The Role of Inorganics in Preeclampsia Assessed by Multiscale Multimodal Characterization of Placentae

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Preeclampsia is one of the most dangerous diseases in pregnancy. Because of the hypertensive nature of preeclampsia, placental calcifications are believed to be a predictor for its occurrence, analogous to their role in cardiovascular diseases. However, the prevalence and the relevance of calcifications for the clinical outcome with respect to preeclampsia remains controversial. In addition, the role of other inorganic components present in the placental tissue in the development of preeclampsia has rarely been investigated. In this work, we therefore characterized inorganic constituents in placental tissue in groups of both normotensive and preeclamptic patients (N = 20 each) using a multi-scale and multi-modal approach. Examinations included elemental analysis (metallomics), sonography, computed tomography (CT), histology, scanning electron microscopy, X-ray fluorescence and energy dispersive X-ray spectroscopy. Our data show that tissue contents of several heavy metals (Al, Cd, Ni, Co, Mn, Pb, and As) were elevated whereas the Rb content was decreased in preeclamptic compared to normotensive placentae. However, the median mineral content (Ca, P, Mg, Na, K) was remarkably comparable between the two groups and CT showed lower calcified volumes and fewer crystalline deposits in preeclamptic placentae. Electron microscopy investigations revealed four distinct types of calcifications, all predominantly composed of calcium, phosphorus and oxygen with variable contents of magnesium in tissues of both maternal and fetal origin in both preeclamptic and normotensive placentae. In conclusion our study suggests that heavy metals, combined with other factors, can be associated with the development of preeclampsia, however, with no obvious correlation between calcifications and preeclampsia.

Keywords: calcifications, hypertension, electron microscopy, preeclampsia, elemental analysis (chemical), histology
INTRODUCTION

Preeclampsia (PE) is a severe pregnancy-associated disease and a major cause of morbidity and mortality affecting 3–8% of all pregnancies worldwide, accounting for over 50,000 deaths of mothers and their offspring annually (1). PE is characterized by hypertension (with systolic blood pressure $\geq$ 140 mmHg and diastolic blood pressure $\geq$ 90 mmHg) accompanied by proteinuria ($\geq$ 300 mg/24 h) (2, 3). In clinics, a distinction is made between early onset and late-onset PE, with 34 weeks gestation being the cut-off value (4). An early sign of early onset PE is the abnormal placentation of cytotrophoblasts in the first trimester. In healthy pregnancies, the maternal spiral arterioles are re-modeled over time to allow a higher blood flow, accounting for the increasing nutrient demand (5). In early onset PE the spiral vessels do not complete the transformation from a proliferative epithelial to an invasive endothelial subtype, causing the diameters of the arteries to be narrower, subsequently leading to relative placental ischemia (6). Late onset PE is mainly induced through limited intervillous perfusion leading to chronic hypoxic placental tissue (7). Both early onset PE and late onset PE are associated with increased levels of proinflammatory and antiangiogenic factors in the maternal blood, including increased concentration of circulating soluble fms-like tyrosine kinase 1 (sFlt-1), which exacerbate the clinical symptoms of the mother (8).

The pathophysiology of PE and its manifestations in the placenta remain poorly understood. Research on PE has focused primarily on histoanatomical (9) and biochemical (5, 10) approaches, however, several studies indicate the potential importance of inorganics in the development of the disease, including macro and micro-nutrients, as well as (toxic) trace elements. For example, elevated concentrations of Cr, As, Pb, Ni, and Hg may contribute to the development of a proinflammatory and antiangiogenic environment by induction of oxidative stress, hence contributing to the development of PE (11–13). Most prior works have examined element concentrations of preeclamptic patients in the blood or urine. Studies analyzing preeclamptic placental tissue for elemental composition (major and/or trace elements) and minerals/calcifications, however, are relatively scarce (14, 15).

