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Pulsed laser ablation and incubation of nickel, iron and tungsten in liquids and air

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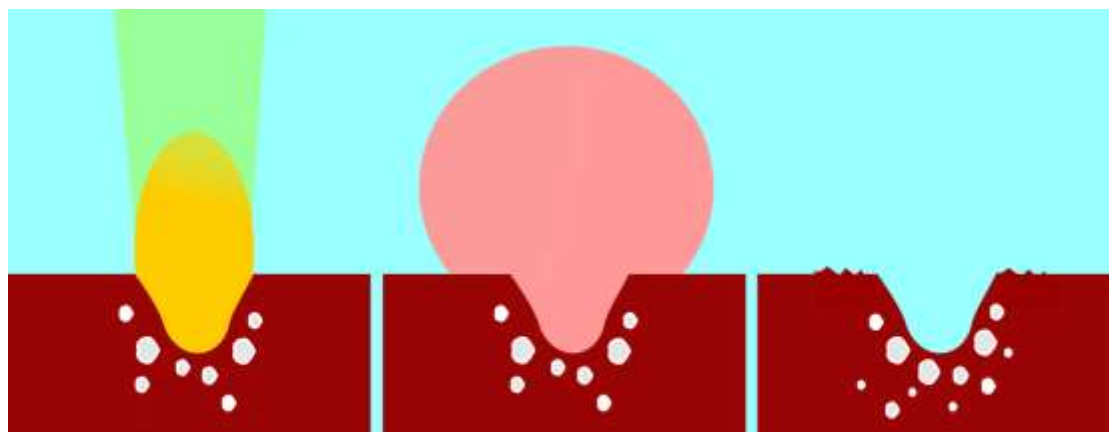
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Graphical Abstract



Highlights

- Laser ablation incubation at air is controlled by thermal properties of the metal.
- Incubation in liquid contact is determined by the mechanical impact on the solid material by the bubble cavitation and the ultimate tensile stress of the metal.

Abstract

Incubation effects in the nanosecond laser ablation of metals exhibit a strong dependence on the thermal and mechanical properties of both the target material and the background gas or liquid. The incubation in air is controlled mainly by thermal properties such as the heat of

vaporization. In liquid, the correlation of the incubation and the ultimate tensile stress of the metals suggests that incubation may be related to the mechanical impact on the solid material by the cavitation bubble collapse, causing accumulation of voids and cracks in the subsurface region of the ablation craters. At high ultimate tensile stress, however, the low sensitivity to the environment suggests that the mechanical impact is likely to play a negligible role in the incubation. Finally, the correlation between the incubation and the carbon content of alcoholic liquids may be explained by an absorptivity increase of the cavity surfaces due to carbonaceous deposits generated by laser-induced pyrolysis, or by the mechanical impact of long-living bubbles at higher dynamic viscosity of liquids.

Keywords

Laser ablation, incubation, metals, fluid interfaces

1. Introduction

Laser processing in liquids has attracted attention because of potential applications in surface cleaning, etching, welding, drilling, cutting, and micromachining of metals, alloys, polymers, semiconductors, glasses and ceramics [1, 2]. Laser ablation of metals in liquids enables the in-situ study of corrosion and repassivation processes [3, 4] as well as the production of biocompatible nanoparticles for medical and catalytic applications [5-7]. The pulsed laser generation of colloidal metal nanoparticles (e.g., Ag, Au, Pt, Pd, Cu, Fe, Ni, W) in distilled water and organic solvents attracted much attention in the past two decades [7-17]. The complex mechanism of laser ablation in liquids involves a series of steps extended over many orders of magnitude in time and involving ablation, plasma expansion inside a cavitation bubble, the penetration of condensed nano-sized phases into the liquid, as well as secondary beam-colloid interaction [18-20].

The dependence of ablation rates and threshold fluences on the number of laser pulses and the background medium, the so-called incubation, has not been understood for metals in contrast to dielectric materials. Therefore, this phenomenon needs further investigations as has been endeavoured in this work.

In order to evaluate ablation and incubation phenomena it is necessary to correlate quantitative ablation and incubation data with the physico-chemical properties of the target materials. In the regime of nanosecond-pulse laser interactions, the behaviour of the ablation is generally

controlled by optical properties and/or thermal conductivity of the target material. In this case, the threshold fluence can be roughly estimated as follows:

$$F_{th} \approx \Delta U l_{eff} (1-R)^{-1}, \quad (1)$$

where ΔU is the energy density needed to heat, melt, and vaporize the target material, R the reflectivity, and l_{eff} is the effective depth of energy deposition [21-23]. In the nanosecond laser interaction with metals, l_{eff} is close to the heat diffusion length $l_T \approx (\delta\tau)^{0.5}$, where τ is the laser pulse duration, $\delta = \kappa/(\rho c_p)$ the thermal diffusivity of the material, κ the thermal conductivity, ρ the density, and c_p the heat capacity. The energy density ΔU can be approximated by the heat of vaporization, $L_v \times \rho$ (Table 1), which accounts for more than 95% of ΔU for any material considered in this work.

