Vivianite and Its Oxidation Products in Mammoth Ivory and Their Implications to the Burial Process

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ABSTRACT: The phase composition and distribution characteristics have been obtained from two mammoth ivory samples with typical blue and yellowish-brown outer layers. The results reveal that hydroxyapatite, newberyite, organic matter, and quartz exist in all structures of mammoth ivory. Vivianite and santabarbaraite mainly contribute to the blue and yellowish-brown oxide layers of mammoth ivory, respectively. Meanwhile, metavivianite also occurs and partly influences the appearance of oxide layers. Vivianite is a common and complex product that can be formed by the interaction of gradually infiltrated Fe$^{2+}$ and the original PO$_4^{3-}$ in mammoth ivory. At the later stage, vivianite can be oxidized into metavivianite and santabarbaraite. As a result, mammoth tusks present dark bluish-green and yellowish-brown appearances. The multi-colored oxide layers are formed by different contents of vivianite and its oxidation products, which also provides valuable information on the relative burial intensity and time in different structures. It is inferred that the burial intensity increases in the sequence of yellowish-white dentin → blue outer layer → yellowish-brown outer layer. These observations are hopeful to be widely used in evaluating the changeable burial environment and exploring historical events that occurred on mammoth ivory.

INTRODUCTION

In paleontology and archeology, researchers have observed the chemical composition and physical characteristics of mammoth ivory to obtain information about environmental evolution in the Upper Paleolithic. Based on the symbolic ancient mammoth ivory tools and artifacts, the origin and evolution of ancient technologies and cultural differences of different communities were identified. The woolly mammoth is an extinct species of the Elephantidae with its tusks mostly buried in ice sheets or frozen soil in the northern hemisphere. The curved mammoth tusks generally present a light yellow to white color with a yellowish-brown surface darkened by burial. With the wide application of mammoth ivory in gemology, its mineral composition is gradually understood by researchers. The dentin of mammoth tusks is composed of hydroxyapatite (Ca$_5$(PO$_4$)$_3$(OH)), newberyite (MgHPO$_4$·3H$_2$O), organic matters, and other minerals. After the outer layer of tusks became yellowish brown by burial, the crystallinity of hydroxyapatite increased, and the organic components partly lost, which can reflect the mineralization degree of mammoth ivory. Recently, the disseminated blue substances have been found on the surface of unearthed mammoth ivory. However, their mineral and chemical composition characteristics have not been reported in detail. In this paper, two typical mammoth ivory samples from Yakutsk, Russia, were taken as the research objects. Each mammoth ivory contains yellowish-white, blue, and yellowish-brown regions (Figure 1a). The mineralogy, chemical composition characteristics, and their influence on the appearance of representative mammoth ivory samples were investigated by petrography, spectroscopy, and chemical composition analyses. The results provide evidence to indicate the burial diagenetic processes in the blue and yellowish-brown outer layers of mammoth ivory.

MATERIALS AND METHODS

Eleven mammoth ivory samples with the yellowish-white dentin and blue and yellowish-brown outer layers were unearthed from the permafrost in Yakutsk, Russia, by Wannian Jixiang (Beijing) Jewelry Co., Ltd. Two representative polished samples, i.e., cross section and longitudinal section, were selected as the experimental samples in this paper, numbered MI-1 and MI-2, respectively. The transition layer between the yellowish-white dentin and yellowish-brown outer layer...
presents blue and bluish-green appearances. To describe this transition layer more conveniently, it is represented by the blue outer layer because the blue color is the most representative.

The scraps of the yellowish-white dentin and blue and yellowish-brown outer layers were separately pulverized to 300 mesh powders for powder X-ray diffraction (XRD) analysis. The powder XRD data were collected with a Rigaku D/Max-RC diffractometer. The instrument was equipped with a conventional copper target X-ray tube set to 40 kV and 200 mA and a graphite monochromator. The scanning speed was 7.2°/min in the test range of 3° < 2θ < 70°. The data were processed by using the Search-Match software package with the International Center of Diffraction Data (ICDD).

The infrared spectra (IR) of the yellowish-white dentin and blue and yellowish-brown outer layers were collected using a Tensor 27 Fourier infrared spectrometer. The acquisition was set at 32 scans in the range of 4000–400 cm⁻¹. The resolution was 2 cm⁻¹. KBr compression and transmission modes were used in the experiment.

