of CAS and may influence the performance of soil system models that assume a spatially homogeneous distribution of CAS. In this study, we present a new method to quantify the abundance

and 3-D distribution of CAS in undisturbed soil that allows for investigating CAS densities with distance to the soil

- macropores. We used X-ray imaging with Ba²⁺ as a contrast agent. Ba²⁺ has a high adsorption affinity to CAS and is widely
 used as an index cation to measure the cation exchange capacity (CEC). Eight soil cores (approx. 10 cm³) were sampled from three locations with contrasting texture and organic matter contents. The CAS of our samples were saturated with Ba²⁺ in the laboratory using BaCl₂ (0.3 mol L⁻¹). Afterwards, KCl (0.1 mol L⁻¹) was used to rinse out Ba²⁺ ions that were not bound to CAS. Before and after this process the samples were scanned using an industrial X-ray scanner. Ba²⁺ bound to CAS was then visualized in 3-D by the difference image technique. The resulting difference images were interpreted as depicting the Ba²⁺
- 20 bound to CAS only. The X-ray image-derived CEC correlated significantly with results of the commonly used ammonium acetate method to determine CEC in well-mixed samples. The CEC of organic matter rich samples seemed to be systematically overestimated and in the case of the clay rich samples with less organic matter the CEC seemed to be systematically underestimated. The results showed that the distribution of the CAS varied spatially within most of our samples down to a millimeter scale. There was no systematic relation between the location of CAS and the soil macropore
- 25 structure. We are convinced that the here proposed approach will strongly aid the development of more realistic soil system models.

1 Introduction

Soil particle surfaces possess functional groups that are negatively charged. These interact electrostatically with cations and

- bind them reversibly (cation adsorption sites, CAS). Typically these CAS are positively correlated with the clay and organic 30 matter (OM) content (Murphy, 2015). The amount of CAS per mass of soil is referred to as the cation exchange capacity (CEC). It is commonly measured in cmol(+) kg⁻¹ soil using Ba²⁺ or NH₄⁺ to replace retained cations. The CEC determines agricultural soil fertility and the nutrient retention capacity (Cardoso et al., 2013; Murphy, 2015) Furthermore, it influences heavy metal retention (Bhattacharyya and Gupta, 2008; Gomes et al., 2001) and the mobility of cationic pollutants in soils
- 35 (Figueroa-Diva et al., 2010; Gevao et al., 2000; Porfiri et al., 2015).

The content and quality of clay or OM varies within soils depending on the bedrock, soil type, pedologic history, land use practices, vegetation cover, climate factors and biological activity (Guo and Gifford, 2002; Horn, 1987; Leue et al., 2010; Teferi et al., 2016). Illuvial clay mineral accumulation in argic horizons on aggregate surfaces and macropore walls and the formation of clay cutans lead to a spatially heterogeneous distribution of clay (FAO, 2014). Horn (1987) attributed a

- pronounced increase in fine material and CEC at aggregate surfaces to the aggregates shrinking and swelling activities, 40 leading to separation of fine and coarse material. Biopores (e.g. earthworm borrows, root channels) often create preferential flow paths and hot-spots of biological activity with higher OM contents compared to the surrounding soil matrix, resulting in a spatially heterogeneous distribution of OM and CAS often in relation to the location of macropores (Bundt et al., 2001a, 2001b; Kögel-Knabner et al., 2008; Nielsen et al., 2015). Thus, spatial variation of clay minerals and OM content in the soil
- profile affects the spatial distribution of CAS (Bundt et al., 2001b; Ellerbrock and Gerke, 2004; Horn, 1987). A higher 45 density of CAS along macropores can be expected, which originates from biological activity (biopores) or is influenced by clay accumulation. The effect of locally higher or lower CEC on soil chemical transport processes, is little investigated and still debated in literature (Jarvis, 2007). Considering a scenario in which solute transport bypasses 90 % of the bulk soil volume, as found by (Koestel and Larsbo, 2014), the location of CAS will influence the performance of mechanistic soil
- system models, especially when modeling short distance solute transport. 50

3-D X-ray imaging techniques bear great potential to study and illustrate relationships and feedback mechanisms between structural, hydrological and biochemical soil properties within undisturbed soil (Peth et al., 2014; Thieme et al., 2003; Tracy et al., 2010). Soil pore structure provides a spatial boundary that determines the accessibility of water, air and nutrients as well as separating biological processes within soils by compartmentalization (Ruamps et al., 2011; Young et al., 2008). These

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 - influences of 3-D soil structure on local processes can be studies using 3-D X-ray imaging. Larsbo et al. (2014) used 3-D Xray scanning to investigate the influence of macropore network characteristics on preferential solute transport. They found that soils with a well-connected and large macroporosity decrease preferential solute transport due to increased diffusion and exchange with the soil matrix. Naveed et al. (2016) investigated the soil structure by X-ray imaging to test prediction models

of biopore and matrix dominated water and air flows. Hapca et al. (2015) used a statistical approach to map the 3-D
distribution of several elements in undisturbed soil by combining 3-D X-ray images with 2-D SEM-EDX data. Mairhofer et al. (2016) visualized plant root interactions in undisturbed soil and Ahmed et al. (2016) visualized interactions of plant roots, phosphate fertilizers and soil structure over time in 3-D using X-ray imaging.