Nanosecond pulses show higher threshold values than sub-picosecond pulses [24-28]. Long pulse durations (\geq ns) induce plasma shielding and scattering so that radiation is lost for the absorption process [27], and/or heat diffusion into the bulk materials increases, and, thus, the threshold increases as well (acc. to Eq. 1). The experimental determination of the ablation threshold can be based on the evaluation of the squared diameter of the ablated zone, D^2 , related to the Gaussian beam radius (w_0), the fluence F , and the threshold fluence F_{th} with the assumption of a Gaussian beam profile [22, 29]:

$$D^2 = 2w_0^2 \ln (F/F_{th}). \quad (2)$$

When the ablation exhibits incubation behaviour, the threshold fluence as a function of the number of pulses, $F_{th}(N)$, can be represented as [22, 29]

$$F_{th}(N) = F_{th}(1) N^{\xi-1}, \quad (3)$$

where ξ is a material-dependent incubation coefficient, $F_{th}(1)$ is the single shot ablation threshold, and N is the number of pulses. In this simple phenomenological model, it was suggested that the incubation is related to the accumulation of mechanical damage. If $\xi = 1$, incubation is absent, whereas for values between 0 and 1 the material is weakened by defect accumulation [22, 29]. For metals, the factor ξ is typically between 0 and 1. The origin of the incubation in metals is still under debate. It was suggested that incubation might be related to fatigue damages [29, 30]. A theoretical expression for the incubation coefficient is not existent, nor is a description of which materials should exhibit a larger incubational behaviour. There have been few attempts to quantify the influence of the target material conversion and structural modification in a multi-pulse experiment by phenomenological [22, 29, 31-33] and physical models based on reflectivity changes due to roughness evolution at low pulse numbers [34] and

bulk defects [35, 36]. In the later model, incubation was attributed to high-density defects (HDDs) in the absorption volume of the solid.

The finite incubation of metals in air may be correlated with the formation of voids and/or cracks in the resolidifying layers that remain after the ablation process [7, 37, 38]. The subsurface regions affected by the void/crack formation are expected to exhibit significantly lower heat diffusion lengths, l_T , than the homogeneous bulk material due to strongly reduced heat diffusivities δ . Thus, the heat dissipation is reduced and the ablation threshold decreases (Eq. 1) with the void accumulation. No incubation investigations exist so far for laser ablation in liquids.

In this work, models considering various materials properties and structural modifications [7, 35-38] - beyond the well described optical changes (comp. dielectrics) - are discussed and semi-quantitatively correlated with the observed contrasting incubation behaviour of Ni, Fe, and W in various media.

2. Experimental

The target materials were nickel (Alfa Aesar; purity $\geq 99.5\%$, $50 \times 50 \times 2$ mm), iron (Goodfellow; purity 99.5%, tempered, round disk, diameter 25 mm, thickness 0.5 mm), and tungsten (Goodfellow; 99.95%, tempered, as rolled, $20 \times 20 \times 0.5$ mm). The liquids were distilled water, ethanol, butanol, and isopropanol (Sigma-Aldrich; p.a.). The metal platelets were cleaned by ultrasonication in ethanol, and were positioned in a glass cell with an optical window allowing the horizontal access of the laser beam. This was focused by a plano-convex lens with a focal length of 92 mm yielding a depth of focus of 1.5 mm. The cell was positioned on a motorized XY-scanning stage. The energy attenuation was performed by a polarizer with a half-wave plate. A power meter (OPHIR Photonics) could be positioned after the polarizer (THORLABS). The focus position in air and various liquid media were experimentally evaluated by microscopically measuring the ablation area on a silicon target (Zeiss AxioVision software) as a function of the distance of the focusing plano-convex lens. A Q-switched Nd:YAG laser system was employed emitting at a wavelength of 532 nm (Spectra Physics GCR-130, ≤ 1.2 W, pulse duration of 5 ns, repetition rate of 20 Hz, beam diameter of about 5 mm). The images of the modified target regions were recorded by a CCD camera connected to an optical microscope (OLYMPUS, STM-MJS). The recorded ablated areas were evaluated by Zeiss AxioVision software. From this, the average diameter D and D^2 were calculated in order to evaluate the $(D^2 - \ln F)$ -relationship [22, 29]. In order to evaluate the Gaussian beam radius, the so-called

cutting edge technique was applied [39, 40]. Imaging of various morphologies of the laser-ablated craters was performed by scanning electron microscopy (SEM; Zeiss Supra 55 VP).

