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Article

Investigation towards Laser Cleaning of Corrosion Products from Lead Objects

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Abstract: In this work lasers with micro-, nano-, pico- and femtosecond pulse durations were used to clean atmospheric corrosion products from the fragments of a 19th-century lead outdoor sculpture. The state of the surface was studied by optical microscopy, Raman spectroscopy, X-ray photoelectron spectroscopy and scanning electron microscopy. It was shown that for all lasers used there is no self-limiting cleaning effect, and the metal damage threshold is lower than the corrosion removal threshold. Using the XPS method, it has been demonstrated that the effect of turning a metallic lead surface blue after irradiation is associated with an interference effect in the PbO film. Raman spectroscopy indicated no phase changes in the corrosion layer after laser cleaning with 8 ns, 75 ps and 100 fs pulses, which makes these lasers useful for the layer-by-layer cleaning of archaeological objects.

Keywords: laser cleaning; atmospheric corrosion; lead; Raman spectroscopy; SEM



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1. Introduction

Lead is one of the first metals that became known to humans and many museums around the world store lead objects in their collections. Lead is quite well preserved in atmospheric conditions as it forms a thin oxide film which protects the base metal from any damaging influence of the environment. The main cause of lead corrosion is the presence of humidity condensed on its surface. Water serves as an electrolyte and provides a medium for dissolving the entire spectrum of the atmosphere's gases [1,2]. By entering a condensed water drop, carbon dioxide leads to the formation of lead carbonates, primarily cerussite (PbCO₃) and hydrocerussite (Pb₃(CO₃)₂(OH)₂). Upon further exposure to the atmosphere, carbonates react with sulfur dioxide (SO₂) and sulfur trioxide (SO₃) and form a film, which consists of a mixture of scotlandite (PbSO₃) and anglesite (PbSO₄). Subsequently, the main part of scotlandite is displaced, forming a protective film of anglesite [2,3]. The build-up of corrosion products eventually leads to the cracking of the surface, thereby increasing the access of contaminants to the underlying metal [4]. Atmospheric composition affects both the relative amounts of corrosion products formed and the order of their formation [5].

In the case of museum storage, volatile organic acids, mostly acetic and formic, can initiate and/or catalyze the corrosion process [6,7]. These acids come from oak or coniferous wood, used in the manufacturing of museum showcases [8]. In the presence of acetic acid vapors, lead forms lead acetate, which then turns into basic carbonate-cerussite under the action of atmospheric carbon dioxide. This process is accompanied by a further release of acid, which in turn causes a cyclic corrosion reaction of the metal up to its complete destruction. In the case of organs, the use of oak wood in their construction led to such a damaging corrosion processes that, in order to preserve organ heritage, the research project COLLAPSE (Corrosion of Lead and Lead-Tin Alloys of Organ Pipes in Europe) was realized [1,9,10].

Lead objects in museums need regular monitoring, stabilization and cleaning to extend their lifetime. Corrosion processes weaken the metal, making it brittle and sensitive to mechanical stress. Metallic lead is softer than its corrosion products: according to the Mohs scale of mineral hardness, the hardness of lead is 1.5, while the hardness of laurionite (Pb(OH)Cl), anglesite (PbSO₄) and cerussite (PbCO₃) is in the range of 2.5–3.5 [11]. Abrasive or harsh manual cleaning can cause a loss of an object's fragments and leave tool marks on the surface. Even cleaning with compressed air or a pressure washer can damage soft metal [12]. With regard to the conservator, the mechanical cleaning of lead objects can be hazardous, given that lead dust is poisonous [1]. In view of all the above factors, the mechanical cleaning of lead is a rather complicated restoration procedure.

In the case of chemical cleaning, the reagents are difficult to remove completely from an object. As a result, traces of chemical agents activate processes of metal destruction [1]. Electrochemical cleaning has been successfully applied to small lead objects [7]. However, the method is quite time-consuming [13] and requires the full immersion of the object in a solution, which is impossible for monumental sculptures. An additional constraint is that only materials that have a non-aggressive nature in relation to an object, and are also safe both to a restorer as well as the environment, are recommended for restoration procedures [1].

Shortcomings of the current suite of methods dictate the need to investigate alternative techniques. The application of laser cleaning for heritage objects is very promising, since the effect of radiation is strictly localized, well controlled and can be immediately stopped. Radiation parameters can be selected individually, there is no mechanical pressure on the treated surface and this is a chemical-free technique.

Only a few papers on the laser cleaning of heritage lead objects have been published, although lead was one of the first metals that laser cleaning was applied to [14]. In the pioneering work of John Asmus in the early 1970s, a ruby laser was used to clear a section of lead strip from a XII century sarcophagus. In the work [12], a Phoenix Nd:YAG laser was used to clean a gilded lead statue, operating in a Q-switched mode (pulse duration of about 10 ns) at a wavelength of 1064 nm, which showed high efficiency. The authors claimed that the achieved results demonstrate a good level of preservation of the lead monument. In another study [15], the laser cleaning of lead coins from calcareous encrustations was carried out. The authors recommend operating with a laser fluence of 0.4–1 J/cm² and a pulse frequency of less than 10 Hz for 2–10 ns, using a 1064 nm Nd:YAG laser. None of the three papers contain any surface examination, either with optical microscopy or SEM.

