Investigation of the Pigments and Glassy Matrix of Painted Enamelled Qing Dynasty Chinese Porcelains by Noninvasive On-Site Raman Microspectrometry

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Received: 24 July 2020; Accepted: 14 August 2020; Published: 17 August 2020

Abstract: A selection of 15 Chinese painted enameled porcelains from the 18th century (Qing dynasty) was analyzed on-site by mobile Raman and XRF microspectroscopy. The highly prized artifacts are present in the collections of the Musée du Louvre in Paris and Musée Chinois at Fontainebleau Castle in France. In the painted enamels, pigments such as Naples yellow lead pyrochlore, hematite, manganese oxide and carbon and opacifiers such as lead arsenates were detected. The glassy matrices of the enamels mainly belonged to lead-rich and lead-alkali glass types according to the Raman spectra obtained. The glaze and body phases of the porcelain artifacts were also analyzed. The detection of lead arsenate apatite in some of the blue enamels was significant, indicating the use of arsenic-rich European cobalt ores (smalt) and possibly mixing with Asian cobalt. This characteristic phase has also been identified in French soft-paste porcelains and glass decor and high-quality Limoges enamels from the same period. Based on the shape of the Raman scattering background, the presence of colloidal gold (Au nanomaterials) was identified in red, orange and pink enamels. Different types of Naples yellow pigments were also detected with Sb-rich, Sn-rich and mixed Sb–Sn–(Zn, Fe?) compositions in the yellow enamels. The results were compared to previous data obtained on Chinese cloisonné and painted enameled metalware and Limoges enamels as well as French enameled watches.

Keywords: porcelain; enamels; China; 18th century; Raman microspectroscopy; pXRF; pigments; arsenic; cobalt; Naples yellow

1. Introduction

Technological aspects of ceramic production in the past were not only affected by the mastery and innovation of the local craftsmen but the transfer of technology coming out of cultural interactions between societies. This is well-reflected in the choice of the raw materials used and the firing conditions as well as in decorative processes such as enameling. Regarding porcelain production in the Far East, a long-distance technological transfer is known to have been conducted by Portuguese Jesuit missions in Japan at the beginning of the 17th century [1–3]. Historical records of the Chinese court and Jesuit mission [4–8] also both demonstrated the importation of European enamels and ingredients as well as
the venue of European craftsmen who had expertise in enameling techniques at the Qing court at the end of 17th century. These enamels are called “falangcái” (琅彩 from falang, a homonym of the “French” character in Chinese) and/or “yangcái”, (洋彩 “foreign colors”) [9], which are deposited generally over the glaze, the latter being fired beforehand with the porcelain body. Although porcelain production was first achieved in China historically, the craft of enameling is generally suggested to have benefited with Western contacts as early as the 14th century via the Silk Road [10–15].

Scientific analyses of various Chinese [10–15] and European [11,16–18] enameled artifacts have previously confirmed the use of a blue enamel precursor (usually called “smalt”) based on European cobalt ores and Naples yellow lead pyrochlore pigments typical of European recipes in enameled metalware and porcelains produced during the Qing dynasty. Furthermore, the use of the European “Cassius purple” method to prepare pink to purple porcelain enamels (the so-called Famille rose) with colloidal gold had been well-established for decades [19]. Due to the rareness of artifacts produced at the Beijing Imperial Palace Workshop (falangzuo (珐琅作), enamel workshop) and, consequently, the lack of fragments of such artifacts, most of the studies have been conducted with noninvasive techniques, in particular on-site with Raman and X-ray fluorescence (XRF) mobile instruments [10–14,16–18,20], which are well-suited for the in situ study of outstanding objects in their secure locations [20–24].

We present here an on-site noninvasive Raman study of Chinese porcelain masterpieces assigned to have been produced at the beginning of the 18th century (Yongzheng (1723–1735) and beginning of Qianlong (1735–1796) reigns) belonging to the collections of the Louvre Museum and Chinese Museum at Fontainebleau Castle in France. One artifact has also been analyzed with a mobile XRF microspectrometer. A comparative approach was carried out with the first Raman study focused on porcelains produced at the end of the Kangxi reign (1661–1722) and the beginning of the Yongzheng reign [12] and with painted enameled metalware of the same collections [14] as well as other types of European enameled artifacts [11,16–18].

2. Methods and Artifacts

2.1. Artifacts

The porcelains analyzed are listed in Table 1 and shown in Figures 1–5. They belong to the collections of the Musée du Louvre (Department of Fine Art Objects) in Paris and Musée Chinois at Fontainebleau Castle, which is located close to Paris. Artifacts from the Fontainebleau Chinese Museum were part of the collection of Napoléon III (1808–1873, president of the French Republic from 1848 to 1852 and then emperor up to 1870) and the Empress Eugénie (1826–1920), former president of the French Republic. Indeed, following the Universal Exhibitions, the European elites of the second part of the 19th century took a great interest in Chinese and Japanese ceramics and built up collections rich in exceptional pieces. Most of the artifacts studied here were assigned, based on a stylistic examination of their shape and decor, to have been made during the Yongzheng reign (1722–1735) or just afterwards. The corpus studied includes four cups, five plates, one milk pot, one saucer lid, one teapot and three bottles, which were analyzed by Raman microspectrometry. Only the TH457 bottle was also analyzed by portable XRF (pXRF). Some of the artifacts have rather similar counterparts in other museum collections (Table 1). Four artifacts display painted enamel decor with exceptional quality, such as the plates with roosters (TH487, Figure 1) and tigers (R1056, Figure 2) and especially the TH457 (Figure 3), F1371C (Figure 4) and F1341C (Figure 5) bottles. Close-up views of the painted flower-and-bird decor on the TH457 bottle and the flower decor on the F1341C bottle and F1429C teapot display the high sharpness of the drawing without any diffusion of the colors outside the areas delimited by (black) lines. The F1371C bottle shows a technique inspired by cloisonné enameled metalware in which the decor is first made by incision in order to avoid diffusion of the colors, somewhat similar to the sgraffito technique.
Table 1. Information about the studied artifacts regarding their collections and time assignments along with special remarks (see [25-27]).

| Artifacts | Inventory Number | Collection | Artifacts | Inventory Number | Size | Artifacts |
|-----------|------------------|------------|-----------|------------------|------|-----------|
| R1041 cup | 1722–1735        |            | R1045 cup | 1730–1745        | H:4  | Similar cup saucer at Rijksmuseum (AK-NM-13720-A) [25] |
|           |                  |            | R1048 cup | 1740–1760        | H:4  | China, Guangzhou workshop? |
| R1135 cup | ca. 1750–1760    |            | R1006 dish | 1740–1760?       | D:10 | China, Guangzhou workshop? |
| R1177 sactivex lid | Louvre Museum |            |           |                  | D:11.5 | Series with the coat of arms of an English family, S. Jones de Stepney or J. Bromwell Jones or his son or brother [25,26]. |
| SN284 milk pot |                   |            |           | ca. 1735         | H:11 |                |
| R1056 plate | 1722–1735 |            | R1175 (2) plate | 1735 | H:1.7, L:13.2 | With the coat of arms of an English family (see above) |
| R1025 dish | 1740–1750        |            |           |                  | H:3.2 | With CFB monogram |
| TH487 dish | 1730–1735        |            |           |                  |      | With 2 roosters drawn |
Table 1. Cont.