3. Results and Discussion

Laser ablation in various fluids in comparison to air environment was performed at various F - and N -values. The phenomenon of incubation relies on laser-induced conversion, modification of target microstructure, and phase changes. Therefore, scanning electron microscopy with secondary electron emission (SE-SEM) detection was performed on ablation craters of Ni (Fig. 1) and Fe (Fig. 2) in both water and ethanol, and on W in water (Fig. 3) in comparison to air environment. SE-SEM was chosen to achieve surface-sensitive information (secondary electron escape depth < 20 nm) on the morphology and possibly also on conversion products. Charging effects and/or varying secondary electron emission rates can support this analysis [41, 42].

The SE-SEM image of a nickel crater generated in air shows strong charging in contrast to the pristine surface of the sample (Fig. 1A). This is indicative of the conversion of Ni to a much thicker NiO_x film inside the crater as compared to the untreated surface. Actually, NiO_x exhibits an extremely low conductivity at room temperature only comparable to fully oxidized iron oxide, Fe_2O_3 [43]. When the ablation takes place under water, less formation of NiO_x can be expected. Only few thicker isolated oxide regions appeared as “flakes” in the crater bottom (Fig. 1B). This suggests that the mixture of water and Ni vapour during the bubble lifetime does not provide as much reactive oxygen for a thick coverage of the crater with this low-conductivity NiO_x in contrast to air (Fig. 1A). Actually, gas chromatography showed that during laser ablation in water, water splitting led to both hydrogen and oxygen in the bubbles [44]. Both the presence of hydrogen and the relative shortage of oxygen in the bubble result in the formation of less NiO_x (Fig. 1B) as compared to the air case (Fig. 1A). The ablation crater under ethanol also shows much less charging than in the case of air (Fig. 1A) but still exhibits a pronounced morphology with increased SE emission that is not restricted to the edges (Fig. 1C). That may indicate the formation of conductive non-oxidic conversion phases, such as nickel carbide due to the carbonization of ethanol [44-48]. Further investigations on this issue are under way.

An SE image of an iron ablation crater in air (Fig. 2A) does not exhibit the type of charging as observed with Ni (comp. Fig. 1A). Fe certainly converted to an oxide after laser treatment. Oxidation of iron leads to different iron oxides (FeO_x) with various phases such as hematite (α - Fe_2O_3), maghemite (γ - Fe_2O_3), wüstite (FeO) and magnetite (Fe_3O_4). Hematite is the most stable

phase of FeO_x with an optical band gap of ~ 2.2 eV [49-51]. However, the presence of a plasma containing a finite concentration of free electrons may provide a reducing function [52]. Thus, the resulting Fe oxides contain lower oxidation states, e.g. in FeO or Fe_3O_4 , which show metallic conductivity [43], and therefore no charging in SEM. Apparently, the FeO or Fe_3O_4 mixture typical for iron in air has been formed. The exception of this finding is observed at the crater edges, where strongly charged “flakes” appeared (Fig. 2A). Actually, air oxygen could access the heated iron target freely outside the plasma region resulting in completely oxidized iron, i.e. Fe_2O_3 , which has at least 20 orders of magnitude lower conductivity than the Fe^{2+} containing oxide phases [43]. The craters under water (Fig. 2B) and ethanol (Fig. 2C) showed no charging suggesting the absence of Fe_2O_3 , because the fluid confined plasma protected the crater surfaces from complete oxidation, leading only to highly conductive species FeO and Fe_3O_4 .

Tungsten ablation in air showed no charging of oxide phases (Fig. 3A) both at the edges and in the crater. The crater walls exhibited a smoother morphology than the original target surface due to melting and resolidification followed by oxidation. It should be considered that WO_3 under atmospheric conditions is intrinsically ‘self-doped’ by native oxygen vacancy point defects resulting in coloured WO_{3-x} films with transparencies depending on the level of oxygen vacancies [53-57]. The plasma containment under water led to surface modification by the strong cavitation impact extending outside the irradiated area (Fig. 3B).

The evaluation of the ablation thresholds and the incubation behaviour is based on the measurement of the crater diameters D by optical microscopy and fitting the data points to Eq. 2. In order to document this procedure, but also the collateral modification features around the craters, representative images of the respective ablation craters of Ni (Fig. 4) and Fe (Fig. 5) in both water and ethanol, and of W in water (Fig. 6) are presented.