The aim of the present work is to study the processes that occur during the interaction of laser pulses of various durations with lead and its corrosion products in order to create a guideline for conservators. Laser cleaning of museum objects is rather demanding, and must meet a number of requirements. Firstly, laser irradiation should not create new artificial colors on the surface, since this changes the historical authenticity of the object. Secondly, the laser must not damage the bulk metal during corrosion removal, since this interferes with the original surface [16,17] and can activate a corrosion process. In this work a number of laser systems with different pulse lengths were used to find the proper cleaning regime and to detect a potential self-limiting cleaning effect.

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2. Materials and Methods

3.4. Materials

Fragments of an outdoor statue of an angel by sculptor I. P. Prokofiev, dated 1823 from the dome of Saint Catherine Church of Vasilyevsky Island in Saint Petersburg, were used from the dome of Saint Catherine Church of Vasilyevsky Island in Saint Petersburg, were used as experimental samples. The 3-8 meter high sculpture was made of wood and covered with 2-mm thick lead plates (Figure 1). Due to the destruction of both the wooden base and tred with 2-mm thick lead plates (Figure 1) but to the destruction of both the wooden base and the lead sheets, the sculpture was dismartled and replaced with a copy. Our experiments base and the lead sheets, the sculpture was dismartled and replaced with a copy. Our experiments were carried out on small fragments of the original lead sheet.



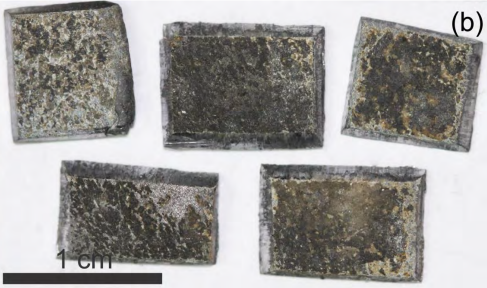


Figure 1: (a) Sculpture of an angel, showing intense corrosion features (photo provided by the figure 1: td. restoration workshop). (b) Macro-photo of lead plate fragments.

Nasledie Ltd. restoration workshop). (b) Macro-photo of lead plate fragments.

2.2. Analytical Methods and Laser Systems

Optical microscopy was used to observe the surface dranges ((color, tapography)) after irradiation. Wisual appearance is in the national important parameter to must use object bleet, because to prepare the love ging late that content that its extragent for the manuscripe color of the perfect medical single Crist States is 160m Hiose H

The color change of the irradiated surface occurred for multiple neasons, such as the formation of a new substance, a change in the crystall's size or a phase modification of initial corrosion products. To analyze the phase composition of comosion deposits Raman spectroscopy was used. Raman spectra were cobtained with a Schatter (B(BlackeBilleSiA), MA, USA) Raman spectrometer, conjugated with an Olympus BX-51 microscope, using a 785 nm solid state laser. Due to a relatively strong luminescence, the obtained spectra were baseline corrected and normalized. X-ray photoelectron spectroscopy (XPS) was used to analyze the thin top-most surface layer on the irradiated surface. The XPS measurements were conducted with an Escalab 250Xi (Thermo Fisher Scientific, Waltham, MA, USA) photoelectron spectrometer with Al-Kα radiation (photon energy 1486.6 eV). Spectra were recorded in the constant pass energy mode at 50 eV, using XPS spot size 650 μm. Peaks were fitted with the product of asymmetric Gaussian and Lorentzian line shapes. The surface micro-topography and elemental composition analysis was performed using a TESCAN

solid state laser. Due to a relatively strong luminescence, the obtained spectra were baseline corrected and normalized. X-ray photoelectron spectroscopy (XPS) was used to analyze the thin top-most surface layer on the irradiated surface. The XPS measurements were conducted with an Escalab 250Xi (Thermo Fisher Scientific, UK) photoelectron spectrometer with Al-K α radiation (photon energy 1486.6 eV). Spectra were recorded in the constant pass energy mode at 50 eV, using XPS spot size 650 μ m. Peaks were fitted with the product of asymmetric Gaussian and Lorentzian line shapes. The surface micro-topography and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed using a TESCAN LYRA3 micro-lyra and elemental composition analysis was performed usin

The lasers used in the experiments varied in pulse duration, and are littled in ducreasing order in Fahl Table 1.

Table 1. Technical characteristics of the lasers used. **Table 1.** Technical characteristics of the lasers used.