| Artifacts | Inventory Number | Collection | Artifacts | Inventory Number | Size    |
|-----------|------------------|------------|-----------|------------------|---------|
| TH457 bottle |                   | A. Thiers Coll. |           | 1735–1796       | H:30.9  |
|           |                   |            |           |                  | D:12    |
| F1371C bottle |                 |            |           | 1730/35–1750/60  | H:19    |
|           |                   |            |           |                  | D:30    |
| F1429C teapot |                | Napoléon Emperor Coll. | Fontaine bleau Castle (Musée Chinois) | Qianlong period (1736–1795) | H:13    |
|           |                   |            |           |                  | D:12    |
|           |                   |            |           |                  | A:19    |
| F1341C bottle |                |            |           | End of Yongzheng period (~1730–1735) | H:30    |
|           |                   |            |           |                  | D:15    |

Figure 1. The cups and plate analyzed. The remote head with optic fibers connected to the laser source and the spectrometer is shown at the bottom. A laser focusing on the R1041 cup exhibits the high translucency of the porcelain (see Table 1 for details).
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Figure 2. The detailed images of the plates and milk pot analyzed (see Table 1 for details).

Figure 3. The detailed images of the TH457 bottle (the neck has undergone a restoration process; see Table 1 for details).

Figure 4. The detailed images of the bottle (F1371C; see Table 1 for details).
Different grades of the hue were obtained by adding minute dots of color in the TH457 bottle (Figure 3), as also observed in the painted enameled decor of 18th-century French watches [18]. This indicates the use of very small pencils. Consequently, the thickness of the painted enamel decor remains very thin. The technique used appears to be different from the common pottery technique of adjusting the hue of enameled decor by superimposing enamel layers relatively thickly, as is visible for the R1006, R1175, R1025, R1041, R1045 and SN284 artifacts (Figures 1 and 2). The technique used for the F1429C teapot (Figure 5) is an intermediate between the two techniques.

An examination of the section of porcelain shards exhibiting similar decor by optical microscopy or of the whole objects by optical coherence tomography studies (OCT) is needed to go further in the description of the enameling technique.
2.2. Mobile Raman Microspectrometry

The high value of the porcelain artifacts made it necessary to conduct the noninvasive Raman analyses in the exhibition or storage rooms of the Musée du Louvre (Paris) and Musée Chinois (Fontainebleau Castle). A mobile Raman setup from HORIBA Scientific Jobin Yvon (Longjumeau, France) was used for the on-site measurements. This setup was composed of an HE532 spectrometer (920 lines/mm grating; resolution ~4 cm\(^{-1}\)), a remote optical device called SuperHead (Figure 1) and a 532 nm 300 mW Ventus laser source (Laser Quantum, Fremont, CA, USA), which were connected to each other by optic fibers. The SuperHead incorporates long working distance x50 (surface analyzed: ~10–20 µm\(^2\); in-depth: ~5–20 µm) and x200 (surface analyzed: ~1 µm\(^2\); in-depth: ~1–2 µm) microscope objectives for the green laser illumination and the collection of scattered light. Details about the procedures have already been published [13,17,18,20].

2.3. Mobile X-ray Fluorescence Microspectrometry

Portable XRF (pXRF) measurements were performed without any contact between the instrument and the artifact using an ELIO instrument (ELIO, XGLab/Brucker, Italy). This setup consisted of a miniature X-ray tube system with a Rh anode (max voltage of 50 kV, max current of 0.2 mA, a 1 mm collimator determining a rather similar surface) and a large-area silicon drift detector (SDD) (50 mm\(^2\) active area). Details about the procedure have already been published; it should be noted the information thickness during the analysis of the enamel was estimated to be ~4 µm at Si K\(_{\alpha}\), 130 µm at Cu K\(_{\alpha}\), 220 µm at Au L\(_{\alpha}\) and ~2.5 mm at Sn K\(_{\alpha}\) [18].

3. Results and Discussion

3.1. Body Phase Identification

Raman spectra (Figures 6–13) have been recorded on the glaze-free body for the different artifacts (see Figure 6A (R1045), Figure 7A (TH457), Figure 8B (R1175) and Figure 9A,B (R1006 and SN284)). In these spectra, the narrow main peak of quartz (SiO\(_2\)) was observed at ca. 460 cm\(^{-1}\) [28,29], shifted down to lower wavenumbers than that of the isolated quartz mineral (~463 cm\(^{-1}\)) due to the compressive stress of the glassy matrix on the quartz grains. A broader component at ca. 480 cm\(^{-1}\) was also observed (see e.g., Figures 7A and 9B), characteristic of a glassy aluminosilicate phase forming the porcelain body in between acicular mullite crystals forming a 3D network [30,31]. The mullite phase was generally not detected with a mobile Raman spectrometer due to the operating conditions of the mobile setup and poor intensity of the mullite Raman spectrum. Only in the body spectrum of the R1045 cup (Figure 6A(a)) could very small features at ~960 and 1130 cm\(^{-1}\) be considered to come from the mullite phase [28]. The Raman spectrum of the glaze, fired at a high temperature with the porcelain body, was rather similar, but the intensity of the ~490 to 505 cm\(^{-1}\) broad component was stronger, and the quartz peak was mostly weaker, as observed for the glaze spectra of the R1045 and R1041 cups (Figure 6A(b),C(b)).
spectra, the narrow main peak of quartz (SiO$_2$) was observed at ca. 460 cm$^{-1}$. Results and Discussion

In the body spectrum of the R1045 cup, not detected with a mobile Raman spectrometer due to the operating conditions of the mobile setup. Only in the body spectrum of the R1045 cup, not detected with a mobile Raman spectrometer due to the operating conditions of the mobile setup. Noted the information thickness during the analysis of the enamel was estimated to be ~4 µm at Si.

The Raman spectrum of the glaze, fired at a high temperature with the porcelain body, was rather similar, but the intensity of the ~490 to 505 cm$^{-1}$ peak corresponded to very short counting times, and 500 accumulations); and SN284 pot (a,a’,a’’), red, (e,e’) black (line); (d,d’,d”), blue, (b,b’,b”) red (eye), (c,c’) green areas.

Representative Raman spectra recorded on the body, glaze and painted enamels of the TH457 cup (a,a’,a’’), blue, (b,b’,b’’), red (eye), (c,c’), green areas. Representative Raman spectra recorded on the body, glaze and painted enamels of the TH457 cup (a,a’,a’’), blue, (b,b’,b’’), red (eye), (c,c’), green areas.