In water, Ni shows a modification zone with reduced reflectivity next to the crater edges increasing with fluence, i.e. from ca. $50\text{ }\mu\text{m}$ at 100 Jcm^{-2} up to ca. $300\text{ }\mu\text{m}$ at 1100 Jcm^{-2} (Fig. 4a). A closer inspection of the respective SE-SEM image (Fig. 1B) shows no major NiO_x growth within the cavity zone. The origin of this phenomenon is likely to be related to the cavitation bubble dynamics, although the mechanistic details are not yet established. The zone of reduced reflectance is much smaller in ethanol (Fig. 4b), i.e. $< 10\text{ }\mu\text{m}$ at 100 Jcm^{-2} up to ca. $50\text{ }\mu\text{m}$ at 1100 Jcm^{-2} .

The optical images of craters on Fe in water (Fig. 5a) showed the roughened region outside the craters similarly to the SE-SEM images (Fig. 2B). This kind of cavitation regions in ethanol

exhibited higher optical reflectivity (Fig. 5b). Possibly carbonaceous conversion products due to the pyrolysis of ethanol [45-48] may be the cause. The optical images of W craters in water (Fig. 6) show again roughening outside the cavities.

The multi-pulse incubation behaviour quantified by the material-dependent incubation coefficient ξ was evaluated according to Eq. 3 based on a D^2 - $\log F$ evaluation of the values of the threshold fluence (Eq. 2). Representative results are depicted for Fe in ethanol (Fig. 7 and 8). Due to the difficulties with the evaluation of diameters for shallow craters generated at low pulse numbers N and low fluences, only the data points at higher N and F were considered in fitting to Eq. 2. All evaluated incubation coefficients ξ are presented in Fig. 9. The incubation of the substrates in air depends strongly on the metal type, with Ni exhibiting negligible and W the highest incubation.

All investigated metals exhibit a moderate, but finite incubation in air, i.e. $\xi < 1.0$ (Fig. 9). The incubation in metals may be attributed to the formation and accumulation of voids and/or cracks in the subsurface layer within the laser spot. A porous region generated due to the void formation is expected to exhibit a drastically reduced heat diffusion length l_T and thermal diffusivities δ as compared to the homogeneous bulk material. Thus, the heat dissipation can be reduced and the ablation threshold may decrease (comp. Eq. 1) due to the sub-surface void accumulation [3, 7, 37, 38]. This qualitative hypothetical model is correlated to a recently developed concept for sub-picosecond laser interactions with metals, where nanovoids are observed to form due to rarefaction processes in the near-surface region of the target [37, 59]. Further research on the applicability of this model to the nanosecond irradiation regime is under way.

Ni in air exhibited negligible incubation ($\xi \approx 1$, Fig. 9) suggesting that the before-mentioned qualitative void/crack formation mechanism in the surface-near bulk is of minor importance. In water, however, a substantially higher incubation ($\xi = 0.52$) with respect to air environment is observed. Possible reaction products, such as oxides, should not have a strong effect on the irradiation absorption. Therefore, the possibility of an accumulation of defects, such as voids and cracks (comp. [37, 38]), below the cavities due to the cavitation impact under water may be considered as cause for the increased thermal containment with increasing N . A contribution of the reduction of the thermal diffusivity by the admixture of oxide phases into the metal matrix may also contribute to the thermal containment.

The comparison of incubation coefficients ξ in water and alcohols, which exhibit strongly varying surface tension γ (22-25 mN m⁻¹ for alcohols [60, 61] and 72 mN m⁻¹ for water), shows

a lack of clear correlation between ξ and γ . In contrast to acoustically generated cavitation bubble dynamics [62], laser induced bubbles may be driven by overpressure with minimal contribution from surface tension [63]. However, the estimations of the vapour pressure at the bubble wall based on the values of surface tension and dynamic viscosity of water and alcohols and the equation given in Ref. [64] reveals a correlation between the pressure and incubation.

Moreover, the incubation in Ni significantly increased (ξ is reduced) with increasing number of carbon atoms in the alcohol molecules (Fig. 9). Solvents with a higher number of carbon atoms generally show a pyrolysis on hot metal surfaces [45-48]. One can assume that each additional laser pulse leads to more carbonaceous products on the metal target. This is supported by recent μ -Raman experiments to be reported elsewhere [65]. Thus, increasing the absorption at the metal-liquid interface may result in what can be understood as incubation.