In addition to toxic trace elements, mineral deposits such as calcifications may play a key role in PE. Calcifications can be formed by metastatic, dystrophic or physiologic processes. Metastatic calcification is defined as the deposition of calcium in healthy tissue as a result of supersaturated calcium serum concentrations (16). Dystrophic calcification occurs in necrotic tissue while physiologic calcification is seen in bone and teeth formation (17, 18). The extent and tissue-specific patterns of vascular calcification are well-known predictors of cardiovascular morbidity and mortality. Indeed, placental calcifications similar to those found in aortic tissue have been reported recently (19). Despite the fact that PE is a hypertensive disease, placental calcifications are scarcely researched in this context. While calcifications are often observed in both healthy and diseased placentae, their relevance for clinical outcome remains controversial, in particular in PE (20–22). In a large study assessing placental histological sections of preeclamptic patients, Ezejigwe et al. found a correlation between placental infarcts and calcifications, however, failed to show a significant relationship between increased placental calcification and preeclampsia (23). In contrast, Grannum III calcifications were suggested to be a useful indicator of PE development later in the pregnancy (24). Furthermore, Chen et al. indicated that the presence of placental calcifications and weight gain during pregnancy might serve as a warning sign and requires closer surveillance for maternal and fetal well-being (25). Note, however, that most of the placental calcification research to date has focused on clinical sonography imaging and histological tissue analysis with poor sensitivity for mineral characteristics. A recent review by Wallingford et al. has stated that investigation is needed to delineate associations between preeclampsia, placental calcification, and vascular calcification in order to evaluate the potential diagnostic value of placental calcification in both acute and long-term cardiovascular health (26).

In this work, we therefore assess inorganic constituents in healthy and preeclamptic placentae with varying degrees of severity by conventional clinicopathological methods enriched by metallomics analysis and cutting-edge nano-analytical characterization. We present a route to augment the detection capabilities of histology by correlation with whole slide micro-X-ray fluorescence ($\mu$XRF) and backscattered electron (BSE) imaging in the scanning electron microscope (SEM), allowing the unambiguous identification and (chemical) characterization of placental solid inorganics (incl. calcifications) with nanometric resolution.

MATERIALS AND METHODS

Patient Cohort and Placentae Harvesting

The study was approved by the Ethics Commission of Eastern Switzerland (EKOS 2020-01387). All placentae were collected directly after delivery at the Cantonal Hospital of St. Gallen, Switzerland, after obtaining informed consent from the patients. Placentae ranging from 25 + 6 weeks of pregnancy to 40 + 5 weeks of pregnancy were included. The corresponding control group included 20 term births between weeks 37 + 0 and 41 + 2 of gestation by healthy, normotensive women.

Ultrasound Examination of the Placentae

Immediately after collection, the degree of placental calcification was assessed by an $ex vivo$ ultrasound scan. The evaluation was carried out in analogy to the Grannum score which distinguishes between a smooth chronic plate (Grade 0), occasional parenchymal calcification (Grade I), deeper indentations of the chronic plate (Grade II) and significant basal plate calcification with indentations interrupting the chorionic plate (Grade III) (27). All ultrasound examinations were performed by a qualified obstetrician using a Voluson 730 (GE Medical Systems, Zipf, Austria) equipped with a transabdominal transducer operating in the frequency range between 2.8 and 10 MHz.
Computed Tomography

Placenta were imaged within 1 h after delivery at the Department of Radiology Cantonal Hospital St. Gallen using a dual-source CT system (Somatom Force, Siemens Healthineers, Forchheim, Germany). Images were collected in single source mode using the following scanning parameters: tube voltage: 120 kilovoltage peak (kVp, peak potential applied to the x-ray tube, which accelerates electrons from the cathode to the anode); tube current-time product: 380 mAs; collimation: 192 × 0.6 mm; gantry rotation time: 250 ms; and pitch: 0.9. Tomography data were reconstructed using a slice thickness of 0.4 mm and an increment of 0.2 mm with a sharp convolution kernel (Ur77). Image data were reconstructed on a dedicated radiological workstation and evaluated by an experienced radiologist using commercially available CT analysis software (Extended Multi-planar Reconstruction plugin, impax ee, AGFA HealthCare, Mortsel, Belgium). In each CT dataset a threshold-based segmentation algorithm was used to quantify (i) the volume of the placenta by counting the volume of all voxels with a CT density between −150 and 150 Hounsfield units (HU), and (ii) the volume of calcified deposits by counting the volume of all voxels with a CT density of 180 HU or higher. The Hounsfield unit (sometimes also referred to as the “CT-unit”) is a relative measurement of the attenuation of X-rays by the specimen in computed tomography (CT) in relation to water and air, where water is arbitrarily set to 0 HU and air to −1,000 HU at standard temperature and pressure (28). In CT images, the Hounsfield scale is then displayed as gray tones.