A common method in X-ray computed tomography is to visualize the objects or processes of interest by enhancing their X-ray photon attenuation using contrast agents. Peth et al. (2014) imaged the OM distribution in soil using osmium as a
contrast agent, which is known to bind strongly to OM. Roscoat et al. (2014) proposed a method to visualize biofilms in porous media using chloronaphtalene as a contrast agent. Koestel and Larsbo (2014) used iodide to increase the X-ray photon attenuation of water when studying water flow in an undisturbed soil column. A suitable contrast agent should contain element(s) with a higher atomic number than other common elements present in the soil. Furthermore, it must have an affinity to bind to or to dissolve in the object of interest (Van Loo et al., 2014). Such an agent increases the electron
density and the X-ray photon attenuation locally by absorbing or scattering a proportion of the primary X-ray beam

- (Wildenschild et al., 2002). Ba²⁺ is a suitable contrast agent in X-ray computed tomography of soils (Van Loo et al., 2014). It has a relatively high electron density (54) and a high affinity to replace other cations and bind to CAS. Furthermore, it is present in natural soils in rather low amounts and used in standard methods to measure the CEC. Therefore, it is a prime candidate for labeling CAS for 3-D X-ray imaging in undisturbed soil.
- 75 To our knowledge spatial variations in the CEC of undisturbed soil cores were not yet imaged in 3-D. Therefore, the objectives of this project are (i) to visualize the CAS of undisturbed soil samples in 3-D using an industrial X-ray scanner and Ba²⁺ as a contrast agent, (ii) investigate the spatial distribution of CAS with respect to the distance of soil macropores and (iii) to relate the CEC as determined by 3-D image analyses of the Ba²⁺ subjected soil sample with the CEC determined by a common laboratory analysis using ammonium acetate.

80 2. Material and methods

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2.1 Site description and sampling

Eight undisturbed, natural soil samples were taken at the end of November 2015 with aluminum columns (height: 5 cm, diameter: 2.2 cm) such that approximately half of the column volumes were filled with soil. Samples numbers (SNO) 1–4 were taken from an agricultural long term field-trial, the Swedish soil fertility experiments (Kungsängen, R3-9001; Table 1; Kirchmann, 1991). The Kungsängen site was established in 1963 near Uppsala and the soil was classified as a gleyic cambisol (Holmqvist et al., 2003). SNO1 and SNO3 were sampled from a depth of 3–5 cm and SNO2 and SNO4 from a depth of 35–38 cm corresponding to the plow pan after tillage. These samples had a high clay content (Table 1) and showed earthworm activity. SNO5–7 originated from a soil located in a marshy depression which was periodically water logged

(depth 5–8 cm). This soil is high in organic matter (OM) and classified as heavy clay (depth 5–8 cm, Table 1 and Table 2).

90 SNO8 originates from an organic matter-rich soil developed in loamy sand under a pine forest (Table 1 and Table 2). In addition, two artificial samples of fine sand containing several clay and peat aggregates were included in the experiments (SNO9 and SNO10) to visualize the difference in Ba²⁺ binding efficiency of clay and organic matter separately. The sand columns were filled under dry conditions with clay and peat aggregates as such that they were separated from each other by sand. No additional compression was applied.

95 2.2 Laboratory analyses

To evaluate the precision of the CEC_{Ba}²⁺ determined through the method described below, the CEC_{NH4}⁺ was determined by the ammonium acetate method (Thomas 1982) for comparison. The samples were sieved (< 2 mm) and air dried and the exchange sites were saturated with ammonium ions at a buffered pH of 7. The CEC_{NH4}⁺ was determined by replacement of NH₄⁺ by K⁺ and measuring the NH₄⁺ contents with a Tecator flow injection analyzer (Foss A/S, Denmark). The soil pH was determined in deionized water for all samples using a PHM 93 pH meter equipped with a Radiometer combination electrode (Radiometer A/S, Copenhagen, Denmark). The particle size distribution was analyzed by sedimentation after removal of carbonates and organic matter by using hydrochloric acid (1 mol L⁻¹) and hydrogen peroxide (30 %), respectively. The soil texture classes were determined according to FAO (2006). The total carbon contents were analyzed by the loss of ignition method according to SS- ISO 10694 for each sample location (TruMac CN, LECO Corporation, MI, USA). The bulk density was obtained by gravimetry after all analyses were completed and the samples were dried at 105 °C. The soil mass was then ralated to the sample volume obtained by the 3 D impage. The bulk densities were appreciated for the calculated Ra²⁺ mass

105 was obtained by gravimetry after all analyses were completed and the samples were dried at 105 °C. The soil mass was then related to the sample volume obtained by the 3-D images. The bulk densities were corrected for the calculated Ba²⁺ mass adsorbed by each sample.