Fe in air exhibited a low ($\xi \approx 0.9$) but somewhat higher incubation than Ni (Fig. 9) suggesting that the before-mentioned qualitative void/crack formation mechanism [37, 38, 59] plays a moderate role. The presence of liquid environment, water or ethanol, led to a substantially higher incubation ($\xi \approx 0.6$) with respect to air environment. By analogy with Ni, the possibility of an introduction of defects (voids and cracks, comp. [37, 38]) below the cavities caused by the cavitation impact may be considered. Thus, the thermal confinement with increasing N may lead to the observed incubation.

W in air (Fig. 7) exhibited the highest incubation ($\xi = 0.75$) of all metals. This finding suggests that the multiple void/crack formation mechanism in the sub-surface region is more effective for W. The presence of water led only to a slight increase in incubation ($\xi = 0.67$) in contrast to the behaviour of Ni and Fe.

Ablation in air is controlled mainly by thermal properties of the metal targets such as the heat of vaporization and the possible formation of voids and cracks near the ablation craters [37, 38, new Balling]. Thus, a correlation between the incubation coefficients ξ with the heat of vaporization L_v [66] could be observed (Fig. 10), i.e. a lower L_v supports an increased void formation.

The incubation in liquid medium generally increases in comparison with air (Fig. 9). There is practically no correlation of ξ in water with L_v (Fig. 10). However, it is correlated with the strength of mechanical impact on the metals by the bubble cavitation. The extent of the increase of ξ depends strongly on the ultimate tensile stress σ_{TS} of the metal [67] (Fig. 11). Incubation of a metal with low σ_{TS} (e.g., Ni) shows a drastic ξ decrease (incubation increase) suggesting the thermomechanical nanovoid/crack formation mechanism in the cavitation process. In

contrast, the mechanical cavitation mechanism plays a rather negligible role in the incubation in a metal with high σ_{TS} (e.g. W) because ξ is almost the same as in air.

A third, optical absorption mechanism may come into play when alcoholic fluids with varying carbon numbers are applied (Fig. 9). An influence of this number on the incubation may suggest a possible absorptivity increase of the cavity surfaces due to carbonaceous deposits generated by laser-induced pyrolysis, particularly at higher carbon contents like in butanol.

4. Conclusions

The ablation and incubation of metals in air is correlated with thermal properties such as the heat of vaporization, and may be related to the accumulation of high-density defects such as voids and cracks below the ablation crater surface. The introduction of a liquid medium changes the cause for high-density defects from a thermal to a mechanical due to bubble cavitation impact, as suggested by the correlation of the incubation and the ultimate tensile stress of the metal. In particular, nickel exhibits relatively low ultimate tensile stress and shows the highest incubation increase in a liquid medium. This is in strong contrast to tungsten that has a much higher ultimate tensile stress than nickel and shows an incubation behaviour barely affected by a liquid environment. An optical absorption mechanism may become important when alcohols with varying carbon numbers are applied: the carbon content could affect the tendency to generate carbonaceous deposits by laser-induced pyrolysis thus increasing the incubation.