Elemental Analysis (Metalломics)

After ultrasound and CT imaging, five full thickness punch biopsies (8 mm diameter) were taken from each placenta in a circular pattern and stored in 70% ethanol. The total placental contents of major (Ca, P, Mg, Fe, K, and Na) and trace elements (Al, Cr, Mn, Co, Ni, Cu, Zn, As, Se, Rb, Sr, Ag, Cd, Cs, Ba, and Pb) were determined. Trace elements were chosen according to previous reports with regard to PE (e.g., Cr, As, and Cd (29), Pb) were then determined. Trace elements were digested in 2.5 mL HNO₃ (Extended Multi-planar Reconstruction plugin, impax ee, AGFA HealthCare, Mortsel, Belgium). In each CT dataset a threshold-based segmentation algorithm was used to quantify (i) the volume of the placenta by counting the volume of all voxels with a CT density between −150 and 150 Hounsfield units (HU), and (ii) the volume of calcified deposits by counting the volume of all voxels with a CT density of 180 HU or higher. The Hounsfield unit (sometimes also referred to as the “CT-unit”) is a relative measurement of the attenuation of X-rays by the specimen in computed tomography (CT) in relation to water and air, where water is arbitrarily set to 0 HU and air to −1,000 HU at standard temperature and pressure (28). In CT images, the Hounsfield scale is then displayed as gray tones.

Histology

Tissue processing, preparation of histological sections and staining were performed using standard methods at the Institute of Pathology Cantonal Hospital of St. Gallen, Switzerland. The pathological examination included a macroscopic description of the location (central, paracentral and peripheral) of calcifications and targeted embedding followed by histological examination based hematoxylin eosin (HE) staining. The placenta were fixed with 4% buffered formalin and the relevant sampled areas were embedded in paraffin blocks. The blocks were cut into 2 µm sections, stained with HE and evaluated for pathological parameters using a conventional light microscope. In HE, calcified areas appear deep blue/purple.

In addition, Von Kossa staining was performed (35). To this end, unstained histological sections were deparaffinized by incubation in xylene for 10 min (2x) followed by rehydration in a decreasing ethanol gradient. After deparaffinization, the samples were incubated with 5% silver nitrate (abcam® ab150687) for 45 min. Throughout the incubation time, the samples were exposed to a 60W incandescent light. Then, samples were rinsed with deionized water and incubated with 5% sodium thiosulfate for 3 min (abcam® ab 150687). The samples were also counterstained with Nuclear Fast Red staining (abcam® ab 150687) for tissue visualization. Finally, a coverslip was placed over the tissue using DPX mounting medium (Sigma-Aldrich 06522). After von Kossa staining, calcified areas appear black due to the reduced (metallic) silver.

Micro-X-Ray Fluorescence Analysis (μXRF)

Unstained, deparaffinized histological sections were mounted on high purity vitreous carbon disks (Micro-to-nano, Netherlands) in order to allow for elemental analysis with virtually only C as a background element. Element distributions maps of the deparaffinized histological thin sections mounted on high purity vitreous carbon disks were recorded on a M4 Tornado micro X-ray fluorescence (μXRF) spectrometer (Bruker Nano GmbH, Berlin, Germany) equipped with an Ag cathode operated at an acceleration voltage of 50 kV. The spot size of the X-ray beam was 20 µm. Each map was scanned with a dwell time of 5 ms per pixel, with a total collection time of 6 h per map. Data were evaluated using Bruker software.
**Scanning Electron Microscopy**

For SEM analysis, the carbon disks were then mounted on conventional aluminum stubs and coated with 10 nm carbon using a CCU-010 coater (Safematic, Switzerland). SEM imaging (secondary and backscattered electrons) and EDX analysis were performed on a Quanta 650 SEM (FEI, Thermo Fisher Scientific) at 10 kV. EDX Data was evaluated using Pathfinder 2.4 (Thermo Fisher Scientific). Large field of view stitch images of entire thin sections (secondary and backscattered electrons) were created on a Magellan 400 SEM (FEI, Thermo Fisher Scientific).