2.3 X-ray computed tomography imaging

The GE Phoenix v|tome|x m X-ray scanner installed at the Department of Soil and Environment at the Swedish University of
Agricultural Sciences, Uppsala was used in this study. It is equipped with a 16 inch monitor (GE DRX250RT) and a 240 kV X-ray tube with a tungsten target. The samples where scanned at a maximum photon energy of 80 kV and an electron flow of 250 µA. The 3-D images where obtained by combining 2000 radiographs taken over a time of 46–90 minutes, depending on the density of the sample, thus the exposure time per radiograph was 333–1000 µs. The radiographs were reconstructed into 3-D tomograms using the GE software datos|x (version 2.1) and exported as 16 bit 3-D Tiff images with a voxel size of 20
µm.

2.4 Experimental setup

First we acquired a 3-D X-ray contrast image of the KCl $(0.1 \text{ mol } L^{-1})$ and the BaCl₂ $(0.3 \text{ mol } L^{-1})$ solutions in separate plastic vials positioned inside an empty aluminum column. This image was used to sample the gray values (Fig. 1) corresponding to the solution and the aluminum wall. These were used to relate gray values to densities and ultimately to

- 120 Ba²⁺ mass. Since the soil matrix will have an effect on the X-ray attenuation by the two solutions (see e.g. Weller et al. 2017) three additional contrast images were taken. For these images the KCl and BaCl₂ solutions were placed in an aluminum column containing air dried soil (silty clay and fine sand) that was packed to different bulk densities. The additional 3 X-ray images were used to investigate the effect of X-ray attenuation by the soil matrix on the gray value distribution within the two solutions (Fig. S1). The soil samples were first placed on sand beds in plastic cups and saturated in a desiccator under a
- 125 near-vacuum with a de-gassed KCl solution (0.1 mol L⁻¹) to avoid air entrapment inside the columns. The samples were slowly saturated from below. Residual ions in the soil columns were washed out by daily removal of the supernatant and replenishment of the KCl solution outside the column (Fig. 2). Furthermore, the soil was given time for swelling. The electrical conductivity (EC) of the supernatant was measured at regular intervals (Device: Cond 3310, WTW GmbH, Weilheim, Germany) and the treatment stopped after the EC in the supernatant had reached the EC of the KCl solution with a
- 130 deviation of max. 2.5 %. Each sample was scanned with the X-ray scanner in 3-D resolution to obtain a reference images for later processing steps. No air entrapment was found in the reference images upon visual inspection. Figure 3 illustrates the sequence of the individual steps undertaken to conduct the experiment.

All samples were then carefully transferred into new plastic cups filled with the 0.3 mol L^{-1} BaCl₂ solution. The samples were slowly saturated with BaCl₂ from the bottom up. In the following, the supernatant was removed on a daily basis and the

- 135 BaCl₂ solution outside the column was replenished (Fig. 2). In this fashion the resident KCl solution was flushed out and cations on the CAS were exchanged with Ba²⁺. During the Ba²⁺ saturation process the ECs of the supernatants were measured and X-ray images taken at regular intervals in order to find the time of Cl⁻ breakthrough and to monitor the spatial distribution of Ba²⁺ within the samples. The treatment was stopped after the EC in the supernatant of each sample had reached the EC of the BaCl₂ solution (max. tolerance of 2.5 %) and the gray values in the 3-D X-ray images showed a
- temporally stable distribution. This was the case after 25 days and removal of 160 mL cumulative supernatant, which corresponds to an average of 15 times the soil columns volume.
 To ensure that all non-adsorbed barium ions were washed out and potential BaSO₄ precipitates were redissolved, all samples

were flushed by a 0.1 mol L^{-1} KCl solution over a period of five weeks and 150 mL KCl solution per sample. After the EC of the supernatant had stabilized at the EC of the KCl solution (tolerance maximally 2.5 %), the KCl rinsing process was

stopped and the final 3-D images where taken of all samples. When the final images were taken the majority of the CAS were assumed to be occupied by Ba^{2+} and that the K⁺ competition was of minor importance. In the following we refer to these images as " Ba^{2+} subjected" images. The average difference in the gray values of the soil solutions in the macropores

between the reference images and the Ba^{2+} subjected images were within a range of 87 gray values thus, the soil solutions were assumed to have the same densities.