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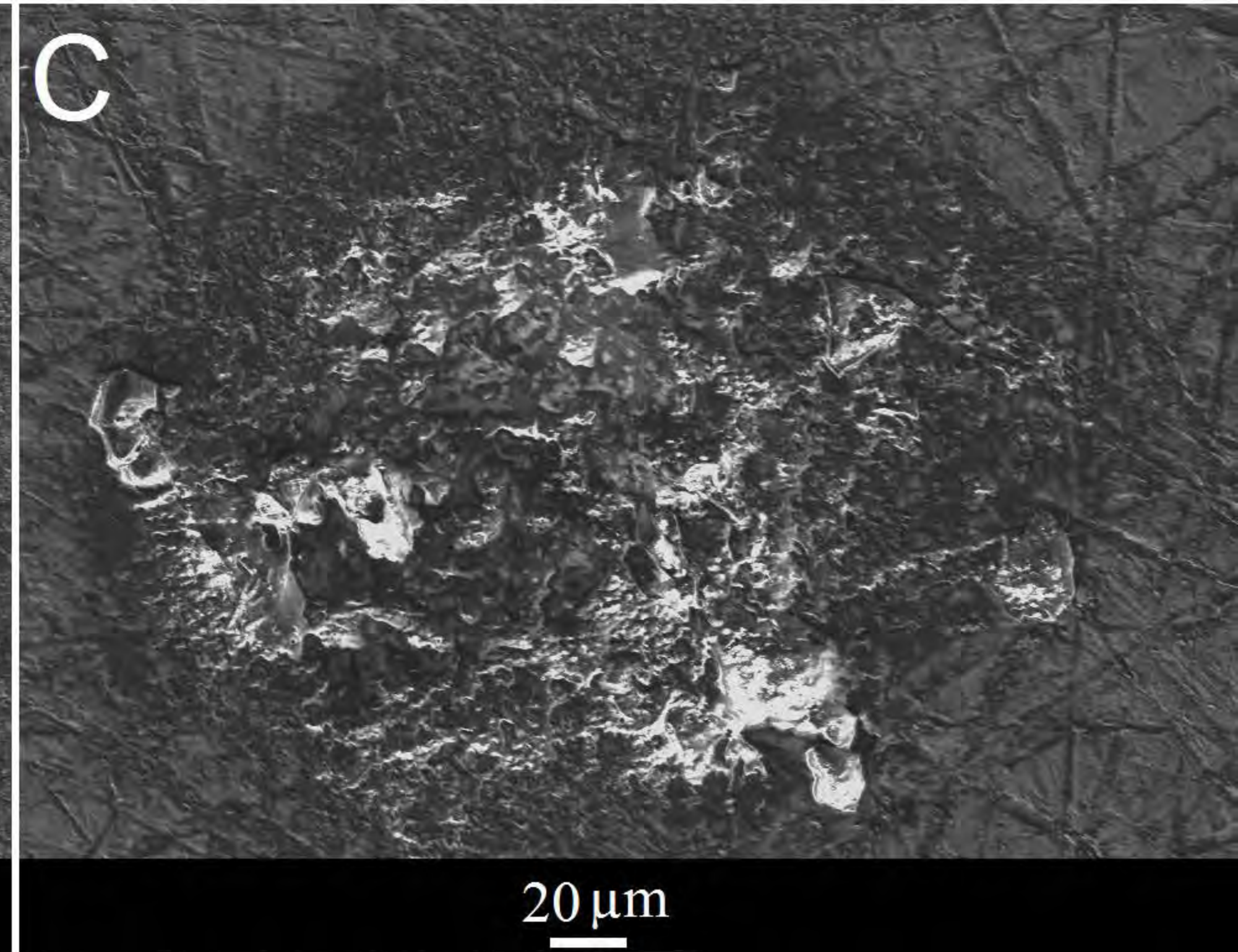