**Figure 6.** Representative Raman spectra recorded on the body, glaze and painted enamels of the porcelains: (A) R1045 cup (a) body, (b) white, (c,c’) blue; (B) R1048 cup (a) white, (b) blue, (c) yellow, (d,d’,d”), red, (e,e’) black (line); (C) R1041 cup (a) blue (flower), (b) colorless glaze, (c) pink; (D) R1135 cup (a,a’,a’’), blue, (b,b’,b’’), red (eye), (c,c’), green areas.

**Figure 7.** Representative Raman spectra recorded on the body, glaze and painted enamels of the TH457 (A) and F1371C bottles (B): (a,a’) white, (b,b’) dark blue, (c,c’) celadon (green), (d) yellow-green.
Figure 8. Representative Raman spectra recorded on the painted enamels of the TH487 plate: (A) (a,a’,a”) (marli) yellow, (b) (marli) blue, (c) orange (coat), (d) red (flower), (e,e’) (marli) black; and small plate (B) (a) blue, (b) green, (c) red (line), (d) body.

Figure 9. Representative Raman spectra recorded on the body, glazes and painted enamels of the R1056 plate: (A) (a) light yellow, (b) blue glaze, (c) colorless glaze, (d) body, (e,e’,e”) black (e’ and e” corresponded to very short counting times, and e, to 500 accumulations); and SN284 pot (B) (a,a’) yellow, (b) green, (c) red (flower), (d) pink (textile), (e) body.

Figure 10. Representative Raman spectra recorded on the painted enamels of the TH487 plate: (A) (a,a’) red, (b,b’) yellow (cock), (c) (marli) light yellow; (B) (a,a’) blue, (b) green, (c,c’,c”) black (c, very short counting time), (d) red.
Figure 10. Representative Raman spectra recorded on the painted enamels of the TH487 plate: (A) (a,a') red, (b,b') yellow (cock), (c) light yellow; (B) (a,a') blue, (b) green, (c,c',c'') black (c, very short counting time), (d) red.

Figure 11. Representative Raman spectra recorded on the glaze and painted enamels of the R1056 tiger plate (A) (a,a') white (tiger belly), (b) pink, (c) glaze; and SN284 pot (B) yellow.

Figure 12. Representative Raman spectra recorded on the F1429C teapot for different colored painted enamels: (A) white (a) and blue (b) enamels; (B) dark green (a,a') and yellow (b,b',b'') enamels; (C) pink (a,a') and red (b,b') enamels; (D) brown (a,a') and black (b,b') enamels.
3.2. Phase Identification in Colored Glaze/Enamels

Different phases were identified by Raman analysis for the different colored areas of the Chinese painted enameled porcelains, such as white, blue, yellow (and green), red (purple to red) and pink (Table 2).

Table 2. Characteristic pigments/opacifiers identified in the different colored painted enamels (overglaze) of the Chinese porcelains analyzed. The decor made with imported European recipes is expressed in bold (NY: Naples yellow, br: broad, nr: narrow, sh: shoulder, s: small).

| Color                  | Artifact       | Identified Phase(s) | Characteristic Raman Peak (cm$^{-1}$) | Remarks      |
|------------------------|----------------|--------------------|---------------------------------------|--------------|
| White (colorless glaze)| R1006, R1041, | quartz             | 455–460                               |              |
|                        | R1056, R1175,  | glassy             |                                       |              |
|                        | TH457, TH487   | aluminosilicate    |                                       |              |
|                        |                |                    | ~485 (br)                             |              |
| White (overglaze)      | F1371C, R1048, | As apatite         | ~813 (nr), ~770 (sh)                  | White type 1 |
|                        | R1056          |                    |                                       |              |
| White-pink (overglaze) | TH457          | arsenate           | 820 (br)                              | White type 2 |
|                        |                |                    |                                       |              |
| Blue (overglaze)       | R1175 (s), R1135, F1371C, F1429C | As apatite         | ~810 (nr), ~775 (sh)                  | Blue type 1  |
|                        |                |                    |                                       |              |
|                        | TH487, R1045, R1048, R1041 | arsenate           | ~815 (br)                             | Blue type 2  |
### Table 2. Cont.

| Color                        | Artifact | Identified Phase(s)                                                                 | Characteristic Raman Peak (cm⁻¹)                                      | Remarks                  |
|------------------------------|----------|-------------------------------------------------------------------------------------|-----------------------------------------------------------------------|--------------------------|
| Blue (glaze)                 | R1006    | glassy silicate (+ quartz)                                                           | 460, 503                                                             | Blue type 3 Smalt?        |
|                              | R1175, TH457 | Sb-rich NY pyrochlore                                                              | ~122/130, ~335, 508                                                  |                          |
| Yellow (overglaze)           | R1175, R1006, R1048, TH487, SN284, R1135 | Sn-rich NY pyrochlore                                                             | ~130–133, ~455                                                     |                          |
|                              | TH487, R1056 | Sn-rich NY pyrochlore + arsenate                                                   | ~133, 327, 440, ~810–820 (broad)                                    |                          |
|                              | F1429C, F1341C?, TH457 | Sb–Sn–(Zn, Fe?) NY pyrochlore + As apatite                                          | 134, 330, 450, 480–490, 518, 810, 770 (sh)                         |                          |
| Yellow-green (overglaze)     | R1135    | Sn-rich NY pyrochlore + As-apatite                                                  | ~133, 330, 815, 770                                                 |                          |
| Green (overglaze)            | R1135    | Sn-rich NY pyrochlore                                                               | ~130–133, 330, 455                                                 |                          |
| Blue-green (overglaze)       | TH487    | Sn-rich NY pyrochlore + arsenate                                                   | ~133, 330, 440, ~810                                                |                          |
| Dark yellow (overglaze)      | TH457    | Feldspar? + hematite                                                                | ~508 + 220, 292, ~1310                                             |                          |
| Red (overglaze)              | TH457, R1175, R1048, R1135 | hematite                                                                            | 220, 292, ~1310                                                   |                          |
| SN284, R1175 (large), TH487 | Au⁺ + arsenate | Fluorescence background peaking at ~500 + ~810                                     |                                                                 |                          |
| Orange (overglaze)           | TH457, SN284 | Au⁺                                                                 | Fluorescence background peaking at ~600                             |                          |
|                              | R1175, R1041, TH487 | Au⁺ + arsenate                                                                     | Fluorescence background + ~820                                      |                          |
| Pink (rose) (overglaze)      | SN284    | Arsenate + Au⁺?                                                                   | ~810 + fluorescence background at ~500                              |                          |
| Black (overglaze)            | R1041, F1429C | Au⁺                                                                              | Fluorescence background at ~500                                      |                          |
| Brown                        | R1006    | Spinel (Mn?) (+ carbon)                                                            | 625 (+1330–1565)                                                   |                          |
|                              | F1429C   | Mn oxide                                                                           | 550, 580, 892                                                      |                          |
3.2.1. Arsenic-Based Phases