Laser System V	Wavelen g VI	ą _{/viell} ęng nm	th, Pulse Duration Duration
EOS Combo (El. En., Firenze, Italy)	1064	1064	39011945 (FRPR) 100 ps (LQS)
SpitLight 2000 (Innolas, Krailling, Germany) Diode-pumpput Nasy (Innolases, Section 1)	1064 1064	1064	76-8 ns
Discherpromagned Lind Salva Glassers (Lassells users) Optical	Sys- 532	1064	6 ₹6 \$ps
Ti:Sa laser (Av esta J.Co. Mots Rus Site sia)	800	532	1003 fsps
Ti:Sa laser (Avesta Ltd., Russia)		800	100 fs

3. Results

331 Resmittes Characterization

3.1. Samplese Signaturi of tion residence includes the cross-section preparation, followed by SEMr FDX signature consultation of the section of the section

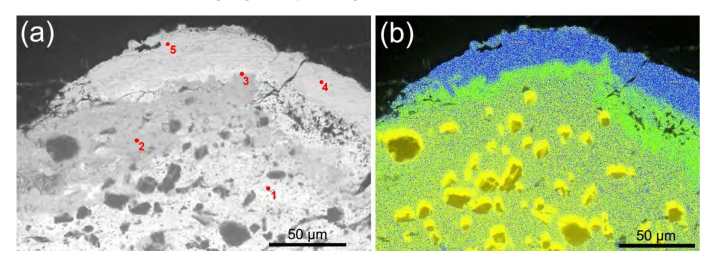


Figure 2. (a) SEM image of the cross-section, numbers correspond to analysis listed in Table 2. (b) SEM-EDX map of the same area, showing the distribution of chlorine (green), sulfur (blue) and silicon (yellow).

Table 2. Chemical composition (wt. %) of the lead alteration products.

Element/№	1	2	3	4	5
0	7.17	7.08	7.34	17.26	17.74
S	n.d.*	n.d.	0.20	7.71	8.04
C1	n.d.	1.18	1.32	0.92	0.34
Pb	92.83	91.74	91.13	74.1	73.88

* n.d.—not detected.

As it can be seen from Figure 1b, corrosion layers differ in color. Each of the observed layers was studied by Raman spectroscopy (Figure 3). The top layer has shades from brown to black. A typical spectrum from this area is shown in Figure 3a. Its characteristic feature is the presence of an intense peak with a maximum at 976 cm⁻¹, which is interpreted as the peak v_1 of the totally symmetric vibration in SO_4^{2-} in anglesite [18–20]. Additional confirmation of the anglesite presence is the detection of bands, corresponding to vibrations v_2 (at 445 cm⁻¹), v_3 (1155 cm⁻¹) and v_4 (608 and 640 cm⁻¹) [18,21]. The Figure 3a spectrum also shows peaks at 272 and 327 cm $^{-1}$. Their intensities change concurrently but independently of anglesite peaks and, as the Figure 3b spectrum shows, they become dominant in areas with the white color. This pair of peaks is consistent with the chlorine-containing mineral laurionite. For this mineral, crystallizing with the orthorhombic space group P_{nma} , the peak at $327\,\mathrm{cm^{-1}}$ was assigned to phonons with the symmetry of the irreducible representation of A_g and B_{2g} , and the peak at 272 cm⁻¹ to B_1 and B_{3g} [22]. Additionally, the spectrum Figure 3b shows peaks at 976 and 1050 cm⁻¹. The 976 cm⁻¹ peak refers to anglesite, while the 1050 cm⁻¹ peak is interpreted as the totally symmetric vibration of v_1 in the CO_3^{2-} ion in hydrocerussite or cerussite [23]. Overall, comparison of the spectra clearly indicates

Heritage 2023, 6, FOR PEER REWIEW the white area (Figure 3b) contains significantly less anglesite than laurionite as compared with the brown area (Figure 3a). The presence of the chlorine-containing mineral laurionite in the white region, as well as small amounts of hydrocerussite, is consistent with SEM-EDX data.



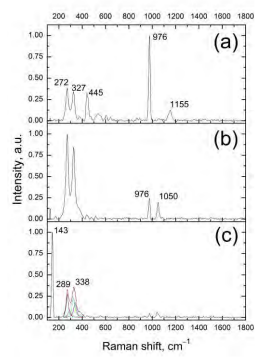


Figure 3. Macro-photo of the cross-section (left). The Raman spectra of different corrosion layers (right) is the figure 3. Macro-photo of the cross-section (left). The Raman spectra of different corrosion layers (right): black-brown (a); white (b); and dark, adjacent to metallic lead (c).

The third area is a visually dark layer adjacent to the metal (Figure 3c). Its spectrum has an intense peak with a maximum at 143 cm⁻¹, which is close to the characteristic peak

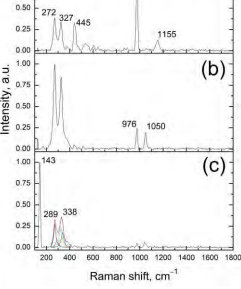
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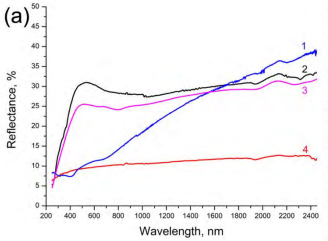
Spectral reflection coefficients we The reflectance spectra for the origin yellow (3) and black color (4) are pre surface, one of the fragments was im Tetra Acetic, C₁₀H₁₄N₂Na₂O_{8·2}H₂C years of outdoor exposure, the sculpti which is hidden by corrosion deposit



10 cm⁻¹ [uations 289 and rge [23]. surface. white (2), al metal Diamine lundred texture, e, where

corroded metal has gained a spongy structure. The curve №1 in Figure 4a corresponds to Figure 3. Macro-photo of the cross-section (left). The Raman spectra of different corrosion layers the reflectance spectrum of this surface. (right): black-brown (a); white (b); and dark, adjacent to metallic lead (c).