**RESULTS**

**Cohort Description**

The entire study population was of Caucasian ethnicity (Table 1), with the majority of patients being over 30 years old. The prevalence of PE was higher in older patients, indicative of the increased likelihood of PE at higher age. All women had a singleton pregnancy and were mostly pregnant for the first time (N = 23). Most patients, namely 85% in the control and 80% in the PE group, were pre-obese (BMI 25–29.9) or obese (BMI > 30) at the time of delivery. In the PE-group, two patients admitted to smoke regularly during pregnancy.

Based on the Grannum score, which is commonly used to assess placental calcifications during gestation, heavy calcifications (Grannum 3) were only found in two patients of the control group. The level of mild calcifications was very comparable between the groups. Thus, in this case the Grannum score was not indicative of PE development or the severity of PE. Not unexpectedly, the placenta weights were significantly lower in the PE compared to the normotensive group due to the preterm deliveries in the PE group (see Table 1).

**Metallomics Analysis of Bulk Tissue**

In order to gain insights into the role of inorganics in healthy and preeclamptic placentae, bulk multi-element analysis was performed on placental tissue. Elemental concentrations in blood and urine may fluctuate during pregnancy due to physiological and metabolic changes (36). Thus, for a more robust survey, major elements (Figure 1) and trace elements (Figure 2) were analyzed directly in the placental tissue after postpartum by elemental analysis using ICP-MS (Supplementary Tables 1, 2).

In general, the median concentration levels of the major elements Mg, P, and Ca in placental tissue were very comparable to results reported in the literature (19) (Figure 1). No significant difference in Na, Mg, P, K, and Ca content between the control and PE group were found on a bulk level (Figures 1A–F). Only for Fe, significantly elevated levels were observed in the PE compared to the normotensive group (median values of 847 (PE) and 758 (control), Supplementary Table 2, ANOVA: p = 0.044).

Especially for Ca, the inter-subject variance was remarkably high with, however, comparable ranges in both groups. Calcium concentration ranged from 764 up to 21,597 mg/kg in the control and from 742 up to 19,496 mg/kg in the PE group (Supplementary Table 1), which corresponds to a 28 and 26 times higher maximum value compared to the minimum, respectively.

With regard to trace elements and heavy metals, levels of Al, Cr, Mn, Co, Ni, Cu, Zn, As, Se, Pb, Sr, Ag, Cd, Cs, Ba, and Pb were assessed in the placental tissues (Figure 2). One factor ANOVA indicates highly significant elevations in Al (p < 0.0001), Cd (p < 0.0002), Ni (p < 0.0001), Co (p < 0.0001) and Mn (p = 0.0003) as well as moderately higher Pb (p = 0.046) and As (p = 0.084) levels in the PE group on a bulk level. In contrast to the aforementioned elements, Rb showed a significantly lower concentration in the PE group (p = 0.034).

**Clinicopathological Analysis Enriched With Nano-Analytical Techniques**

**Computed Tomography**

To assess macroscopic calcifications, all placentae were analyzed by CT. CT analysis allowed quantification of the macroscopic total calcified volume and extraction of information on the crystallinity/density of the calcifications based on the Hounsfield