In the painted enamels analyzed, the detection of two types of arsenic-based phases was significant. The first type refers to lead alkali arsenate apatite, recognized with the narrow (ca. 810–813 cm$^{-1}$) peak and a well-defined shoulder at ~770–775 cm$^{-1}$, characteristic of the As–O stretching mode of apatite, for which a composition close to Na$_{1-x}$K$_x$Ca$_y$Pb$_4$(AsO$_4$)$_3$ ($x = 0.1$ and $y = 0.5$) has been deduced in a French soft-paste porcelain [34]. This type of opacifying phase was found in some of the enamels analyzed, mostly in white and blue as well as light pink and yellow ones [11–14,17,18,34], as observed in Figure 6B(a) (R1048), Figure 6D(a,a') (R1135), Figure 7B(a,a'),(b,b') (F1371C), Figure 8A(b), Figure 8B(a) (R1175), Figure 10A(a), Figure 11A(a,a'), Figure 12A(a,b), Figure 12B(b,b','b") and Figure 12C(a') (F1429C). In the case of blue enamels, lead arsenate apatite typically forms as a result of a reaction between arsenic coming from the cobalt source (smalt) and the lead-based enamel/glaze (see further) during the firing process. Its composition also depends on various fluxing species, such as Na(K) and Ca, present in the enamel/glaze matrix [3]. Raman spectra of these enamels displayed the characteristic signature of lead arsenate apatite along with that of the glassy silicate matrix, indicating that the blue was obtained by dissolution of Co$^{2+}$ ions in the glassy matrix [29]. The bands expected due to the precipitation of cobalt aluminate or cobalt silicate (see further) were also not detected. Cobalt ores with an arsenic-rich composition are characteristic of European sources used between ~1500 and 1800 [17,18,35,36]. In particular, raw cobalt arsenide mined from the Erzgebirge (the Ore Mountains) in Saxony (Germany) were mixed with potassium glass to produce blue smallt, which was then exported as a coloring agent for glasses/enamels and paint for easel paintings during the 16th–17th centuries [3,36,37]. The detection of lead arsenate apatite could also be due to deliberate addition of an arsenic compound for opacification, especially in the case of the white enamels analyzed [34]. This phase has also been identified in the previous analytical studies of 18th–19th-century Chinese porcelains produced during the Qing dynasty [11–14] as well as in the blue decor of 17th- and 18th-century French enameled glass [16], enameled metalware [10,11,18] and soft-paste porcelains [17,34].

The second type of arsenate phase was characterized by a much broader (ca. 815–820 cm$^{-1}$) peak (Figure 6A(c,c') (R1045 blue), Figure 6B(b) (R1048 blue), Figure 6C(a) (R1041 blue), Figure 7A (TH457 white-pink), Figure 10A(c) (TH487 yellow) and Figure 13A(b') (F1341C blue)). An intermediate shape between those of the two arsenate phases was observed for Figure 9B(d) (SN284 pink). This broad arsenate band has already been observed in Kangxi Chinese porcelains for the blue areas [12–14] as well as certain French enameled productions [11,16–18]. The broadening of the As–O stretching modes may have arisen from the poor crystallinity or small size of the arsenate apatite crystals or from structural distortions differentiating the AsO$_4$ tetrahedra (only one strong peak with A$_1$ symmetry is expected for the stretching mode of an XO$_4$ tetrahedron). However, arsenates with different composition and structures from apatites (e.g., feldspar) have also been identified in Italian and Spanish majolica enamels [38–41]. Microdestructive analyses to be performed on porcelain shards, such as μ-diffraction, μXRF and/or transmission electron microscopy analyses, are needed to identify more precisely the arsenic-based phases given the ca. 815 cm$^{-1}$ broad band.

The two signatures observed in the blue enameled areas indicated the use of cobalt ores rich in arsenic, i.e., cobalt ores imported from Europe [11,15–18] or a mixture of Asian (Mn- and Fe-rich) [3,13,36,42–46] and European (As-rich) cobalt ingredients. However, the rareness—and high cost—of European ingredients led Japanese potters, and likely Chinese ones, to mix with Asian ingredients. This point will be discussed further using the XRF spectra obtained.

3.2.2. Lead Pyrochlore (Naples Yellow)

A second important outcome of this study was the detection of at least two types of Naples yellow lead pyrochlore pigments in the yellow and green painted enamels of the porcelains. Naples yellow refers to a variety of synthetic lead antimonate pigments with a pyrochlore structure, prepared by mixing lead oxide with various compounds, including mainly antimony and tin, in the presence of various fluxing agents [1,3,12,13,18,47–54]. This pigment, in the form of Sn- or Sn-rich types,
has been used as a glass and glaze pigment since antiquity [55–57] and as a painting pigment after the Quattrocento/Renaissance [53,54]. Possible variations of the pigment might also include the addition of zinc, iron and silicon elements [14,18,49,50]. The adjustment of the proportions of these compounds at varying firing temperatures allows for the production of modified forms of the pigment displaying different hues. The composition of Naples yellow type pigments is therefore nonstoichiometric with the incorporation of various cations into the lead pyrochlore structure \((\text{Pb}_{2-x}\text{M}'\text{M}_2\text{Y}_7\text{O}_{17-δ})\) with \(\text{M, M}' = \text{Sb, Sn, Fe, Si, Zn; M} = \text{RE}\). This phase is built with two sublattices, one of (big) Pb\(^{2+}\) cations, the second of (small, covalent-bonded) cations forming tetrahedral and octahedral entities sharing some oxygen atoms. During the firing process of the enamels, composition of the pigment used may be further affected with other species present in the enamel matrix (Fe, Si, Sn, etc.) [14,18,50,58,59]. Some of these incorporating elements may also have different speciations (e.g., Sn\(^{2+}\) or Sn\(^{4+}\), Sb\(^{3+}\) or Sb\(^{5+}\), Fe\(^{2+}\) or Fe\(^{3+}\)), and furthermore, the oxygen nonstoichiometry very much depends on the conditions of the firing process (reducing/oxidizing conditions).

In practice, the lead pyrochlore pigment can be prepared as a powder in advance and then mixed with the glass/glaze powder before deposition, the so-called “anime” and “corpo” recipes [57,60,61]. Alternatively, all the ingredients can be mixed together to obtain the yellow glass/glaze powder. In the first method, the grains of lead pyrochlore are rather big (>\(\mu\)m), while in the other, the homogeneous nucleation on cooling gives rise to small precipitates (<a few \(\mu\)m). With a high-magnification microscope objective, a single pigment grain could be analyzed, and the Raman spectrum obtained was strong without a background, and almost no signature of the glassy matrix was recorded (see e.g., Figure 7A—TH457 yellow; see also in ref. [34]). In the case of nucleation, both the signature of lead pyrochlore precipitates and glass were recorded together with a significant background (e.g., Figure 6D(c,c')).