150 Three samples (SNO4, SNO6 and SNO10) were excluded from further analyses. SNO6 and SNO10 had been destroyed after the saturation process in the desiccator by a too rapid rise in air pressure. SNO4 had a very small hydraulic conductivity and all required experiments could not be completed.

2.5 Image processing

- The software ImageJ/FIJI (Schindelin et al., 2015) with the plugin SoilJ (Koestel, 2017) was used for image processing. The resolution of all 3-D images was reduced by a factor of 4 in order to decrease the image-processing time for the subsequent steps. Thus, the analyzed images had a resolution of 80 μm. The first step of the image analysis was to correct the image illumination to ensure that all objects of the same density within individual images and across all images exhibited the same gray values. The mean gray values obtained for the aluminum wall and the KCl solution in the contrast image (Fig. 1) were used as the target gray values for the gray-scale standardization (21,418 and 16,225 respectively). All other gray values were
- 160 scaled accordingly by the linear relationship between the target gray values and the initial gray values of the corresponding image according to Koestel and Larsbo (2014). A 3-D unsharp mask with one pixel radius was applied to all images in order to increase sharpness. Both, the reference image and the Ba²⁺ subjected image were binarized using a threshold gray value of 16,990, which was obtained from the joint histogram of all images (Fig. 4). The binerized images were created to obtain the pore space for both the reference image and the corresponding Ba²⁺ subjected image.

165 **2.6 Creating the difference images**

The images where registered using the ImageJ plugin "descriptor-based series registration (2-D/3-D)" by Preibisch et al. (2010) with the transformation model affine to account for minor changes in particle positions during the sample treatment. Thereafter, the difference images were obtained by subtracting the 3-D Ba²⁺ subjected images from the 3-D reference images.

170 2.7 Relationship between gray values and barium mass

To estimate the Ba^{2+} mass in the difference images as a proxy for the CEC_{Ba}^{2+} the 3-D contrast image of $BaCl_2$, KCl solutions and the aluminum wall was used as a reference (Fig. 1). The mean gray values of the two solutions were subtracted to obtain the maximal contrast in gray values ($\gamma_{SAT} = 2637$) corresponding to the density contrast between the KCl and $BaCl_2$ solutions. The $BaCl_2$ mass ($m_{j,d}$ in mg) was then calculated according to Koestel and Larsbo (2014) using the Eq. (1).

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$$m_{j,d} = \frac{\left(N_{Ba} - N_{K}\right)}{N_{Ba}} \frac{V_{VOX} C_{MAX}}{\gamma_{SAT}} \gamma_{j,d} \qquad \text{Eq. 1}$$

where *j* is the voxel, $N_{Ba} = 56$ and $N_K = 18$ are the atomic numbers of barium and potassium, respectively, V_{VOX} represents the voxels' volume (5.12 * 10⁻⁷ cm³), C_{MAX} the maximal possible increase in tracer solution (41.199 mg cm⁻³) and γ the corresponding gray value. In Eq. 1 we assume that the CAS were predominantly occupied by K⁺ when the reference images were taken and with Ba²⁺ when the Ba²⁺ subjected images were taken. In order to calculate an estimate of the CEC_{Ba}²⁺ in cmol (+) kg⁻¹ soil the sum of the positive charged sites as occupied by Ba²⁺ was calculated and related to the samples volume

180 cmol (+) kg⁻¹ soil the sum of the positive charged sites as occupied by Ba²⁺ was calculated and related to t and bulk density.

2.8 Spatial distribution of cation exchange capacity

To test whether the imaged Ba²⁺ concentrations as proxies for the CECs were elevated in macropore sheaths (400 µm distance from pore surface) as compared to the Ba²⁺ concentrations in the soil matrix, the binarized 3-D images of the pore space for the reference and the Ba²⁺ subjected images were combined using the ImageJs' Image Calculator. This was done to account for any changes in pore space due to disturbances during the saturation periods. The resulting images of the combined pore space were dilated five times using the ImageJ plugin Process and its function dilate 3-D. Subsequently, this dilated binary image was subtracted from the corresponding difference image in a way that the resulting image showed the gray values from the difference image only outside the pore space and the dilation area (only the soil matrix). After inverting the dilated binary image it was subtracted from the difference image and resulting in a second image that represent only the

gray values in the dilation area, thus the soil around the macropores. These images were used to assess the difference of the imaged density of CAS between the soil close to macropores and the soil in the matrix of all natural soil samples.