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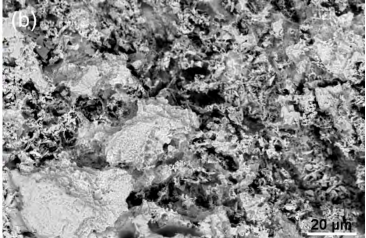


Figure 4. (a) Reflection spectra of various surface areas: chemically cleaned lead (1); white (2), brown **Figure 4.** (a) Reflection spectra of various surface areas: chemically cleaned lead (1); white (2), brown (3) and black (4) corrosion layers. (b) SEM image of the chemically cleaned lead surface. (3) and black (4) corrosion layers. (b) SEM image of the chemically cleaned lead surface.

As can be seen from the graphs of the first 4.), the layers have low reflection in the UNe region of the white (A) and 2 rayer by the control of the metal of the period o

With a multi-analytical approach to the sample study, the following sequence is obtained: a visually dark oxide layer (PbO, litharge) with the inclusion of laurionite (PbClOH) and hydrocerussite (Pb $_3$ (CO $_3$) $_2$ (OH) $_2$) is located closest to the lead. We consider that this layer contains the original surface layer, and so therefore with laser cleaning we need to stop at this depth. Moving outwards away from the metal layer, the dark oxide layer is followed by a white layer, which becomes thicker with a gradual increase in anglesite (PbSO $_4$) content. Furthermore, as the percentage of anglesite increases, laurionite and hydrocerussite decrease. Areas with the highest content of anglesite have brown and black shades. Previously [5,25], the sequence of formation of various lead compounds has been studied, and even though different authors gave different orders of phase formation, the

final substance in all works was anglesite. In our case, the marine air of Saint Petersburg led to the appearance of laurionite in this chain, as a result of the reaction of sodium chloride with lead oxide, as reported in [1,2]. Due to the outdoor exposure, the topmost layer of contamination contains soot/dust deposits.

3.2. Laser Cleaning Experiments

The laser cleaning experiments were carried out according to the following procedure: first, the laser-induced damage threshold of a chemically cleaned lead sample irradiated by a single pulse was determined. SEM was used for the precise detection of this damage threshold. Secondly, the corroded samples were irradiated with respect to the damage threshold of the metal. In the case of under-cleaning, firstly the number of pulses was increased. If the complete removal of corrosion products still could not be achieved, then the laser fluence was gradually increased. Irradiation was carried out in dry conditions and with the addition of water/isopropyl alcohol by wet brush. A portable ventilation system was used in all trials, since the by-products generated by the laser irradiation of lead are poisonous.

Although it is widely known that micro- and nanosecond pulses lead to sufficient heating of the surface, no negative consequences for these durations have been reported in published papers [12,14,15]. From the authors' own experience and the published review on laser cleaning of metals [26], it follows that even similar museum objects can behave differently under the same irradiation parameters. The slight difference in composition accounts for the differences in laser absorption and removal of the corrosion layers. Therefore, irradiation with μ s and ns pulses was performed.

3.2.1. Microsecond Pulses

Experiments began with EOS Combo laser irradiation of the chemically cleaned sample in free-running mode ($\tau_{pulse} = 30\text{--}110~\mu s$) with a pulse energy of 100 mJ. By gradually reducing the laser spot from 10 to 4 mm, we obtained several steps of fluence from 0.12 to 0.80 J/cm². The effect of the laser irradiation was visible to the naked eye starting from 0.35 J/cm². However, the SEM study showed that already at fluence above 0.19 J/cm², molten metal globules were formed on the lead surface (Figure 5a).

A corroded sample was then irradiated: at a fluence of 0.12–0.19 J/cm² no visible cleaning effect was observed even after 100 pulses. The addition of water or isopropyl alcohol did not increase the cleaning efficiency. There was no cleaning at the fluence of 0.35–0.80 J/cm², however black globules appeared on top of the corrosion layer (Figure 5b,c). Raman spectroscopy has shown that the globules' composition is very heterogeneous, and three main groups can be distinguished. The first group consists of synthetic analogues of litharge with a relatively large contribution of anglesite and a small amount of amorphous carbon. The second group is qualitatively similar to the first one, except there are fewer sulfates, and a larger proportion of the amorphous phase. In the third group, globules consist almost entirely of massicot (orthorhombic PbO), with occasional litharge in small quantities.

In an attempt to complete cleaning of the lead surface from corrosion layers, the laser fluence was increased. At 0.8 J/cm², a slight removal occurred. The number of pulses was increased to 100, but this did not lead to a complete cleaning of the metal. With a fluence of 3.2 J/cm², all deposits were removed after 3–4 pulses; however, the surface became heavily melted. SEM image demonstrates that while lead melts, the contaminants partially integrate into melted metal (Figure 5d). The surface, weakened by the corrosion processes, develops cracks and cavities which fill up with the corrosion products. Heating, produced by high energy irradiation, seals these cavities with corrosion products inside. This process is highly undesirable in the cleaning of heritage objects. The threshold for removing corrosion deposits in this mode is noticeably higher than the threshold for lead damage.