In the last decades, several Raman spectrometry studies performed on various pottery glazes, glasses and enamels [1–3,11–18,47–52,57–59] as well as oil paintings [53–56] allowed for the identification of Naples yellow lead pyrochlore pigments. In general, these pigments have a very characteristic Raman signature, which basically includes the strong stretching mode peaks of Pb ions at the low-wavenumber region (~115–145 cm\(^{-1}\)) and M/M’–O stretching modes from ~300 to 600 cm\(^{-1}\). Furthermore, Sb-rich pyrochlore pigment is mainly distinguished by strong peaks at ~110–130 and ~505 cm\(^{-1}\), while Sn-rich-type pigment exhibits stronger peaks at ~130–135, ~335 and 450 cm\(^{-1}\) (the Sn–O stretching mode) with the disappearance of the ~505 cm\(^{-1}\) peak [10,13,18,47–52]. Zn-containing mixed lead pyrochlores exhibit a medium 450–480 cm\(^{-1}\) component [18]. Although Sn-rich lead pyrochlore was observed in most of the yellow and green enameled areas (Figure 6B(c) (R1048), Figure 6D(c,c') (R1135), Figure 9A(a) (R1006), Figure 9B(a,a') (SN284), Figure 10A(b,b') (TH487), Figure 11A(a) (R1056) and Figure 13B(c,c') (F1341C)), Sb-rich compositions were clearly observed in the yellow enamels of the TH457 bottle from the Thiers Collection (Figures 3 and 7A) and the R1175 plate with an English family’s coat of arms (Figures 2 and 8 A(a,a')). Mixed lead pyrochlores were also observed in some cases (Figure 7B(a,a') (F1371C) and Figure 12 B(b,b',b'') (F1429C)). As a second piece of proof, the XRF spectrum recorded on the yellow enamel of TH457 showed both Sn and Sb signals (Figures 14 and 15).
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Figure 14. Representative portable XRF (pXRF) spectra recorded on the TH457 bottle for the glaze and different painted enamels.

Figure 15. Representative pXRF spectra recorded on the TH457 bottle for pink and black enameled areas; bottom: zoom in on the energy window characteristics of Sn and Sb elements for different areas.
3.2.3. Red to Violet Colors

The last important characteristic of the painted enameled porcelains analyzed was the presence of different methods for obtaining red-related colors. The use of hematite or gold nanoparticles was identified in the painted enamels to obtain red, orange to pink/purple hues (Table 2). In only two artifacts, the TH457 bottle (Figures 3 and 7A, white-pink and red) and R1175 plates (Figures 2 and 8A(c,d), orange and red; and Figure 8B(c), red), the use of both methods was observed. Hematite has a well-defined Raman signature, with a strong (ca. 1305–1310 cm\(^{-1}\)) mode (resonance signal of magnon mode under green laser excitation) and narrow peaks between 200 and 600 cm\(^{-1}\) [62,63] (see e.g., Figure 6D(b,b',b''), Figure 7A, Figure 8A(c),B(c) and Figure 12C(b,b')). The magnon mode is very sensitive to the particle size of hematite [63] and the oxygen stoichiometry [62], with small particles giving an orange-red color [29]. Broadening of hematite peaks indicates a partial substitution of iron with other elements, possibly Ti and Al coming from the iron sources [29].

A specific Raman feature referred to the use of gold nanoparticles dispersed in the glassy matrix of the enamel (this type of enamel is at the origin of the label “Famille rose” [15,19], with a characteristic broad fluorescence background peaking at ~500–600 cm\(^{-1}\) under a green laser, corresponding to ~500 nm on the absolute scale [14,18]; see Figure 6C(c) (R1041), Figure 7A (TH457), Figure 8A(d) (R1175), Figure 9B(c) (SN284), Figure 10A(a,a'),B(d) (TH487), Figure 12C(a) (F1429C) and Figure 13C(a) (F1341C)). This method was first experimented with in 17th-century France by the glassmaker Bernard Perrot and then some years later by Johann Kunckel in Germany and certainly by others in Italy before [16,18,64–67]. The presence of gold was also confirmed by pXRF analysis (see arrow on Figure 15—pink, in which the small Au L\(_\alpha\) peak is obvious), with a hardly visible peak well-identified by data fitting. In addition, it is important to note that a very small band characteristic of the As–O stretching band at ca. 820 cm\(^{-1}\) could be detected along with the fluorescence background characteristic of the use of colloidal gold (seen clearly in Figure 10A(a,a') (TH487) and less intensely in Figure 9B(c) (SN284) and Figure 13C(a) (F1341C)). This band corresponds to the use of Perrot’s preparation route for obtaining the colloidal gold precipitate using an arsenic salt, which is different from that of Kunckel, who made use of a tin salt instead [16]. Similar Raman signatures were observed for painted Chinese enameled wares of the same period [14] as well as on French enameled watches [18].

3.3. Glassy Matrix

Raman spectra of the glazes/painted overglaze enamels showed the characteristic signature of glassy silicates, which basically include Si–O bending and stretching bands in the range ~500 and ~1000 cm\(^{-1}\), respectively. A comparison of the Si–O stretching band components and the area ratio of the Si–O bending/stretching band has proved to be very efficient for the compositional classification of different glass/glaze/enamel types based on experimental Raman data obtained for a great deal of ancient glassy objects [68–71] as well as on Density Functional Theory calculations [72]. The glazes were aluminosilicate glazes, with the Si–O stretching bands varying between ca. 900 and 1200 cm\(^{-1}\). The painted enamels analyzed displayed a broad and strong band varying between ~920 and 1070 cm\(^{-1}\) with various shoulders/components, indicating mainly lead-based compositions [69–71], as observed for soft-paste porcelains [17] and enameled metalware [14,18]. In some of the spectra, the intensity of the Si–O bending modes near 500 cm\(^{-1}\) appeared to be equal to or weaker than that of the stretching modes at 950–1050 cm\(^{-1}\) (see e.g., Figure 6A(c),C(a), Figure 10B(b) and Figure 13B(a,a')). This indicates a polymerization ratio close to less than 1, according to firing at temperatures inferior to 900–1000 °C [68,71].