**TABLE 1 | Characteristics of the study population.**

| Characteristic                      | Control group (n = 20) | Preeclampsia group (n = 20) |
|-------------------------------------|------------------------|-----------------------------|
| Age at sampling (years)             |                        |                             |
| 20–24                               | 2                      | 1                           |
| 25–29                               | 6                      | 5                           |
| 30–34                               | 10                     | 5                           |
| 35–39                               | 1                      | 6                           |
| 40–45                               | 1                      | 2                           |
| Mean                                | 30.5                   | 32.6                        |
| Parity                              |                        |                             |
| 1                                   | 9                      | 14                          |
| 2                                   | 6                      | 4                           |
| 3                                   | 4                      | 0                           |
| 4                                   | 0                      | 1                           |
| 5                                   | 1                      | 1                           |
| Mean                                | 1.9                    | 1.55                        |
| BMI at end of pregnancy             |                        |                             |
| <18.5                               | 0                      | 0                           |
| 18.6–24.9                           | 3                      | 4                           |
| 25.0–29.9                           | 10                     | 6                           |
| >30.0                               | 7                      | 10                          |
| Mean                                | 28.21                  | 31.66                       |
| Smokers                             |                        |                             |
| 0                                   | 0                      | 2                           |
| Gestational age (average week)      |                        |                             |
| Placenta weight in g (average)      | 505.2                  | 383.5                       |
| Grannum score                       |                        |                             |
| 0                                   | 0                      | 0                           |
| 1                                   | 10                     | 12                          |
| 2                                   | 8                      | 8                           |
| 3                                   | 2                      | 0                           |
| Mean                                | 1.6                    | 1.4                         |

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Boxplots (A–F) showing the concentration of Na, Mg, P, K, Ca, and Fe in the placental tissue of the control (red) and PE (green) placenta group as determined by ICP-OES. Outliers are displayed as filled black dots. The entire distribution of data points is shown as gray transparent dots. The black line displays the median and notches the 95% confidence interval of the median. For all displayed elements except Fe, control and PE group did not show significant differences in the mean ($p > 0.05$), also indicated by their overlapping notches.

unit (HU) (see Supplementary Tables 3, 4 and Supplementary Figure 3). This analysis showed that both the calcified volume and the CT density of this volume (Ca HU) were significantly lower in the PE group ($p = 0.02$ and $0.0003$, respectively). Representative early onset and a late onset placentae selected based on calcification content and clinical criteria (PE symptoms, Table 2) are displayed as reconstructed tomograms to illustrate the patterns of placental calcifications (Figure 3). Overall, mineral deposits in moderately calcified placentae (including both PE placentae) were predominantly found on the basal plate and in the periphery of the maternal side, with no apparent difference between normotensive and PE placentae. To illustrate the high variability in the extent of calcification, a heavily calcified placenta found in the control group was included in Figure 3. This placenta stemming from a clinically healthy mother (40 + 4 gestational weeks) showed extensive calcification, which reached from the basal plate up to the borders of the cotyledons, forming a cloud-like pattern.

Histology and Correlation With Nanoanalytical Characterization
Following CT, we examined conventional HE- and von Kossa-stained histological sections collected from all the placentae. In agreement with the CT examination, most calcifications were located in the decidua on the maternal side of the placenta. Additional calcifications were also found in the intervillous space, in villi and in the periphery of the placenta, and in the central and paracentral part. Calcifications detectable in histology appeared either solitary, grouped or as large chunks. Selected placentae were additionally characterized with high-resolution techniques with chemical sensitivity, including micro X-ray fluorescence ($\mu$XRF) and whole slide scanning electron microscopy (WS-SEM). $\mu$XRF maps of placental tissue indicated calcium and phosphorus accumulations predominantly on the maternal side in the parenchyma (Figure 4). A reasonably good spatial correlation of the Ca and P detectable in $\mu$XRF with the visible calcifications in the HE and von Kossa staining was found (Figure 4). However, the spatial resolution of histology and $\mu$XRF imaging is limited to $> 1\, \mu\text{m}$, and hence no information on mineral morphology can be extracted. In contrast, WS-SEM enables the investigation of full histological slides with a spatial resolution of around 5 nm. By measuring back-scattered electrons, differences in atomic number (between soft tissue and minerals) can be assessed and mineral deposits can be readily identified as bright (electron-dense) spots. Due to the high spatial resolution of WS-SEM, mineral morphology can be assessed and correlated to histoanatomy.