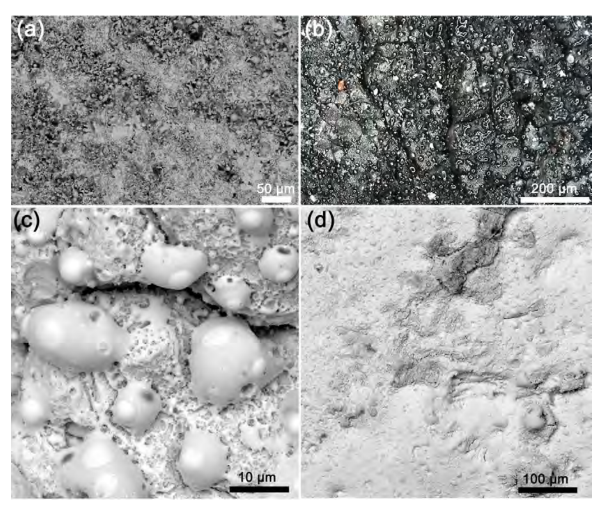


Figure 55. (a) SEMI image of the milially calcanted dead facus from diated a vittgle pingle apollog at 0.1/211/21121. Metrophono philosophilos

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cleaning efforts was when he had after 100 hobos. The he did of the pense of the fluence of pulse alcohol did not increase the cleaning efficiency. There was no cleaning at the fluence of was 150 mJ, and a range of the fluence from 0.20 to 0.76 J/cm² was tested on the chemically 0.35–0.80 J/cm² had a range of the fluence from 0.20 to 0.76 J/cm² was tested on the chemically cleaned sample. SEM study showed fraces of melting already at the minimal fluence 0.50 c), Raman spectroscopy has shown that the globules composition is very heterogene-

cleaned sample. Stavi study showed that the globules composition is very heterogeneobc. Raman spectroscopy has shown that the globules composition is very heterogeneous, and three main groups can be distinguished. The first group consists of synthetic
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mained. The laser fluence was increased to 4.7 J/cm², and all impurities were removed, but the surface became heavily melted (Figure 6a, middle area). SEM study of the irradiated surface highlighted this aspect, which is of great importance during laser cleaning of heritage objects. Laser irradiation fuses the traces of original manufacturing tools, which is extremely undesirable in conservation (Figure 6b). The self-limiting cleaning effect, oc 1301 curring in marble or limestone cleaning [14], was not observed here. In the lead case, after reaching the metal base, ablation intensifies, the plasma jet becomes brighter, and the ablation sound gets louder. The water-assisted cleaning mode, recommended in [15], did in [15], did not prevent surface melting in either EQS Combo laser regimes. Therefore it is not prevent surface melting in either EQS Combo laser regimes. Therefore, it is recommended to avoid both regimes of the EQS Combo laser in lead object cleaning.

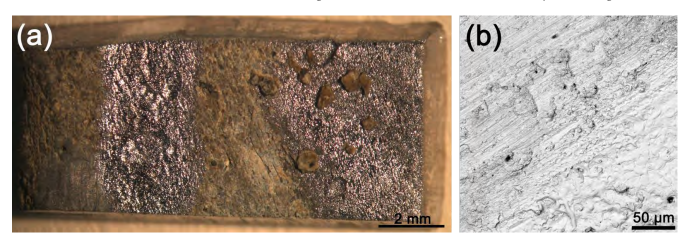


Figure 6. (a) Macro-photo of the sample irradiated in LQS mode. (b) SEM image of the sample surface after several pulses at 4.7 J/cm².

3.3.2.3.66-8-Nanosecond Pulses

In the trials with SpitLight laser, irradiation was performed in the fluence range from In the trials with SpitLight laser, irradiation was performed in the fluence range from 0.02 to 0.4 J/cm² with 20 mJ pulse energy. SEM studies showed that chemically cleaned lead melting started from 0.06 J/cm². When a corroded sample was irradiated at 0.06 J/cm² lead melting started from 0.06 J/cm². When a corroded sample was irradiated at 0.06 J/cm², no cleaning was observed. The number of pulses was increased to 100 but there was no no cleaning was observed. The number of pulses was increased to 100 but there was no cleaning effect. The addition of water or isopropyl alcohol also did not increase the cleaning effect. The addition of water or isopropyl alcohol also did not increase the cleaning ing efficiency. When laser fluence was increased to 0.2 J/cm² a noticeable removal of corrections or isopropyl alcohol also did not increase the cleaning ing efficiency. When laser fluence was increased to 0.2 J/cm² a noticeable removal of corrections or isopropyl alcohol also did not increase the cleaning ing efficiency. When laser fluence was increased to 0.2 J/cm² a noticeable removal of corrections or isopropyl alcohol also did not increase the cleaning ing efficiency. When laser fluence was increased to 0.2 J/cm² a noticeable removal of corrections or isopropyl alcohol also did not increase the cleaning ing efficiency. When laser fluence was increased to 0.2 J/cm² a noticeable removal of corrections or isopropyl alcohol also did not increase the cleaning ing efficiency. When laser fluence was increased to 0.2 J/cm² a noticeable removal of corrections or increased to 0.2 J/cm² a noticeable removal of corrections or increased to 0.2 J/cm² a noticeable removal of corrections of the property of the correction of the property of the correction of the property of the property