The painted enamels can be further classified into three main glass types (Table 3) based on representative glass (and glaze) types [12,13,16,68,69] and previous studies conducted on Chinese cloisonné and painted enamels [10,14,73–76]: (i) lead-rich glass with the strongest component at ca. 920 to 1020 cm\(^{-1}\) (Type I); (ii) lead (earth) alkali glass with the strongest component at ca. 1030 to 1060 cm\(^{-1}\) (Type IIa), with rather similar-intensity components at ca. 980 and 1040 cm\(^{-1}\) (Type IIb); and (iii) lead (earth) alkali glass with the strongest component at ca. 1070 cm\(^{-1}\) (Type III). Some of the yellow, green and black enamels belonged to the lead-rich type of glass, with their strongest component
at low wavenumbers, which indicates a low degree of polymerization and firing at low temperatures (the addition of Naples yellow increases the lead content). Most of the painted enamels (white and blue) belonged to the lead alkali type of glass (the addition of smalt increases the potassium content), with their strongest component shifted to higher wavenumbers (up to ca. 1060 cm$^{-1}$) accompanied by a similar low wavenumber component to the lead-rich glass. The glass compositions of the painted enamels did not show much variety, indicating a limited number of firings over the transparent glaze.

Table 3. SiO$_4$ stretching components of the glassy matrix of the painted enamels analyzed (strongest; shoulder/peak) and comparison with Chinese enameled metalware (Lead alkali glass: L-a; Lead-rich glass: L-rich).

| Artifact | Si–O Stretching Components (cm$^{-1}$) | Color | Glass Type | Observed in Cloisonné Metalware [14] | Observed in Painted Metalware [14] |
|----------|----------------------------------------|-------|-------------|--------------------------------------|--------------------------------------|
| F1371C   | 1025, 1130                             | White, blue | L-a IIa      |                                      |                                      |
| R1048    | 980, 1040, 975, 1040                   | White, Blue | L-a IIa      | Blue and white                      | F1440C                               |
| R1056    | 980, 1035                              | White    | L-a IIa      |                                      |                                      |
| TH457    | 985, 1130, 985, 1070                   | White pink, Yellow | L-rich I | Yellow and black                     | Yellow and turquoise yellow and green |
| R1175    | 985, 1040, 975                         | Blue     | L-a IIa      | See above                           | Green                                |
|          | 905, 970, 1035, 1120                   | Yellow   | L-rich I     | See above                           |                                      |
|          |                                        | Black    | L-a IIb      |                                      |                                      |
|          |                                        | Green, orange, red | L-a IIa |                                      |                                      |
| TH487    | 975, 1050, 970, 1045                   | Blue     | L-a IIa      | See above                           |                                      |
|          | 920, 1140                              | Yellow   | L-a IIa      |                                      |                                      |
|          | 915, 971, 1037                         | Black    | L-rich I     |                                      |                                      |
| R1045    | 975, 1035                              | Blue, white | L-a IIb |                                      |                                      |
| R1041    | 980, 1050                              | Blue, pink | L-a IIa |                                      |                                      |
| R1006    | 1015, 1020                             | Yellow   | L-rich I     |                                      |                                      |
|          | 1020                                   | Black    | L-rich I     |                                      |                                      |
| SN284    | 980, 1060                              | Yellow, red, pink | L-a IIa |                                      |                                      |
| R1135    | 1020                                   | Green, blue, red | L-rich I |                                      |                                      |
| F1341C   | 970, 1035, 980, 1030                   | White, Blue | L-a IIb |                                      |                                      |
|          | 1040, 1090                             | Green    | L-a IIb      |                                      |                                      |
| F1429C   | 980, 1030, 1140                        | White, blue | L-a IIa |                                      |                                      |
|          | 970, 1020                              | Light pink | L-a IIa |                                      |                                      |
|          | 900, 975, 1010, 1130                   | Yellow   | L-rich I     |                                      |                                      |
|          | 900, 975, 1010, 1125                   | Green    | L-a IIb      |                                      |                                      |

3.4. XRF Analysis

One of the porcelain artifacts, the TH457 bottle, was analyzed by pXRF. This artifact exhibits the most sophisticated decor, and its shape is well-suited for XRF analysis using a portable instrument since the analyzed area must be roughly perpendicular to the instrument [18]. Additionally, the distance between the instrument and the artifact must be small (~10 mm), and flat areas are required. Figures 14 and 15 show the representative XRF spectra recorded on the white glaze and the different
overglaze enamels (white, blue, green, red, yellow, pink and black) in the 1–22 keV range, where most of the peaks characteristic of major and minor elements were observed. Figure 15 shows details of the 24–30 keV range, where peaks of tin and antimony elements could be compared visually.

As expected for porcelains, the glaze showed elements characteristic of a glassy aluminosilicate porcelain glaze fired at a high temperature [1–3,28–31] with high levels of potassium, calcium and silicon (Table 4). Significant iron content was also obvious in the (colorless) glaze. The high whiteness of the glaze, in association with some iron impurities, demonstrated a firing process under a reducing atmosphere that imposed Fe$^{2+}$ speciation. These ions lead to a weak blue coloration, reinforcing the “whiteness” of the glaze for human eyes. Strontium, an impurity of calcium, was detected at a significant level. Light elements such as sodium could not be detected due to their low atomic number. All overglaze enamels were lead-rich: all the L$_{\alpha}$, L$_{\alpha}$XRF transitions of the lead element were clearly observed. A comparison between the relative intensity of lead and potassium or calcium peaks indicated the lower content of lead in the black enamels.

Table 4. The glaze and overglaze painted enamels analyzed by pXRF. (Major-minor-traces).

| Glaze          | Elements                  |
|---------------|---------------------------|
| White         | Si, K, Ca, Fe–Sr          |
| Overglaze enamel |                      |
| Blue          | Si, Pb–K, Co, Fe, As–Ni, Mn |
| White         | Si, Pb–K, Fe, Ca, As–Ni   |
| Green         | Si, Pb–Cu, Fe, K, Ca, Sn, Zn–Mn, Ni, (Co?) |
| Yellow        | Si, Pb–K, Sb, Fe, Sn–Ni   |
| Red           | Si, Fe, Pb–K, Ca, Ti–Mn   |
| Pink          | Si, Pb–K, Ca, Fe, As–Au, Ni |
| Black         | Si, K, Ca, Pb, Fe–Mn, Co, Cu–Ni, Zn |

The XRF spectrum of the blue overglaze enamel showed strong lead peaks; pronounced arsenic, iron and cobalt peaks; and weak manganese and nickel peaks. Arsenic was also observed in the white overglaze enamel (flower decor) as well as in the pink one. Arsenic content is visually detected by the small As K$_{\beta}$ peak just before the Pb L$_{\alpha}$ one, the As K$_{\alpha}$ peak being at the same position as the strong Pb L$_{\alpha}$ one. Confirmation was carried out with the software fitting. Note the high intensity of the potassium peak in blue, according to the use of smalt. The presence of nickel and manganese was due to the impurities found in the cobalt ore used (Table 4). The intensity of the iron peak was rather similar for all overglaze enamels. The presence of manganese was thus consistent with the implication that Asian and European cobalt ingredients were mixed. The detection of arsenic was also significant in relation to the type of the cobalt ore. The red overglaze enamel displayed significant iron peaks, in accordance with the Raman identification of hematite (Figure 7A). Minor amounts of titanium and traces of manganese were also observed, possibly coming from the hematite source. The pink overglaze enamel displayed minor amounts of iron, also indicating the use of hematite. In addition to that, the detection of gold traces was significant, confirming the use of colloidal gold for obtaining the desired hue. The presence of arsenic was also in accordance with the use of Perrot’s technique.