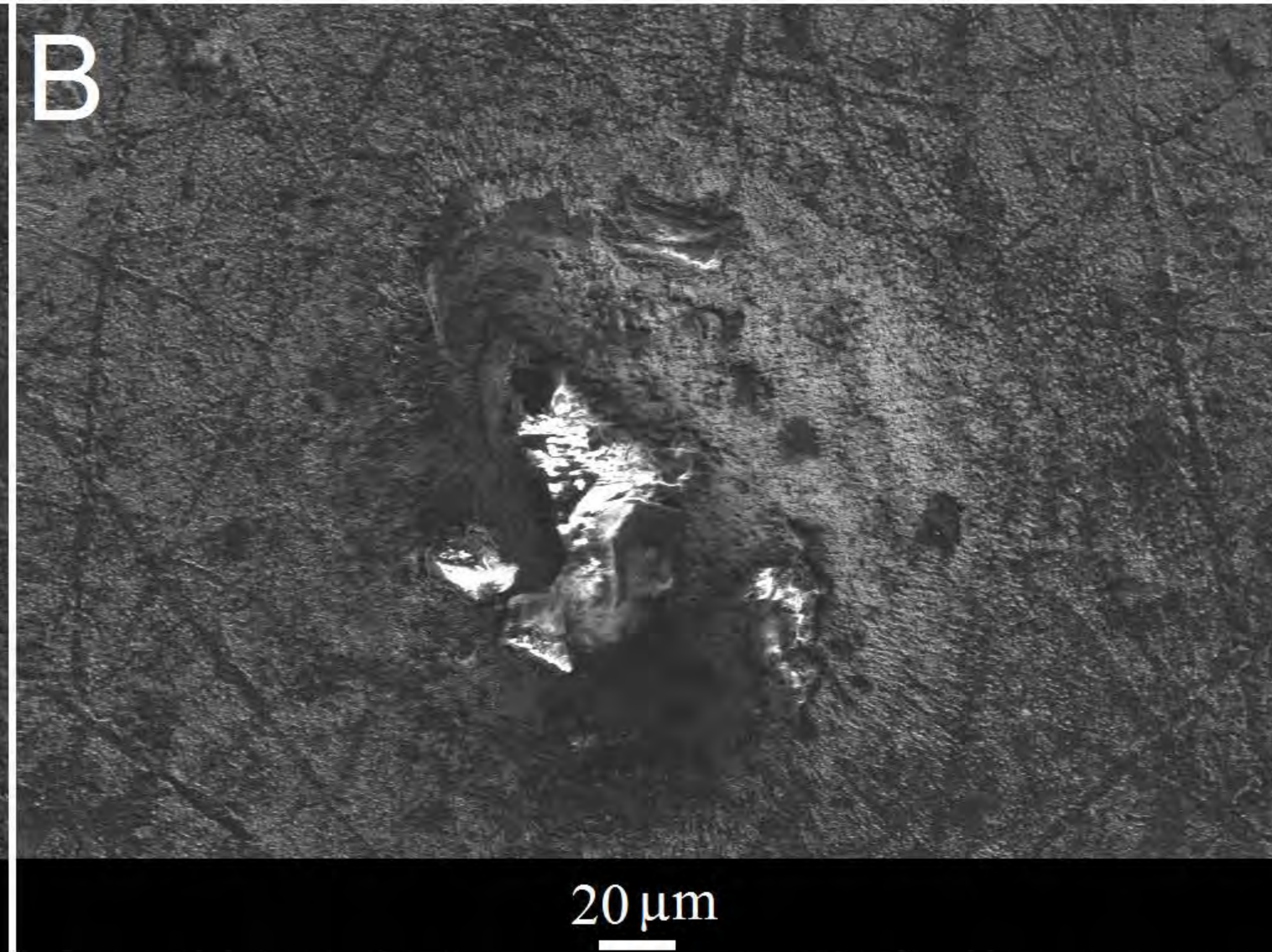
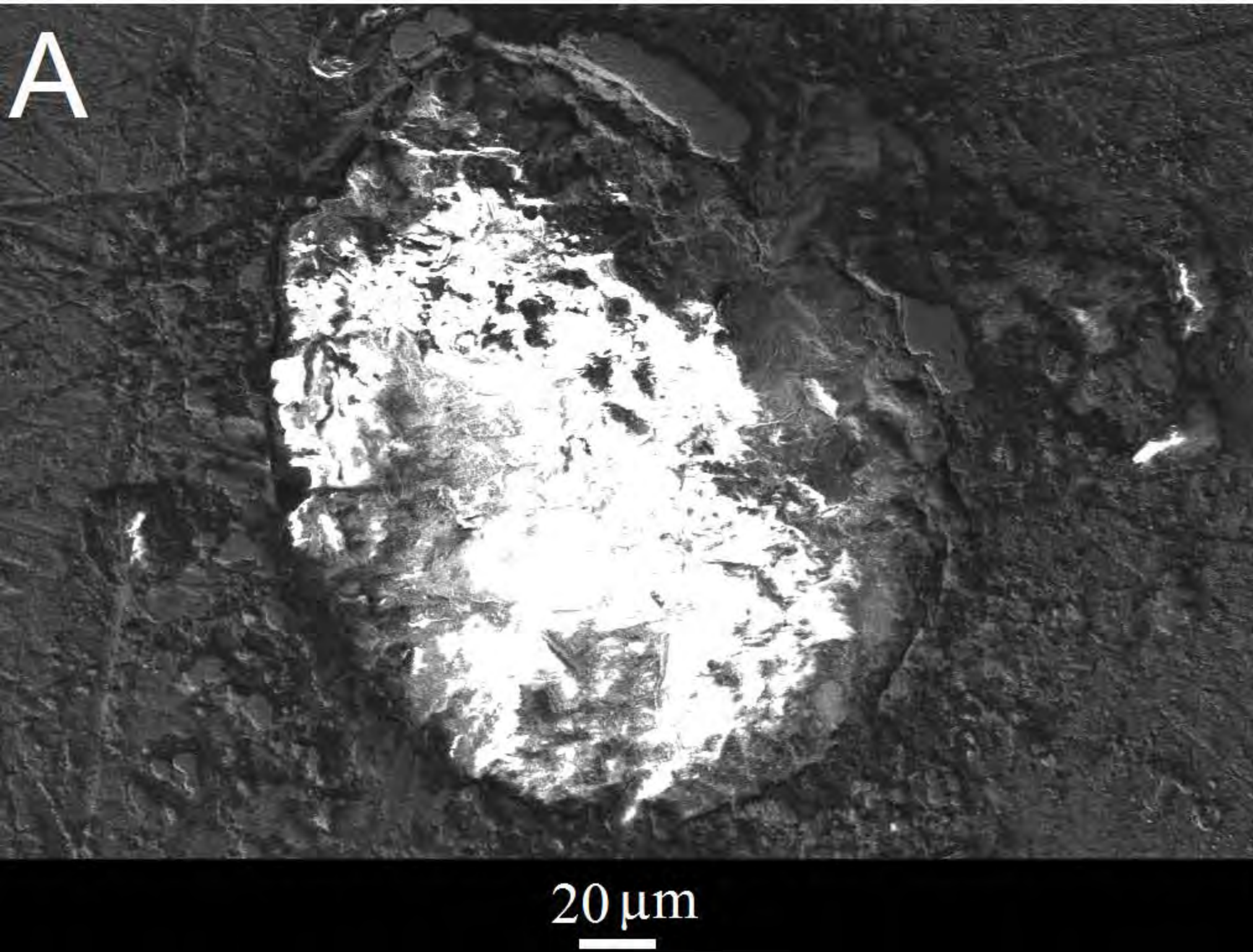