2.9 Data analysis

For statistical analyses the open source software R (v0.98.1x) and RStudio (v3.2.5) was used (R Core Team, 2016). The
 relation between measured CEC_{NH4}⁺ and the CEC_{Ba}²⁺ obtained through image analyses was analyzed by a linear model. The CEC_{Ba}²⁺ distribution around the macropores of the natural soil cores was graphically compared. For 3-D visualization the open-source software drishti was used (Limaye, 2012).

3 Results and Discussion

3.1 Uncertainties and image artifacts

200 An increase in contrast between the KCl and BaCl₂ solutions was observed with increasing density of the material used for the three additional contrast images. The average increase relative to the contrast image of the air-filled column did not exceed 10 % (Tab. S1 and Fig. S2 and S3). Considering that these packed soils exceeded the bulk densities of the samples described in our manuscript this increase in contrast is the maximum expectable increase. As we were using the air-filled contrast image in the following calculations, our results correspond to the lower bond of possible Ba²⁺ mass estimations. 205 Therefore, it would mean that our estimates of the cation exchange capacity (CEC) are underestimated by a maximum of 10 %.

Image artifacts in the difference images can originate from soil movement after the reference image and before the Ba²⁺ subjected image were taken. These will be visible as bright areas, if high gray values in the Ba²⁺ subjected image (e.g. soil matrix) are subtracted from low gray values in the reference image (e.g. soil pore) and in dark areas, if the reverse is the

- 210 case. The areas in the magnified difference images of SNO2 and SNO3 (marked with an X, a circle and arrows in Fig. 5) are due to such shifts. For SNO2 a shift only occurred locally, but for SNO3 shifts were visible throughout the soil column. In Fig S4 in the supplementary material we visualized the global gray value distribution of the soil columns of SNO1, 2 and 3. A difference between SNO1 with very few artifacts and SNO2 and 3 with more abundant artifacts due to particle shifts is apparent. The histograms of SNO2 and SNO3 have two plateaus in gray value abundance at approximately gray values +/-
- 215 1500 to +/-4500 and gray values +/-1500 to +/-3750 respectively (red horizontal lines in Fig. S1). Whereas, the histogram of SNO1 does not show a plateau feature. These plateaus represent local particle shifts within the samples that occurred after the reference images were taken. As a particle shift will lead to large positive gray values at one side of the image feature and large negative values at its opposite side. In future studies, measurements of the size of plateaus in difference images may be investigated as means to quantify the amount of misplaced regions.
- 220 In Fig S4 the small peak around gray value -1300 (black vertical line), corresponds to air bubbles that had formed after the reference images were taken.

Swelling and shrinking of the soil in between the two scanning occasions could lead to artifacts similar to the described above. In this case the most obvious artifacts would present themselves as brighter areas around macropores if the sample was swelling or darker areas if it was shrinking. However, the occurrence of artifacts due to swelling or shrinking was ruled

225 out after scrutinizing the size and shape of respective macropores and it was found that none of them had changed (see Fig. 6 as an example).

In the difference image of SNO1 we found some brighter spots within the soil matrix that correspond to high density areas in the reference image (Fig. 6, circles). We hypothesize that these are porous iron or manganese oxides concretions that either bind Ba^{2+} and therefore are visible as bright spots in the difference image, or represent imaging artifacts resulting from the

230 grayscale standardization. Since the applied grayscale standardization is only valid for imaged densities in between the density of the KCl solution and the density of the aluminum wall. In case the relationship between image gray-value and material density is non-linear or changing with the introduction of a denser material (here Ba²⁺), the resulting difference images may not represent the Ba²⁺ densities only. This would especially be the case when estimating imaged densities larger than the aluminum wall (e.g. iron or manganese oxides).