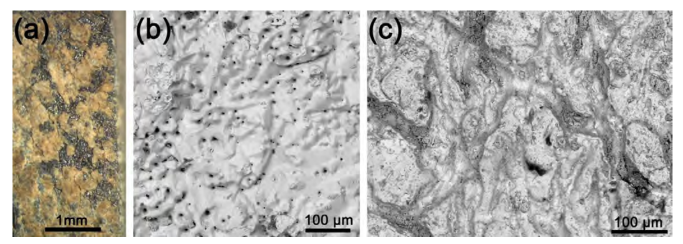


Figure 7. (a) Macro-photo of the area, irradiated with one pulse at 0.2 J/cm², (b) SEM image of the Figure 7. (a) Macro-photo of the area, irradiated with one pulse at 0.2 J/cm² (b) SEM image of sample, cleaned with 100 pulses at 0.4 J/cm² (scale bar 100 μm). (c) SEM image if the corroded samther sample acted with with uses pulses at 0.4 J/cm² (scale bar 100 μm). (c) SEM image if the corroded by the sample acted with with uses pulses at 0.4 J/cm² (scale bar 100 μm). (c) SEM image if the corroded sample, irradiated with 50 pulses were needed at 0.14 J/cm².

The irradiation experiments demonstrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and achieves the herapsutrate that lead corrosion layers have different local densities and layers are considered to the herapsutrate that lead corrosion layers have different local densities and layers are considered to the herapsutrate that lead corrosion layers have different local densities and layers are considered to the herapsutrate that lead corrosion layers have different local densities and layers are considered to the densities and adbasion to the motal surface. The very food laxes of contaminants which are formed by soft user nion paredy control south dust is all usion a can be remaked in wite easily with neveral neveral to the person where it contaminants are nothered have twe akead hosion to thneveraliston sayfas a combreil grantfully a leavest by Jeser ainced in tom I nither process the combre cannot clean the surface completely.

3.2.4. Picosecond Pulses

532 nm, 63 ps

In the experiments with picosecond laser, the first trials were performed at 532 nm.

the corrosion layers are thicker, harder and have stronger adhesion to the metal, the laser cannot clean the surface completely.

3.2.4. Picosecond Pulses

532 nm, 63 ps

In the experiments with picosecond laser, the first trials were performed at 532 nm. The laser spot diameter was constant (3 mm), while pulse energy varied. The chemically cleaned sample was irradiated in the fluence range of 0.004–0.030 J/cm². SEM studies showed that surface melting starts from 0.030 J/cm². When the corrosion layer was irradiated, only at a fluence of 0.065 J/cm² did partial removal begin. An increase in the number of pulses or wetting of the surface did not improve cleaning efficiency. Thus, only by doubling the metal damage threshold was the removal of the corrosion layer achieved.

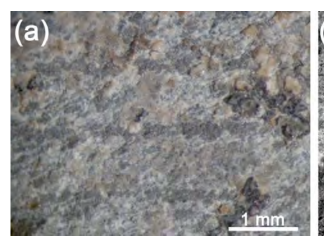
1064 nm, 76 ps

At a wavelength of 1064 nm, when the chemically cleaned sample was irradiated, SEM studies showed melting at 0.01 J/cm^2 . The corroded sample was irradiated in the range from $0.005 \text{ to } 0.070 \text{ J/cm}^2$, and only at 0.06 J/cm^2 did deposits begin to be removed. For the complete cleaning of the surface, around 50 pulses were needed at 0.14 J/cm^2 . Under these conditions, melting and flattening of the surface was observed by SEM (Figure 7c).

The femtosecond laser system has a constant pulse frequency of 10 Hz. To determine

3.2.5. Femtosecond pulses

the threshold for metal damage, it is necessary to obtain a single crater; therefore, the sample was moved by a motorized translation stage at a 2 mm/s speed. By reducing the beam energy with a discrete attenuator, 7 crater tracks were obtained with a laser fluence Heritage 2023, 6, FOR PEER REVIEW 0.07–1.10 J/cm². With SEM studies, it was shown that the single pulse at 0.07 J/cm² did not cause the lead to melt. When a corroded sample was irradiated at 0.07 J/cm², no cleaning was observed. As the initial irradiation of the corrosion layer under these conditions did not reveal When signorious editings the specifical and layer and signorious code to was where the local when signorious editings the specifical and the strain of the corrosion layer under these conditions are specifically signorious editions. The specific spe



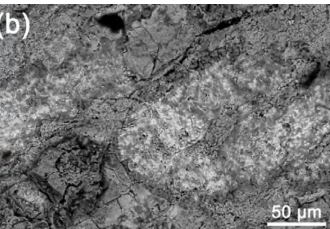


Figure 8. (a) Macro-photo of the horizontal tracks of the laser irradiation of the corrosion layer at 0.1 Figure 8. (a) Macro-photo of the horizontal tracks of the laser irradiation of the corrosion layer at mm/s speed. (b) SEM image of irradiated (0.07 J/cm²) spot. 0.1 mm/s speed. (b) SEM image of irradiated (0.07 J/cm²) spot.