Using WS-SEM and Density Dependent Color-SEM (DDC-SEM), four types of distinct morphologies of calcifications were identified (Figure 5 and Supplementary Figures 1, 2).
These subtypes can be described as needle-like structures (Figure 5I), large spherical structures formed by concentric rings merged together forming larger minerals (Figure 5II), rough spherical particles (Figure 5III), which in some cases were also accumulating to form larger calcifications, and large calcification blocks of no distinct morphology (Figure 5IV). Again, and in line with CT and histological analyses, calcifications were found in the inter/intravillous space and the basal plate. Importantly, also much smaller deposits were readily detected and identified in WS-SEM, way beyond the CT detection limit. For example, micron-sized particles were found in the decidua beneath the subepithelial tissue of the amniotic membrane in portions of fetal tissue (Figure 6). Energy-dispersive X-ray spectroscopy showed that all types of calcifications were predominantly composed of Ca, P and O and contained Mg, albeit at variable amounts (Figure 5 and Supplementary Figure 2). X-ray emission related to N, S and Na stemmed mostly from surrounding/underlying tissue, whereas C originated from both tissue and sample support (Supplementary Figure 2).

**DISCUSSION**

Our analysis of major and trace elements in the placenta suggests that several metals (As, Cd, Ni, Pb, Al, Mn, and Co) are likely
to be associated to the occurrence of PE as they were elevated in PE placentae. In the case of Rb, a deficiency in this element might be related to PE occurrence. Essential major elements (Ca, P, Mg, Na and K) and all other analyzed trace elements (Cr, Cu, Zn, Se, Sr, Ag, Cs, and Ba) were comparable in both groups. The comparable concentrations of Ca, P, and Mg in the two groups indicates a subordinate role of mineralization in PE, despite the disorder’s hypertensive origin. As total elemental analysis does not discriminate between particulate Ca, P and Mg and their freely dissolved, adsorbed or complexed states in the tissue, we further characterized the placentae using CT. Strikingly, both the total volume as well as the overall crystallinity/density of the calcifications were significantly higher in the control group. This indicates, that the development of calcifications in our study population was predominantly related to tissue aging rather than to the occurrence of PE.

Further nanoscale examinations of calcifications enabled identification of four distinct general types of calcifications [(I) needle-like structures, (II) large spherical calcifications formed by concentric rings, (III) small spherical rough particles, (IV) large block calcifications of no specific internal structure] which occurred in tissue of both normotensive and preeclamptic patients. In addition, the elemental composition of these deposits was very comparable between the study groups. Additionally, type III particles were found in both maternal and fetal tissue regions, including the decidua beneath the subepithelial connective tissue of the amnion. This gives an indication that not all calcifications in the placenta might be related to tissue aging.

Taken together, our results consistently suggest a subordinate role of calcifications in PE and pinpoint to a significant contribution of several trace elements and thus environmental factors to the development of PE.

While studies using bulk tissue analysis of the placenta are scarce, previous work on the role of several environmentally relevant metals in preeclampsia allows contextualization of our findings. Our data are in line with a previous study by Wang et al., which found that higher maternal blood concentrations of multiple metals, including Cr, As, Hg, and Pb, were associated with PE (11, 37). In particular, arsenic has previously been associated with adverse pregnancy outcomes, including low birth weight, anemia and spontaneous abortion (38). Also Cd is well-known to accumulate in the placenta (39). Higher levels of Cd have previously been also observed in placentae with parenchymal calcifications, regardless of smoking habit,