A HORIBA HR-Evolution Raman spectrometer was adopted to measure the Raman spectra for the different color regions of MI-1 under the conditions of 50× magnification objectives and 50–1250 cm⁻¹ test range. The system was equipped with 532 and 785 nm solid stage lasers. The acquisition time was 20 s, the accumulation was 3 scans, and the minimum pixel was 1 cm⁻¹.

The elemental maps of MI-1 and MI-2 were obtained using a Bruker Micro-XRF spectrometer (M4 TORNADO). The spectrometer was equipped with an X-ray tube (Rh anode) with polycapillary X-ray optics. The system worked at 50 kV and 599 μA, with chambering in air at 1.5 mbar and an acquisition time of 10 ms per pixel. The data were processed by the mineral analyzer software for M4 TORNADO.

The O (1s), P (2p), and Fe (2p) spectra of the yellowish-white dentin and blue and yellowish-brown outer layer powder...
samples were acquired using a Thermo Fisher Scientific Escalab 250Xi spectrometer configured with the 650 μm square test regions and an Al Kα2 radiation source (photon energy 1486.6 eV). The C (1s) peak (284.6 eV) was used to calibrate the position of binding energy. The equipment worked at a vacuum of 1 × 10−7 mbar and a constant energy of 50 eV. The CasaXPS software, Shirley background, and Gaussian:Lorentz function = 60:40 were used to fit the XPS data.

### RESULTS

#### XRD

In this paper, the representative sample MI-1 was selected for the XRD test. The powder XRD data for three regions of MI-1 are shown in Figure 2. Mammoth ivory is mainly composed of amorphous hydroxyapatite and organic material, so the obtained spectra present an unavoidable noise. The yellowish-white dentin and yellowish-brown outer layer of MI-1 are consistent with the characteristic diffraction peaks of hydroxyapatite (ICDD PDF 09-0432) in the (110), (020), (200), and (211) crystal planes, respectively. An additional weak reflection peak at \(d_{(111)} = 2.80 \text{ Å} \ (\theta = 31.99^\circ)\) is attributed to hydroxyapatite. It indicates that the blue outer layer of MI-1 is mainly composed of vivianite and smaller amounts of hydroxyapatite. There are two other peaks at \(d_{(100)} = 4.25 \text{ Å} \ (\theta = 20.86^\circ)\) and \(d_{(101)} = 3.34 \text{ Å} \ (\theta = 26.65^\circ)\) for the yellowish-white dentin and blue outer layer of MI-1. These peaks are consistent with the data for quartz (SiO₂) (ICDD PDF 83-2465), showing that quartz exists in both regions.

#### Microscopic Observation

The developed fissures and irregular curves are presented on the rough and dry surface of MI-1 (Figure 3a). The blue, bluish-green vivianite and other substances are exposed after the loosely structured yellowish-brown outer layer was shed (Figure 3a). The colors of the cross section from inside to outside are yellowish-white, blue, and bluish-green, and yellowish-brown in that sequence (Figure 3b). The dentin is light yellow to white, with a roughly parallel wavy texture. Vivianite and other substances invade the dentin in a point distribution. They are mostly concentrated in the blue outer layer or the gaps connected with the external environment. The yellowish-brown outer layer of mammoth tusks continuously and uniformly form external to internal spreads (Figure 3b).

In the above three regions, the local fibrous structural units are the first-order (1st) dark gray interference color and parallel extinction under cross-polarized light (CPL). They are deduced to be hydroxyapatite (Figure 3c). Vivianite presents the 1st yellowish-white interference color and an obvious pleochroism of blue (Np)-colorless (Nm)-light brown (Ng) under CPL. In addition, it shows the moderate relief under plane-polarized light (PPL) (Figure 3d). The results show that vivianite is mainly distributed in the blue outer layer, and it is also found in the yellowish-white dentin (Figure 3c,d). Quartz cannot be accurately identified under a polarizing microscope because of its low content and similar interference color with hydroxyapatite.

#### Infrared Spectra

The IR spectra of the yellowish-white dentin and blue and yellowish-brown outer layers of MI-1 are shown in Figure 4. In the three IR spectra, a band at around 3250 cm⁻¹ is observed. It is attributed to the O−H stretching vibration, which is associated with the adsorbed water, vivianite, and its oxidation products. The peak at 1450 cm⁻¹ and the overlapping band centered at 1000 cm⁻¹ are due to the P−O symmetric and antisymmetric stretching modes in each spectrum, respectively. Furthermore, the three IR spectra all exhibit the P−O shear bending components at 560 cm⁻¹.