The yellow overglaze enamel showed distinct peaks of iron together with minor amounts of antimony and tin, indicating the use of a complex (Fe) Sb-rich Naples yellow lead pyrochlore pigment (see Section 3.2). A similar type of this pigment also seemed to have been used for the green overglaze enamel, with a higher amount of tin and some zinc. Antimony was detected in the low-energy range, but a magnification of the 24–30 keV range (Figure 15) showed the variable content of tin and antimony elements more clearly. The green enamel additionally included copper, which contributes to color formation, together with manganese, nickel and zinc. It was difficult to see if cobalt was also present.
Iron was clearly observed in the black overglaze enameled areas (Table 4). Black enamel was obviously obtained with a mixture of iron and manganese in the presence of copper (and cobalt?), according to the Raman identification of manganese-rich oxide (spinels) (see e.g., Figure 6B(dd′) (R1048), Figure 8A(e) (R1175), Figure 9A(e,e′,e″) (R1006) and Figure 10B(c,c″) (TH487)). It is important to note that relative intensities of lead and silicon elements measured on the black lines were intermediate between those measured on other overglaze enamels and that of the glaze. This is due to the limited thickness of the black lines: the XRF spectrum (Figure 15) both displayed the contribution of the black line plus those of adjacent areas, the glaze on one side and the colored overglaze on the other side. All the data obtained by pXRF were in perfect agreement with the Raman data.

4. Comparison with Painted Enameled Metalware

The characteristics of porcelain and metal enameling were expected to be very much linked to each other in terms of the pigments/opacifiers used and the firing conditions employed. At the end of the 17th and early 18th century in China, enameling workshops both in charge of metal and porcelain enameling were established in the Forbidden City by the Jesuit mission to satisfy the demand of the Kangxi Emperor [4–8]. These workshops are considered to have been in close contact with each other [27]. The present study has shown that the material characteristics and production technology of 18th-century Chinese painted enameled porcelains and the painted enameled metalware of the same period are very similar. Pigments/opacifiers of European origin, such as Naples yellow lead pyrochlore pigment (yellow to green enamels), colloidal gold (red, orange and pink enamels) and lead arsenate phases (white and blue enamels) as well as similar lead-based glass compositions were identified in both of the enameled ware types. However, with the present state of knowledge, some differences can be noted on the diffusion of these new technologies in China. Tables 5 and 6 chronologically compare the appearance of European technologies for blue, white, yellow/green and red/pink enamels of porcelain and metalware, respectively.

| Period (Assignment Based on Decor Style) | Porcelain | Specific Technologies | Remarks | Refs |
|----------------------------------------|-----------|----------------------|---------|-----|
| Final period of Kangxi reign (<1722)   | G1710 vase | Ming blue H-red Sn-N.y.(s) | “Biyu tang zhi” mark Famille verte/Famille noire J.p.k. | [12] |
|                                        | G822 dish  | Sn-N.y. (s) H-red     | “Da Qing Kangxi nian zhi” mark Famille verte J.I.F. | [12] |
|                                        | G5696 bowl | Ming blue H-red Sn5b-N.y.(s) | “Da Qing Kangxi nian zhi” mark doucai J.I.F. | [12] |
|                                        | G5250 bowl | As-blue; Ming blue Sb-N.y.; Sn-N.y. Au° Cassiterite? | “Kangxi yu zhi” mark huafalang Pw. | [12,17] |
|                                        | G3361 water dropper | As-blue As-white Sn-N.y. Au° | “Da Qing Kangxi nian zhi” mark Famille rose J.I.F. | [12] |
| Period (Assignment Based on Decor Style) | Porcelain | Specific Technologies | Remarks | Refs |
|----------------------------------------|-----------|-----------------------|---------|------|
| **Yongzheng reign (1723–1735)**       | R1006 cup | *Ming blue* Sn-N.y.   | This work |      |
|                                        | R1056 dish| As-white Sn-N.y.      | This work |      |
|                                        | R1041 cup | As-blue Au°           | This work |      |
|                                        | MG4806 bowl | *Ming blue* Sb-N.y. (s) | “Yongzheng yu zhi” mark huafalang/jiangcai J.I.F. | [12] |
|                                        | MG913 bowl | As-blue Sn-N.y. (s) H-red | “Yongzheng yu zhi” mark huafalang/jiangcai J.I.F. (?) | [12] |
|                                        | MG7368 bowl | *Ming blue* N.y. (s) H-red | “Da Qing Yongzheng nian zhi” mark, doucai J.p.k. | [12] |
| **Qianlong reign (>1735–1750)**       | R1175 dish | As-blue (A) CaF (?), Au° Sn/Sb-N.y., H-red | This work |      |
|                                        | TH487 dish | As-blue Sn/Sb-N.y. Au° | This work |      |
|                                        | F1341C bottle | As-blue Sn-N.y. Au°; H-red | This work |      |
|                                        | SN284 milk pot | As-blue (A) Sn-N.y. Au° | This work |      |
|                                        | SN284 cup | Sn-N.y. Au°           | This work |      |
|                                        | Shard bowl | As-blue As-white H-red Sn-N.y. (yellow) Sb-N.y. (green) | From Palace excavation | [13] |
| **Qianlong reign (>1735–1750)**       | R1045 cup | As-blue               | This work |      |
|                                        | FI429C teapot | As-blue(A) Sb-Sn-Y.y. Au° | This work |      |
|                                        | TH457 bottle | *Ming blue* Sn-N.y. Sb-Sn-N.y. Au° H-red | This work |      |
|                                        | F1371C bottle | As-white As-blue (A) Sn-N.y. | This work |      |
|                                        | R1025 dish | As-blue (A) Au°      | This work |      |
|                                        | R1048 cup | As-white, As-blue Sn-N.y., H-red | This work |      |
Table 5. Cont.

| Period (Assignment Based on Decor Style) | Porcelain       | Specific Technologies | Remarks       | Refs |
|-----------------------------------------|-----------------|-----------------------|---------------|------|
| Qianlong reign (1750–1800)              | MG3668 teapot   | As-blue, Au\(^+\) As-white Sn-Sb-N.y. | Yixing p.k.   | [12] |
| MG9604 teapot                           | As-blue As-white Sn-N.y. | | Yixing p.k. | [12,17] |

P.w.: palace workshop; J.I.F.: Jingdezhen Imperial Factory; J.p.k.: Jingdezhen private kiln; As-blue: European cobalt; Sb-N.y.: Sb-rich Naples yellow; Sn-N.y.: Sn-rich Naples yellow; s: small intensity; ancient Ming technology: Asian blue; hematite red; MGxxxx: Musée national des arts asiatiques—Guimet inventory number.