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**FIGURE 3** | Computed tomography images of three representative placentae (a late onset PE placenta, an early onset PE placenta and a calcified control placenta, see Table 2 for their characteristics) in maternal, fetal and sagittal view. Soft tissue is shown in red while calcifications appear in white.
gestational age, or birth weight (39). It is known that Cd can cause oxidative stress and may thus contribute to the development of PE (40). Indeed, increased placental tissue Cd levels have also been observed in PE cases in previous studies (14, 15). Cd levels in human placenta in studies between 1976 and 2011 ranged between 1.2 and 53 ng/g (39). Studies from 1978 to 1991 report concentrations of 2.6 up to 32 ng/g (39). Mean Cd concentration in placentae collected in TN, United States was 19 ng/g with a maximum value of 84 ng/g (41). Hence, these values are well in line with the ones observed in our study (median 6–30 ng/g control group and 10–83 ng/g PE group, Supplementary Table 1). Previous studies showed increased levels of Pb and Al in urine and serum of PE patients (32, 42, 43). Our results for the first time show that Al and Pb are also increased in placental tissue of PE patients. Pb and Al are known for inducing vasoconstriction and increasing the circulating levels of endothelin, which can induce PE (37, 42–44). Manganese plays a crucial part during the development of the nervous system and higher manganese concentration in maternal blood had previously been associated with lower risk of preeclampsia (44). Interestingly, in our cohort, Mn was higher in the PE group on a bulk level. Previous studies have indicated the strong concentration-dependent effect of Mn. Both deficiency in Mn as well as overexposure can increase reactive oxygen species generation and contribute to oxidative stress (45). With regard to Ni and Co, the concentrations were also elevated in the PE group compared to the normotensive control. There are few data regarding Ni and Co and their influence on preeclampsia. However, it is known that Ni can also cause oxidative stress and Co can cause pregnancy-associated hypertension, which can ultimately also lead to PE (46, 47). Rubidium as the only element that was reduced in the PE group has previously been associated with reduced fetal weight and pathological uterine Doppler (48).

Regarding distribution, mineral composition and ultrastructure of calcific deposits we found comparable results in both groups. Interestingly, spherical calcium phosphate deposits similar to those we detected in both maternal and fetal tissue were previously also found in osteocyte lacunae (49) and aortic tissue (16) and might thus be a physiological component of tissue.

The above findings suggest only weak trends of increased placental calcification in line with studies investigating calcifications in PE based on conventional clinicopathological studies (23, 50, 51). While several studies concentrated on ultrasound findings and correlated the amount of calcifications to the outcome of the pregnancy (20, 22) and drew a connection to increased stillbirth if the calcifications were seen before the
FIGURE 5 | Whole slide scanning electron micrographs (WS-SEM) and corresponding HE histology micrographs from three representative placentae. From left to right: late onset PE, early onset PE and calcified control placentae. High magnification DDC-SEM illustrate the four characteristic subtypes of calcifications observed in the tissue samples: (I) needle like structures, (II) large spherical calcifications formed by concentric rings, (III) small spherical rough particles, (IV) large block calcifications of no specific internal structure (for higher detail images see also Supplementary Figure 1). Their localization is indicated by red boxes on the WS-SEM images. Energy-dispersive X-ray spectra (EDXS) indicate comparable composition of the particles (location of analysis indicated by white boxes), which mainly consist of Ca, P, and O with traces of Mg. Spectra obtained from surrounding non-calcified tissue support this statement (see Supplementary Figure 2).

36th gestational week (20), these results cannot be compared to our study as they concentrate on calcifications in general and not in preeclampsia in particular. A recent review again showed associations between Grannum 3 calcifications and PE (52). Although the numbers of examined placentae in the reviewed studies is higher compared to this work, their results are based exclusively on the Grannum classification, which is known to have severe limitations, including observer bias (53). Also in our study, Grannum classification was not consistent with the quantitative data obtained from CT analysis (see Supplementary Figure 4). Thus, overall, we do not consider Grannum Score to be a suitable endpoint for assessing the extent of placental calcification for scientific purposes due to its aforementioned limitations. Sonographic 2D/3D imaging of the entire placenta may be a more suitable alternative. Alternatively, magnetic resonance imaging might offer additional insight and can also be performed during pregnancy (54), in sharp contrast to x-ray CT which is limited to postpartum analyses of the placenta due to radiation.

While the association of calcifications and PE remains controversial, and some studies suggest a positive correlation, we found no such association in our study. This could either be related to the significantly lower sample number, or a more direct analysis of calcifications by our materials characterization methods, making the quantification of minerals less prone to artifacts.