235 **3.2** Spatial heterogeneity of cation adsorption sites

Figure 7 shows cuboids extracted from the center of the reference images and the Ba²⁺ subjected images from regions of interest with a quadratic horizontal cross-section (edge-length 14.4 mm) of SNO1 (height 17.76 mm), SNO3 (height 14.8 mm), SNO7 (height 20.32 mm) and SNO8 (height 20 mm). The Ba²⁺ distribution is visualized by translucent areas (no or little Ba^{2+} , green (medium Ba^{2+} mass) and blue (high Ba^{2+} mass). Note that the color-scales in Fig. 7 are optimized for depicting the 3-D structures and therefore semi-quantitative. The samples from the agricultural silty clay soil (SNO1 and 240 SNO3) show a relatively uniform distribution of adsorbed Ba^{2+} with some spots of high values within the matrix and some elevated values around macropores in SNO1. The heavy clay soil (SNO7) shows a high spatial heterogeneity in Ba²⁺ densities with large areas of high values and large areas of very low values. The loamy sand soil (SON8) shows plenty of areas with no or little Ba^{2+} adsorbed and some with locally distinct higher values. Figure 8 shows cross sections of the reference images for the seven intact soil samples and Fig. 9 shows the corresponding difference images, here the gray value 245 scale is quantitative. In Fig. 9 it is easy to differentiate between pores and soil matrix. Thus, most of the soils possess CAS abundant enough to be visualized by this method. The adsorbed Ba²⁺ on CAS and its horizontal variances in the cross sections were particularly large in the heavy clay soils SNO5 and SNO7. In contrast, the other four undisturbed samples (SNO1, SNO2, SNO3 and SNO8) have lower contrasts and less Ba^{2+} adsorbed. This pronounced difference between the samples reflects the variation in CEC_{NH4}^+ and OM contents between them (Table 1 and Table 2). The artificial sample 250 (SNO9) shows the highest gray values for OM aggregates, which is easy to differentiate from the clay aggregates and the sand. The difference in gray values of the clay aggregates and the surrounding sand is less distinct than the gray value difference of the OM aggregate and the sand. This reflects the variation of the respective CEC_{NH4}^+ (Table 2). In Fig. 10 the arithmetic means and standard deviations of the gray values in each horizontal cross section is plotted against depth for all soil samples. This confirms the observations made in Fig. 9 that SNO5 and SNO7 show the highest density of Ba²⁺ adsorbed 255 to CAS and SNO7 the highest heterogeneity in a vertical profile. All the soils from the agricultural field-trial and the forest soil show lower Ba²⁺ mass adsorbed and little vertical heterogeneity.

3.3 Cation adsorption sites and macropore space

- The sample SNO1 shows brighter gray values around biopores. A magnification of these brighter areas shows that they surround the entire pore walls (with an approximate thickness of 0.4 mm, Fig. 6). This is likely due to a locally higher CEC_{Ba}²⁺ caused by accumulation of organic matter. Figure 6 shows a biopore in SNO1 that was created by an earthworm, judging on its shape and size. The Ba²⁺ mass distribution calculated from the extracted gray values of the 3-D matrix space and the 3-D space in the macropore sheaths of the difference images are presented in Fig. 11. For SNO3, SNO7 and SNO8 the median Ba²⁺ masses within the matrix are lower than those within the sheaths. SNO2 and SNO5 show the opposite trend and no difference between the Ba²⁺ masses of matrix and sheath is observed for SNO1. Hence, the observed increased Ba²⁺ mass around the biopores depicted in Fig. 6 (middle) was specific to this macropore but was not observed for other
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macropores in this sample. In general, there is little difference between the Ba^{2+} mass distributions of the individual samples from the agricultural field-trial. This may be explained by the fact that the samples were taken from the plow layer in autumn after the field had been plowed and biopores were very rare or freshly formed and not yet coated with sufficient amounts of

- 270 OM. With the exception of SNO5 the soils from the unplowed sites (marshy depression and forest soil) show higher Ba²⁺ masses adsorbed within the macropore sheaths compared to the matrix. SNO7 originate from a heavy clay soil, where biopores formed by faunal activity or root growth are not destroyed by plowing and therefore, the OM within them is redistributed to a lesser extent. SNO8 was sampled in a loamy sand soil, here the difference between the Ba²⁺ mass distributions is likely due to the fact that the sand grains are surrounded by OM and fine material and therefore, more likely
- 275 located within the matrix. In contrast, the macropore sheaths tend to contain higher amounts of OM and fine material. Sand grains typically exhibit a low CEC and seen as darker objects in the difference image compared to OM and fine material that usually possess a higher CEC and are able to adsorb more Ba²⁺. This can be seen in Fig. 11, where the median Ba²⁺ mass of the matrix is below zero and its third quartile is exceeded by the median Ba²⁺ mass of the macropore sheaths. Ba²⁺ mass values below zero can be explained by a very low Ba²⁺ mass in general and possibly in combination with small shifts in soil

structure. In addition, the Ba²⁺ contrast will be underestimated if the KCl treatment did not lead to a complete exchange of

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cations that are heavier than K^+ and were exchanged by Ba^{2+} later.