The peaks at 1650 and 1538 cm⁻¹ separately appear for spectra 1 and 2, which are both caused by the N−H bending vibration and C−N stretching vibration of amide. The shoulder peak at 475 cm⁻¹ for spectrum 2 is associated with the Fe−O stretching vibration. Meanwhile, the peaks at 777 and 475 cm⁻¹ of spectrum 3 are attributed to the Fe−O−H vibration and Fe−O vibration, respectively. These peaks demonstrate that iron phosphate exists in the blue, yellowish-

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Microscopy images of MI-1: (a) surface, (b) Cross section with the yellowish-white dentin (1), blue outer layer (2), and yellowish-brown outer layer (3). (c,d) MI-1 thin-section under CPL and PPL. Cross symbols label the Raman test locations.

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Infrared spectra of the yellowish-white dentin (1), blue outer layer (2), and yellowish-brown outer layer (3) in MI-1.
brown outer layers. The presence of metavivianite (Fe$^{2+}$Fe$^{3+}$$_2$(PO$_4$)$_2$(OH)$_2$·6H$_2$O) in the yellowish-brown outer layer is indicated by the Fe···O–H bending vibration at 777 cm$^{-1}$. In addition, the band near 1635 cm$^{-1}$ is attributed to the bending vibration of the adsorbed water, proving that the yellowish-brown outer layer contains santabarbaraite (Fe$^{3+}$$_3$(PO$_4$)$_2$(OH)$_3$·5H$_2$O). Raman Spectra. The Raman spectra recorded for the yellowish-white dentin, blue outer layer, and yellowish-brown outer layer of MI-1 are shown in Figure 5. In each of the four Raman spectra, the antisymmetric stretching vibration of P–O is observed in the 1090–970 cm$^{-1}$ region, while a strong band at around 950 cm$^{-1}$ can be attributed to the P–O symmetric stretching. Furthermore, there are two bands near 500 and 320 cm$^{-1}$ for the four Raman spectra. These bands fit the previously published results for the P–O in-plane and out-of-plane bending, respectively. In spectrum 1, the weak protein features could be observed at the bands of 890, 870, 855, and 815 cm$^{-1}$. In addition, the vibrations at 206, 148, and 124 cm$^{-1}$ were ascribed to the Fe–O stretching components in the three Raman spectra of the blue outer layer and yellowish-brown outer layer.

Based on the above results of peak assignments, spectrum 1 shows characteristic bands that are associated with phosphate and organic matters. Spectra 2-1 and 2-2 correspond to vivianite and metavivianite, respectively. Spectrum 3 corresponds to metavivianite. The bands will overlap each other when santabarbaraite exists in the yellowish-brown outer layer. Determining the presence of santabarbaraite needs to combine with other test methods, such as XPS.

Elemental Mapping by $\mu$-XRF. The blue and yellowish-brown regions disseminate toward the direction of dentin in MI-1 and MI-2. To obtain the chemical composition information of different structures, $\mu$-XRF mapping was carried out on the regions with different structures coexisting shown in Figure 1b,c, and the results are shown in Figures 6 and 7, respectively.

Remarkably, the distribution of oxygen corresponds well to that of silicon in MI-1 and MI-2. Based on the XRD and IR results, it is speculated that quartz exists in all three regions of MI-1 and MI-2, and it is more concentrated in the yellowish-brown outer layer. As can be seen on the mappings, magnesium and phosphorus are mainly distributed in the yellowish-white dentin of MI-1 and MI-2, indicating the distribution characteristics of newberyite (MgHPO$_4$·3H$_2$O). In the case of calcium, it is also concentrated in the yellowish-white dentin followed by the yellowish-brown outer layer and blue outer layer in that sequence. In conclusion, it is consistent with the distribution characteristics of hydroxyapatite.

In contrast, manganese and iron are centered in the blue and yellowish-brown outer layers, and they also present enrichment along the cracks. It is ascribed to the presence of vivianite and its oxidized products. It is also referred that manganese ions may replace a part of iron ions into the mineral lattice. In addition, manganese and iron show an inverse correlation pattern with calcium.

The environmental titanium is concentrated in the yellowish-brown outer layer and oriented along the cracks in MI-1 and MI-2. XPS. The XPS spectra were collected on the typical dentin and blue and yellowish-brown outer layers of MI-1 shown in Figure 1b, and the results are given in Figure 8. The existence of surface adsorbed oxygen is revealed by the peak at 532.6 eV.

Figure 5. Raman spectra for representative sites in the yellowish-white dentin (1), blue outer layer (2-1) (2-2), and yellowish-brown outer layer (3) of MI-1.