4. Discussion

It was found that for all lasers used in the study, the metallic lead melting threshold is lower than the corrosion layer removal threshold. This may be the result of one or a combination of the following factors:

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4. Discussion

It was found that for all lasers used in the study, the metallic lead melting threshold is lower than the corrosion layer removal threshold. This may be the result of one or a combination of the following factors:

- The spectral reflection coefficients at 1064 nm for metal lead and corrosion layers are almost the same, and approximately 30% (Figure 4). This means that they absorb laser radiation equally well;
- Some part of the incident radiation is effectively scattered and reflected at the surfaces
 of corrosion layer grains; this leads to a decrease in the absorbed energy;
- Lead has a low melting point (327.5 °C). As the bonding energy of the corrosion layers
 is quite high, the laser energy used is not sufficient to remove the corrosion layers but
 generates enough heat to melt the metal;
- The heat capacity of the irradiated materials is quite different. The heat capacity of lead is quite low—0.13 kJ/(kg·K), and a relatively small amount of heat is required to melt it. For comparison, the heat capacities of cerussite and anglesite are (330 J/(kg·K)) and (344 J/(kg·K)), respectively [28];
- The thermal conductivity of lead is relatively low. During the laser heating, the nearsurface layer gets overheated and, as the heat does not have time to transfer to the inner layers, this results in metal melting. The thermal conductivity of the corrosion layer is even lower, but due to the optical properties, absorption occurs mainly in the metal.

Melting of the lead surface during laser cleaning destroys the protective lead oxide layer. As a result, the surface becomes chemically active and corrosion processes can accelerate. To avoid this, the surface should be passivated after cleaning. The surface details, including the traces of the manufacturer's tools, can also be destroyed, which is highly undesirable in the case of cultural heritage objects. In our experiments the µs and 100-ns pulses caused a significant melting effect. In addition, the lead fumes produced are harmful to the operator, so a proper ventilation system should be always in place.

In all the irradiation experiments it was impossible to achieve the selective cleaning of the corrosion layer, based on its composition. The non-uniform cleaning occurs due to an uneven density of corrosion deposits and their different adhesion to the bulk lead. With the lasers used it was also impossible to stop the treatment on the original surface, since there was no perceptible difference for the laser beam.

The atmospheric corrosion of lead results in the formation of caverns. Corrosion products anchor tightly in these caverns. During laser cleaning at high energy densities, the remaining corrosion products fuse into the metal base, which is highly undesirable. To prevent this, high energy densities should be avoided, or chemicals should be used in the final stage of surface cleaning after laser irradiation.

4.1. Surface's Discoloration

4.1.1. Blue Discoloration

With all lasers, straight after melting, the surface acquires a blue color (Figure 9, left). This blue coloring has not been described in previous publications on lead laser cleaning. Since Raman spectroscopy detected only metallic lead in the blue areas, XPS was used to investigate the chemical composition of the irradiated regions (Figure 9, right). The XPS spectrum shows a pair of peaks, characteristic for the spin-orbit splitting of the 4f level of lead and corresponding to the quantum numbers 7/2 and 5/2. The peak energy (138.47 eV for Pb $4f_{7/2}$) indicates that the entire surface layer in the blue region consists of the native lead oxide PbO [29]. Therefore, the blue color of the surface appears not due to the presence of colored chemical substance(s), but rather is caused by light interference in the thin lead oxide film. In a few months after irradiation, the blue color became less noticeable, and disappeared completely on several samples.

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0.9

spectrum shows a pair of peaks, characteristic for the spin-orbit splitting of the 4f level of lead and corresponding to the quantum numbers 7/2 and 5/2. The peak energy (138.47 eV for Pb $4f_{7/2}$) indicates that the entire surface layer in the blue region consists of the native lead oxide PbO [29]. Therefore, the blue color of the surface appears not due to the presence of colored chemical substance(s), but rather is caused by light interference in the thinglead oxide film. In a few months after irradiation, the blue color became less noticeable, and disappeared completely on several samples.

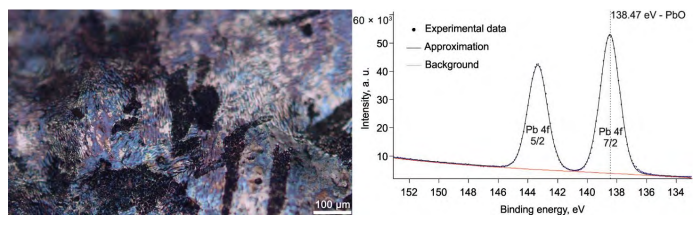


Figure 9. Macro-photo of the surface after irradiation with ns pulses (**left**). XPS spectrum of the 4f **Figure 9.** Macro-photo of the surface after irradiation with ns pulses (**left**). XPS spectrum of the 4f level of lead in the bluish region (**right**). level of lead in the bluish region (**right**).