Table 6. Chronological summary of the innovative technologies evidenced in Chinese enameled metalware [14].

| Period                          | Painted | Specific Technologies | Cloisonné | Specific Technologies | Remarks                   |
|---------------------------------|---------|-----------------------|-----------|-----------------------|---------------------------|
| Final period of Kangxi reign (<1722) |         |                       | F1448C    | Sb-Sn-N.y. Sn-N.y. Cassiterite Au\(^+\) | Palace workshop? |
| Qianlong reign (1735-1750)      | R957    | As-white               | As-blue   | Sn-N.y.               | Guangzhou                |
| Qianlong reign (1750-1775)      | F1440C  | As-blue (A)            |          | Sn-N.y.               | Qianlong mark            |
|                                 |         |                       | F1735C    | Sb-Sn-N.y. Sn-N.y.    |                           |
|                                 | R958    | As-white               | As-blue (A) | Sn-N.y.     | Guangzhou                |
|                                 |         |                       |          | Au\(^+\)               |                           |
|                                 | R975    | As-blue                | Sb-Sn-N.y. | Au\(^+\)   | Guangzhou                |
|                                 |         |                       |          |                       |                           |
|                                 | F1501   | As-blue                | Sb-Sn-N.y. Sn-N.y. Au\(^+\) | Palace workshop |
|                                 |         |                       |          |                       |                           |
|                                 | F1467.1/2 | Sb-N.y.               | Sn-N.y.   |                        | Qianlong mark            |
|                                 |         |                       |          |                        |                           |
|                                 | F1698C  | As-blue                | Sn-N.y.   |                        | Guangzhou                |

Regarding the blue overglaze enamels studied so far, the change from Ming blue (made with Asian cobalt ores) to As-rich blue (pure European cobalt ores or mixing with Asian ones) is suggested
for the artifacts assigned to the end of the Kangxi reign on the basis of the decor style. It seems that both types of blue were used in the MG5250 “Imperial” bowl [12]. This bowl also exhibited the Raman spectrum characteristic of the use of colloidal-gold-based (Famille rose) enamel. Naples yellow pyrochlore, tin-rich, was the first European technology imported, but the addition of antimony was expected in the MG5696 “Imperial” bowl [12]. A weak signal of cassiterite was even detected in a similar shard [17]. It can be also noted that the MG5250 bowl is assigned to have been made at the palace workshop; the same innovative technology of enameling was observed for the MG3361 pencil water dropper assigned to have been made at the Jingdezhen Imperial Factory [12]. This indicates a very rapid transfer of technology from the palace workshop managed under the Jesuits’ guidance to the imperial kilns at Jingdezhen. The Sn-rich Naples yellow (in yellow and green enamels) appears to have been used more frequently than the Sb-rich homologue up to the mid-18th century. The addition of Naples yellow increased the lead content. The use of cassiterite seems be exceptional, and its use must be confirmed by an examination of shard sections of similar artifacts.

It is worth mentioning that cassiterite was identified in the painted enamels of the metalware, as an opacifier, for a Kangxi cloisonné metalware piece and the exceptional painted and cloisonné Qianlong ewers from the French emperor’s collection. Regarding enameled porcelain, Sn- and Sb-rich Naples yellow pyrochlorites appear to have been used since the final period of the Kangxi reign. The same conclusions can be drawn for the use of colloidal gold for red to pink colors. By comparison, it appears that almost all enameled metalware was prepared with the European technologies, including As-rich cobalt, although enamel traditional recipes such as Ming blue were used simultaneously with new ones (Ming blue was used for underglaze decor). More artifacts must be analyzed to confirm this point.

5. Conclusions

The present on-site Raman study of intact Chinese painted enameled porcelains from the 18th century sheds light on the materials (pigments/opacifiers, colorants, glass types of glazes/enamels and body phases) and the technology used during their production. The study also provides a comparative approach for the technical aspects of painted enameled porcelains and contemporaneous enameled metalware regarding the different reigns of the Qing dynasty. The use of the same on-site analytical protocol for these artifact groups is advantageous for comparing the data obtained. The outcomes are significant in terms of the know-how transfer (materials/techniques) from Europe to China by the Jesuit mission at the end of the 17th century.

From the (limited) number of objects studied, it appears that the first innovations were the different types of Naples yellow lead pyrochlore pigments and then the use of European cobalt during the later years of the Kangxi reign for enameled porcelains. Cassiterite appeared first during the same period in certain cloisonné enamel wares. The detection of cassiterite traces in a Kangxi enamel is assigned to (the effect of) the use of Sn-rich Naples yellow that induced precipitation of cassiterite. Its addition to adjust the hue is also possible. More artifact studies are needed to clarify this point. Yongzheng enameled objects made use of European recipes for blue, yellow and rose; the opacification with different As phases was identified. Unfortunately, only one metalware piece from the Yongzheng period has been studied [14]. Studies of a larger number of objects are required, in particular to study the technology transfer from the Imperial Palace to the Jingdezhen imperial kilns and to private kilns in Jingdezhen or Guangzhou [27] to really have a statistical view of the number of porcelains on which the lead-based enamels were deposited directly on the body and not on the glazed body [77].

Author Contributions: Conceptualization, P.C. and B.Z.; methodology, P.C.; validation, P.C. and B.K.; investigation, P.C.; resources, B.Z., J.-B.C., Y.Y. and V.D.; writing—original draft preparation, P.C.; writing—review and editing, P.C. and B.K.; funding acquisition, P.C. and B.Z. All authors have read and agreed to the published version of the manuscript.

Funding: This research was funded by the French Agence Nationale de la Recherche ANR EnamelFC project—19-CE27-0019-02 and by the Partenariat ‘Hubert Curien’ PHC-Cai Yuanpei Campus France–China cooperative program.
Acknowledgments: M. Gironda from ELIO, XGLab/Bruker is kindly acknowledged for recording the XRF data. C. Gougeon from the Louvre Museum and S. Paronetto from Fontainebleau Castle are acknowledged for their help in the preparation of the measurement campaigns. J. Defretin is acknowledged for providing historical information on some artifacts. Yong Lei and Huawen Liu from the Palace Museum, Department of Analysis and Conservation and Professor Ludovic Bellot-Gurlet from MONARIS (Laboratoire de la MOlécule au Nano-objet: Réactivité, Interaction & Spectroscopies) are kindly acknowledged for many discussions. This work has been conducted under the framework of the Laboratoire International Associé CNRS–Palace Museum TrEnamel and Campus France PHC-Cai Yuanpei programs.

Conflicts of Interest: The authors declare no conflict of interest.

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