Figure 6. Elemental maps produced by $\mu$-XRF of the yellowish-white dentin (1), blue outer layer (2), and yellowish-brown outer layer (3) in MI-1.

Figure 7. Elemental maps produced by $\mu$-XRF of the yellowish-white dentin (1), blue outer layer (2), and yellowish-brown outer layer (3) in MI-2.
in the three O (1s) spectra (Figure 8a−c).38 The peak at 531.9 eV is caused by oxygen in Mg=O, while the peak for phosphate and hydroxyl occurs near 531.1 eV.38,39 Thus, according to XRD and μ-XRF mapping results, phosphate, newberyite, and surface adsorbed water can be detected in the three different color regions of MI-1.

In the three XPS spectra of P (2p) (Figure 8d–f), the peak ranging from 132.6 to 133.2 eV corresponds to the phosphorus in PO₄³⁻, and the peak at around 133.3 eV is attributed to the phosphorus in HPO₄²⁻.40,41 It demonstrates that phosphorus exists mainly in the form of hydroxyapatite in the three regions of MI-1. It is also inferred that the yellowish-white dentin has the greatest amount of newberyite.

The Fe (2p) spectra were fitted by using the ferrous and ferric multiplet patterns. The results of XPS and peak fitting (Figure 8g–i) show that the yellowish-white dentin almost contains no iron. The peaks at around 711.7 and 710.1 eV are due to the octahedral coordination of Fe^{3+} and Fe^{2+}, respectively. The satellite peaks of Fe^{3+} and Fe^{2+} occur in the range of 715.4−716.6 and 713.4−713.8 eV.42 The spectra of Fe (2p) indicate the distributions and relative contents of vivianite, metavivianite, and santabarbaraite.

**DISCUSSION**

Two typical mammoth ivory samples from Yakutsk, Russia, with the yellowish-white dentin and blue and yellowish-brown outer layers from inner to outer were investigated by XRD, vibration spectroscopy, μ-XRF mapping, and XPS. Based on the experimental results, the yellowish-white dentin is mainly composed of hydroxyapatite, newberyite, organic matters, and quartz. The blue outer layer is located below the yellowish-brown outer layer and disseminated toward the dentin. Its main constituent minerals are vivianite and metavivianite, with a small amount of hydroxyapatite, newberyite, quartz, and organic matters. Meanwhile, the yellowish-brown outer layer contains santabarbaraite, metavivianite, hydroxyapatite, newberyite, and quartz.

Vivianite is an easily oxidized hydrated ferrous phosphate mineral. According to the oxidation degree from low to high, its oxidation series can be described as follows: vivianite (monoclinic) → metavivianite (triclinic) → santabarbaraite (amorphous). This process is accompanied by color changes from blue to bluish green then to yellowish brown.43−45 In archeology and geology, researchers can judge the sedimentary environment and the formation process of vivianite on the basis of the spatial distribution of vivianite and its oxidized products.44−51 According to the chemical composition and
oxidized degree of vivianite, some scholars speculated the origin and preservation integrity of works of art.\textsuperscript{52,28,53} The results of $\mu$-XRF mapping and XPS show an apparent iron content in the blue and yellowish-brown outer layers. To further analyze the oxidation degree of vivianite in these two layers, the Fe (2p) peak fitting data are given in Table 1. It demonstrates that a part of vivianite has been oxidized in varying degrees. The Fe$^{2+}$ and Fe$^{3+}$ account for 47.3 and 52.8%, respectively, in the blue outer layer, consistent with the result that there are vivianite and metavivianite simultaneously. The values of 19.4% Fe$^{2+}$ and 80.6% Fe$^{3+}$ are detected in the yellowish-brown outer layer. It can be explained by the presence of santabarbaraite and metavivianite in this region. The content difference of Fe$^{2+}$ and Fe$^{3+}$ between the blue and yellowish-brown outer layers reflects the composition distinction of vivianite and its oxidation products. According to their distribution characteristics and relative content differences, we conclude that there was almost no oxidation in the yellowish-white dentin. Meanwhile, the blue outer layer is oxidized to a certain extent, and the yellowish-brown outer layer has the highest oxidation degree. The yellowish-brown outer layer is closer to the external environment so that it is strongly oxidized by the external burial environment.