4.1.2. Yellow Discoloration

In the case of picosecond pulse irradiation, the cleaning assulted in an apparent substance discoloration of source sympletic issue algore issuently shows a magnature of the exposed to no source sympletic issuently substance is no source of the source of

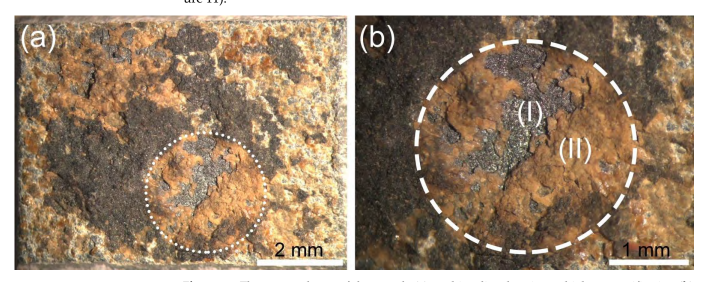


Figure 10. The macro-photos of the sample (a), and irradiated region at higher magnification (b). Figure 10. The macro-photos of the sample (a), and irradiated region at higher magnification (b). The The irradiation parameters: 1064 nm, 76 ps, 10 pulses at 0.06 J/cm².

(c)

The spectrum for area I in Figure 10b has a relatively intense peak with a maximum 200 at 498 cm⁻¹ (Figure 1118). It is unsymmetrically broadened from the low frequency side. Compared to the spectrum for the non-irradiated lead (Figure 11a), a shift to the low-0.9 143 frequency area by about 5 cm $^{-1}$ and an additional peak at 323 cm $^{-1}$ area between which 0.6 may be due to the thermal effect of irradiation on the lead oxide film [23]. Additionally, the 0.3 Spectrum in Figure 11b shows the emergence of intense bands characteristic ${rak d} f$ amorphous carson, namely peaks D (1312 cm $^{-1}$) and G (1590 cm $^{-1}$); the black spectrum corresponds to 0.0 the area with oxide layer and the red spectrum collesponds to the area with high amorphous 0.9 138 carbon content. This is a typical result of organic matter undergoing therrial damage, which 0.6 in the Raman spectrum of the non-irradiated lead gave weak unidentified beaks with a 0.3 mm 0.0

Figure 10. The macro-photos of the sample (a), and irradiated region at higher magnification (b). The irradiation parameters: 1064 nm 76 ps. 10 pulses at 0.06 J/cm² strong luminescence. The background was subtracted during the data processing for better visualization of the cional

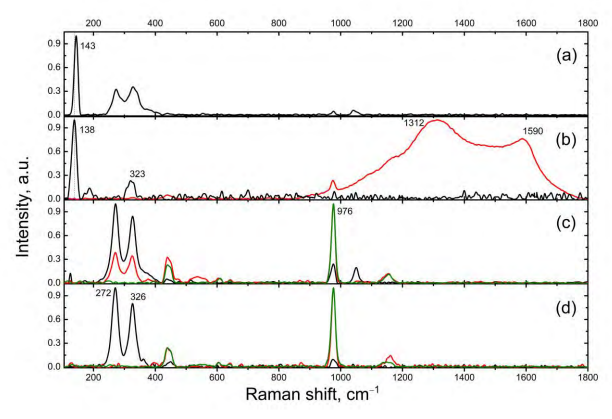


Figure 11. Ramanspettramenallalleadadusuckaseithotttokat dad andra (beir (beir

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5. Conclusions

This study addresses the effects of laser irradiation on a XIX-century outdoor lead sculpture. The effects of pulse duration and laser fluence on the morphology, composition and color of corrosion layers and the base metal were investigated.

It was found that for all lasers used in the study, the metallic lead damage threshold is lower than the corrosion layer removal threshold. Furthermore, there is no self-limiting cleaning effect for the lead atmospheric corrosion deposits and it is impossible to stop the ablation at the original surface. Laser irradiation performs a non-uniform surface cleaning due to the inhomogeneous density and adhesion to metal of the corrosion layers.

In the case of μ s and ns pulses, black globules of mixed composition form on the top of the corrosion layer. The shorter laser pulses do not change the phase composition of the corrosion layer, which is confirmed by Raman spectroscopy studies. This aspect may be important in laser cleaning of archaeological objects, which are covered with a dense layer of corrosion deposits. The benefit of lasers for archaeological lead object cleaning is that, unlike cleaning with hand tools, it does not exert mechanical pressure on the surface being cleaned.

Author Contributions: Conceptualization D.P. (Denis Prokuratov), methodology, D.P. (Denis Prokuratov); writing—original draft, D.P. (Denis Prokuratov) and D.P. (Dmitry Pankin); writing—review and editing, D.P. (Denis Prokuratov), D.R. and V.B.; investigation, A.S. (Andrey Samokhvalov), D.P. (Dmitry Pankin), O.V., N.K., A.P., A.S. (Alexander Shimko), A.M., R.B., A.R. and O.S.; funding acquisition, O.S. All authors have read and agreed to the published version of the manuscript.

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