3.4 Comparison with ammonium acetate method

The correlation of the CEC_{NH4}^{+} measured in the laboratory with the CEC_{Ha}^{2+} obtained from the difference images shows a significant relationship ($R^2 = 0.87$; p <0.01; Fig. 12). It confirms that the applied method captured the trend in CEC across the soil samples and strengthens the validity of our results obtained by the difference image analysis. CEC_{R0}^{2+} underestimated 285 the low CEC levels by the silty clay soils from the agricultural field and the loamy sand soil under forest (SNO1, SNO2, SNO3 and SNO8) and overestimated the heavy clay soil that is rich in OM (SNO5 and SNO7). An underestimation of the CEC by our difference image analysis can result from multiple factors. The laboratory analysis of CEC_{NH4}^+ is based on experiments with sieved, finely grinded soils, whereas our results of CEC_{Ba}^{2+} are based on undisturbed soils. Some soil surfaces in pores not accessible to the BaCl₂ solution in the undisturbed samples may not have been saturated with Ba²⁺ and 290 did not contribute to the CEC_{Ba}^{2+} . When investigated as sieved soils these surfaces get exposed and more easily accessible, thus participating in ion exchange. Our method may estimate the physically accessible CAS more accurate than the CEC_{NH4}⁺ method and therefore underestimates the CEC of the dense clay samples compared to the CEC_{NH4}^+ method. Using the two different index cations (Ba²⁺ and NH₄⁺) to estimate the CEC can also lead to different results (Ciesielski et al., 1997; Jaremko 295 and Kalembasa, 2014). In the presence of 2:1 clay minerals like smectites, NH_4^+ may be trapped in the interfoliaceous cavities (Essington, 2004a; Pansu and Gauthevrou, 2006). Furthermore, competition and displacement of Ba^{2+} on clay

mineral surface adsorption sites by K^+ can be rather strong due to their similar ionic radii (Kabata-Pendias, 2010). Therefore, the KCl rinsing process may have (i) reduced Ba²⁺ densities locally as well as the X-ray photon attenuation on clay mineral surfaces and (ii) underestimated CEC_{Ba}^{2+} of clay-rich samples in contrast to organic matter rich samples where Ba^{2+} can be bound in OM complexes (Bodek et al., 1988; Bradl, 2004; Lee et al., 2007; Pichtel et al., 2000). The low CEC_{Ba}^{2+} values of SNO8 seem to contradict this hypothesis. However, SNO8 originated from a loamy sand soil under a pine forest, and its soil OM appeared little humified, therefore the density of functional groups and the potential to develop a larger CEC may not have been fully reached (Essington, 2004b). Additionally, the low pH of this sample may have led to an underestimation of the CEC_{Ba}^{2+} compared to the buffered CEC_{NH4}^{++} (Skinner et al., 2001).

305 4 Conclusions

We have shown that a modern industrial X-ray scanner is capable of providing 3-D images that can be used to map the cation adsorption sites (CAS) in undisturbed soil cores by difference image analyses. Furthermore, Ba²⁺ provides enough contrast to assess the 3-D distribution of CAS in soil. All undisturbed soil samples showed some degree of a spatially heterogeneous distribution of CAS most of them down to a millimeter scale. However, no clear relationship between the location of CAS and the macropore structure was found. Even though our method deviated from the common ammonium acetate method to some extent, the results correlated significantly. This deviation may be due to several factors. The most likely are a lower accessibility of CAS in the undisturbed soils used here compared to the ammonium acetate method that is based on sieved soils. A competition between K⁺ and Ba²⁺ on the ion exchange sites on clay surfaces that may have led to a reduction in local

 Ba^{2+} densities in the clay rich samples. To further assess the discrepancy between these two methods a larger set of soil

- 315 columns could provide detailed information on systematic deviations due to physical or chemical soil properties. It could also provide us with further valuable information on the accessibility of CAS and aid the development of more realistic soil system model. Our here proposed method could also be used to map organic matter distribution in 3D. For this purpose the KCl rinsing process should be prolonged and the KCl concentration could be increased. This would make it more likely that most of the Ba²⁺ bound to clay surfaces and other exchange sites is replaced by K⁺, whereas the Ba²⁺ bound in complexes to organic matter would stay in place. Alternatively heavy anions could be used as contrast agents for imaging the soil organic
- 320 organic matter would stay in place. Alternatively heavy anions could be used as contrast agents for imaging the soil organic matter instead of Ba^{2+} (e.g. I⁻, Br^- , WO_4^{2-} or MoO_4^{2-}). When used on soils from temperate climate regions these may have the advantage that the CEC is not biasing the results.

The difference image resolution could be improved by reducing the sample size. The smaller the sample the better the resolution. Note that the maximum resolution also depends on the hardware used (X-ray scanner and computer) and its

325 configuration. After some preliminary tests we found that the scanner used in this study (GE Phoenix v|tome|x m) is capable of taking images at a resolution down to 5 μm at a soil column with a diameter of 8 mm. Others have reported resolution down to 1 μm when using X-ray scanner optimized for smaller sample sizes (e.g. Tippkötter et al., 2009). By using a monochromatic X-ray source Voltolini et al. (2017) imaged soil micro-aggregates with a sub-micron resolution.