The experimental results of XRD and $\mu$-XRF mapping show that quartz exists in the three different color regions of mammoth ivory. Also, it is mainly concentrated in the yellowish-brown outer layer. The hydroxyapatite matrix is porous in mammoth ivory, which is beneficial to the infiltration of chemical substances from the surrounding environment.\textsuperscript{57} It is speculated that quartz gradually penetrated into mammoth ivory from surrounding rocks or soil. The formation of vivianite requires specific material sources and environmental conditions. The spatial distribution of vivianite shows that the Fe$^{2+}$ in vivianite is provided by the external environment, while the PO$_4^{3-}$ in vivianite originates from mammoth ivory under certain burial effects.\textsuperscript{54} In an anoxic frozen soil environment, when Fe$^{2+}$ reaches a specific concentration, it will interact with PO$_4^{3-}$ to form vivianite. With the burial time going on, Fe$^{2+}$ will gradually penetrate into the inside of mammoth ivory and form a gradually thickened vivianite layer. After the formation of the vivianite layer, the burial environment of mammoth ivory, such as the redox conditions, microbial community, temperature, and pressure, may change. Then, vivianite will be oxidized to a series of oxidation products in varying degrees, including dark bluish-green metavivianite, yellowish-brown santabarbaraite, and so on.\textsuperscript{55} These processes are also carried out along the direction from outside to inside of mammoth ivory, so there will be a phenomenon that the oxide layers of different colors coexist. The study on the formation and oxidation processes of vivianite can provide some powerful evidence for inferring the burial environment change. With the burial time going on, the blue and yellowish-brown outer layers will further expand toward the center of mammoth ivory. By analyzing the proportion of the blue and yellowish-brown regions in different mammoth tusks under similar conditions, we can compare their relative burial time and preservation integrity.

### CONCLUSIONS

The cross section of a typical mammoth ivory sample, unearthed from Yakutsk, Russia, presents the yellow-white, blue, yellowish-brown layers from inside to outside successively. The changes of color in different regions correspond to the distinction of the mineral species and content. The yellowish-white dentin is composed of hydroxyapatite, newberryite, organic matters, and a small amount of quartz. The blue outer layer is located under the yellowish-brown outer layer and extends to the dentin. Its main constituent minerals are vivianite, metavivianite, and a small amount of hydroxypatite, quartz, newberryite, and organic matters. The yellowish-brown outer layer contains santabarbaraite, metavivianite, hydroxyapatite, newberryite, and quartz.

Vivianite is formed by the interaction of gradually infiltrated Fe$^{2+}$ and original PO$_4^{3-}$ in mammoth ivory. Its series of oxidation products, such as dark bluish-green metavivianite and yellowish-brown santabarbaraite, can form in the later stage. Their formation is related to the changeable burial environment, such as the oxidation–reduction environment or other conditions. The formation of vivianite and its later oxidation process are carried out from the surface to the center of mammoth ivory. Therefore, there is the phenomenon that the oxide layers of different colors coexist. The burial intensity may increase according to the following sequence: the yellowish-white dentin $\rightarrow$ blue outer layer $\rightarrow$ yellowish-brown outer layer. The difference in contents of vivianite and its oxidation products reflects the distinction of burial intensity in different color regions of mammoth ivory. By analyzing the proportion of blue and yellowish-brown regions in different mammoth tusks under similar conditions, it can provide evidence for the preservation integrity of mammoth ivory.

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**Table 1. Semi-Quantitative Calculation Results of Iron with Different Chemical Valences of the Blue Outer Layer (2) and Yellowish-Brown Outer Layer (3) in MI-1**

| sites   | name       | position (eV) | FWHM (eV) | % area |
|---------|------------|---------------|-----------|--------|
| 2       | Fe$^{2+}$oct2p3/2 | 710.45        | 2.4       | 34.3   |
| 2       | Fe$^{3+}$oct2p3/2 | 711.81        | 1.8       | 36.6   |
| 2       | Fe$^{2+}$oct2p1/2 | 723.82        | 2.0       | 13.0   |
| 2       | Fe$^{3+}$oct2p1/2 | 725.45        | 2.0       | 16.2   |
| 3       | Fe$^{2+}$oct2p3/2 | 709.63        | 1.2       | 9.7    |
| 3       | Fe$^{3+}$oct2p2/3 | 711.50        | 2.3       | 44.9   |
| 3       | Fe$^{2+}$oct2p1/2 | 723.23        | 1.5       | 9.7    |
| 3       | Fe$^{3+}$oct2p1/2 | 724.99        | 2.1       | 35.7   |
| 2 total Fe$^{2+}$ | | 47.3 | |
| 2 total Fe$^{3+}$ | | 52.8 | |
| 3 total Fe$^{2+}$ | | 19.4 | |
| 3 total Fe$^{3+}$ | | 80.6 | |

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