The good agreement with the available literature shows robustness of the metallomics approach in the bulk tissue and makes it a powerful route to profile lifestyle and environmental factors in pregnancy. The placenta as a whole, and especially also the metallomics fingerprint might serve as a time capsule containing highly relevant information on exposure to toxins and nutrient deficiencies encountered during pregnancy. Metallomics profiling thus offers a powerful and readily accessible route to
investigate the implications (including life time risks) of such exposure in a comprehensive manner.

In addition to the bulk tissue analysis, we present a whole slide imaging approach with nanometric resolution to identify and characterize solid inorganic deposits in the placenta in their histoanatomical context with limits way beyond conventional histology. To the best of our knowledge, the present work is the first study to assess the ultrastructure and chemical composition of calcifications found in preeclampsia compared to normotensive placentae. In particular, WS-SEM with nanometric resolution and chemical sensitivity offers a powerful route to overcome the limitations of contemporary histology in assessing mineral deposits in the placenta but also a variety of other tissues prone to calcification. Using our multi-modal and multi-scale characterization approach, we are able to provide the full histoanatomical context to mineral structures, which in this case allowed us to directly compare calcific deposits found in inter/intravillous space regarding their distance to the basal plate (maternal side) and the amniotic (fetal) side and thus to assign them to either maternal or fetal-tissue origin. Further, this study demonstrated the presence of nanometer sized calcific deposits in every analyzed placental tissue that were not detectable by conventional histopathology.

While we have been able to confirm important differences in heavy metal contents, the small study population has an intrinsic risk for type II errors. Additionally, dietary habits and intake of dietary supplements may pose confounding factors, potentially being a root cause for increased concentration of Fe in the PE group. Also the fact that our control group represents healthy placentas at term while the study group includes diseased placentae from fetuses that were delivered prematurely poses a relevant yet unavoidable bias.

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Additionally, the increased heavy metal concentrations found in placental tissue from preeclamptic patients are likely a direct consequence of increased exposure of the mother (55). However, we do not have complete anamnestic data on residence, working environment, passive smoking to conclusively show a correlation.

In summary, our metallomics analysis of normotensive and preeclamptic placentae is in agreement with the literature available for subsets of elements and demonstrates that heavy metals found elevated in blood and urine of PE patients are also detected at increasing levels in placental tissue, placing emphasis on their relation to the occurrence of the multifactorial disease. Thus, from a clinical perspective, the metallomics fingerprint analysis might serve as a time capsule containing highly relevant information on exposure to toxins and nutrient deficiencies encountered in pregnancy, offering a new route to early identification of patients at risk based on metallomics analyses of blood or urine during pregnancy. With regard to calcifications we show that, while no difference in Ca, P, and
Mg was observed on a bulk level, the macroscopic particulate fraction was even lower in PE placentae which is an indication that placental age is the dominating factor leading to high particulate and crystalline Ca fractions. Additionally, our study adds further evidence for the limitations of the Grannum score, as we found a poor correlation between Grannum score and direct quantification of calcifications based on CT data (see Supplementary Figure 4).

In general, the methodology presented here may in future prove highly beneficial in accessing information stored in placental tissue and assessing its implications with regard to health later in life.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Ethics Commission of Eastern Switzerland (EKOS 2020-01387). The patients/participants provided their written informed consent to participate in this study.

**AUTHOR CONTRIBUTIONS**

TR was involved in the study design, collected, and analyzed clinical data. ET performed optical and electron microscopy studies and analyzed the data. YE and SL collected, analyzed, and interpreted radiological data. DB and WJ performed the histopathological analysis of placental tissue. AA developed the elemental analysis sample preparation method. JK and TF helped with sample and clinical data collection. RH contributed to the study design of the clinical study. AG developed analytical procedures for elemental and µXRF analyses. TR together with ET, AG and IH wrote the manuscript. IH conceived the study and supervised the work. All authors reviewed and approved the final version of the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmed.2022.857529/full#supplementary-material
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