The difference image quality could be improved by using the absorption edge technique and a monochromatic X-ray beam 330 as is available on synchrotron facilities. This would eliminate the negative effects on the difference image quality by soil aggregate displacement or a possible change in the relationship between the gray values and object densities, although it would reduce the sample size.

5 Competing interests

The authors declare that they have no conflict of interest.

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340 Table 1. Location of each sampling site, Organic carbon (Org. C), clay, silt and sand contents.

Site	Location	Org. C [%]	Clay [%]	Silt [%]	Sand [%]
Kungsängen (3–5 cm	59.836361, 17.6875	2.17	55.5	41	3.6
depth)					
Kungsängen (35–38 cm	59.836361, 17.6875	2.34	53.9	42.4	3.7
depth)					
Periodically water logged	59.823972, 17.6623	4.78	79.9	20	0.3
soil (5–8 cm depth)					
Forest soil (5–8 cm	59.824194, 17.6633	4.76	8.9	10.7	80.3
depth)					

Table 2. Soil texture classes, sampling site, pH, cation exchange capacity (CEC_{NH4}^+) and bulk density (BD) for each sample number345(SNO).

SNO	Soil texture	Site	рН [H2O]	CEC [cmol kg ⁻¹]	BD [g cm ⁻³]
1	Silty clay	Kungsängen	6.5	20.3	1.14
2	Silty clay	Kungsängen	6.5	19.1	1.30
3	Silty clay	Kungsängen	6.3	19.9	0.87
4	Silty clay	Kungsängen	6.2	20.4	1.20
5	Heavy clay	Periodically water logged soil	7.1	47.9	0.48
6	Heavy clay	Periodically water logged soil	6.8	22.0	0.41
7	Heavy clay	Periodically water logged soil	6.8	22.0	0.56
8	Loamy sand	Forest soil	5.1	13.1	0.96
9 & 10	Fine sand		4.9	3.0	
9 & 10	Clay aggregates		7.0	16.9	
9 & 10	Peat aggregates		6.7	45.7	

Figure captions



350 Figure 1: Contrast image of the aluminum cylinder wall, KCl and BaCl₂ solution and the air (left). The corresponding histogram on the right with the mean values for the aluminum cylinder wall (green), KCl (red) and BaCl₂ (blue) solution and the air (yellow) indicated as dotted lines.



Figure 2: Set-up for the KCl or BaCl2 saturation process.



Figure 3: Experimental time schedule. X-ray scans were performed after KCl treatment to obtain the reference image (Scan 360
1) and after removal of residual Ba²⁺ ions to obtain the Ba²⁺ subjected image (Scan 2).



Figure 4: The joint histogram of the reference and Ba^{2+} subjected images. The red dotted line represents the segmentation threshold that was used for binarization.



Figure 5: Effect of aggregates movement on the difference image of SNO2 (top) and SNO3 (bottom). Reference image (left), difference image (middle) and the image of Ba²⁺ treated soil (right). The red cross and circle indicate the identical coordinates in all three images. The movement of one soil aggregate resulted in very high gray values (red marks) or very low gray values (blue marks) in the difference image. Note that the reference images and the image of Ba²⁺ treated soil share the same gray value calibration bar.



Figure 6: Magnification of a macropore from SNO1. Reference image (left), difference image (middle) and the image of Ba2+ treated soil (right) (right). The X and the circle indicate the identical coordinates in the images. Note that the reference image and image of Ba2+ treated soil share the same gray value calibration bar.





only semi-quantitative.



Figure 8: Cross sections of the reference images of the seven intact soil samples. Depth from the soil surface: SNO1, SNO2 and SNO3 at SNO8 mm, SNO7 and 8 at 9 mm, SNO5 at 15 mm and SNO9 at 5 mm.



Figure 9: Difference images of the seven intact soil samples. The gray scale represents the Ba^{2+} mass (M_{Ba}) in µg per voxel (vx). Depth from the soil surface: SNO1, SNO2 and SNO3 at SNO8 mm, SNO7 and SNO8 at 9 mm, SNO5 at 15 mm and SNO9 at 5 mm.



Figure 10: Vertical profiles of the Ba²⁺ mass distributions of the seven natural soil samples. Arithmetic mean (left) and standard deviation (right).



Figure 11: Comparison of the Ba^{2+} mass distributions within the matrix (red) and within the macropore sheaths (blue) for all natural soil samples.



Figure 12: Relation between the CEC (cmol(+) kg⁻¹) measured with NH₄⁺ and the CEC_{Ba}²⁺ obtained from the difference image analysis of the natural soil samples, blue line represents a linear model with its 95 % confidence interval in gray (p < 0.01). The dotted line has a slope of 1.