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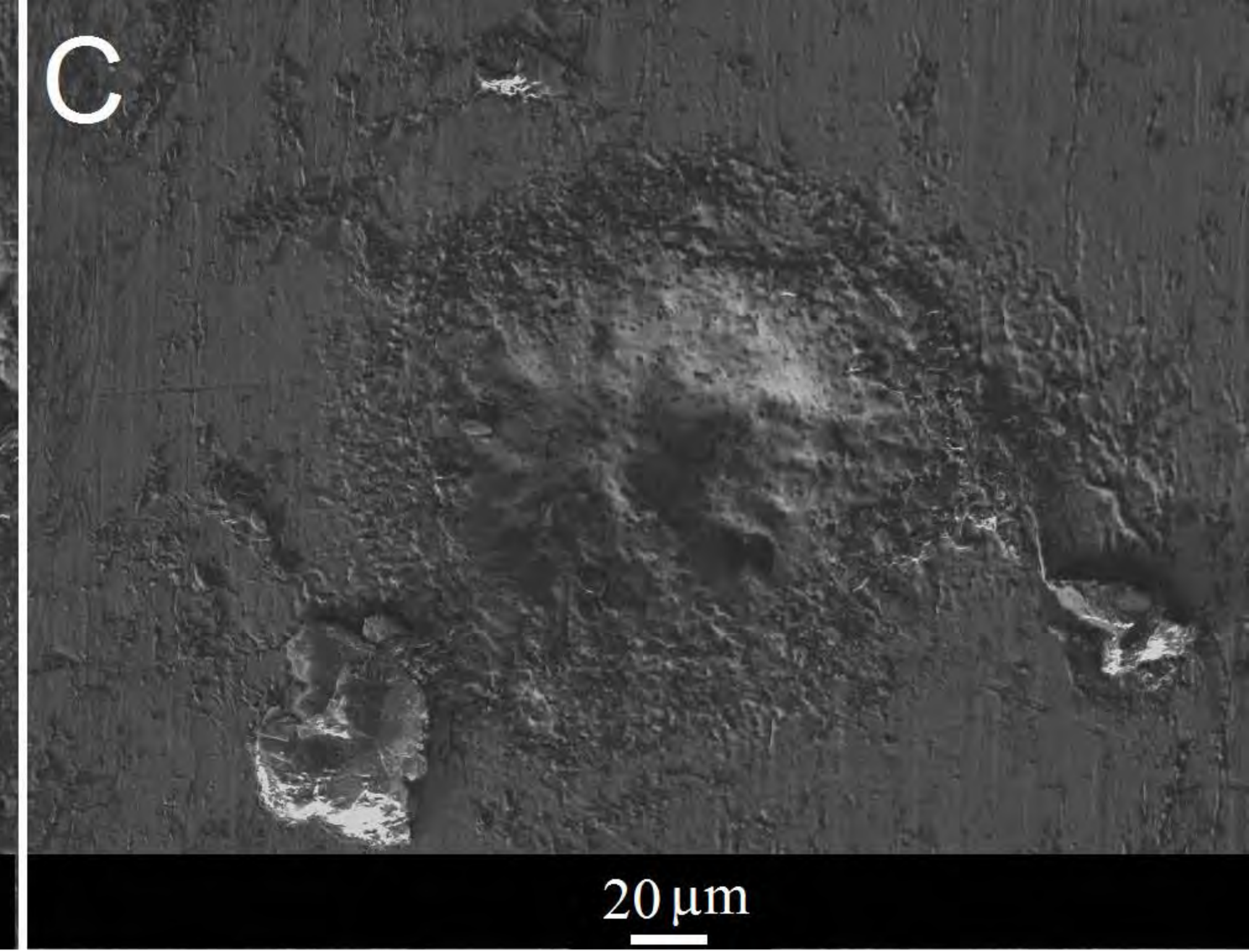
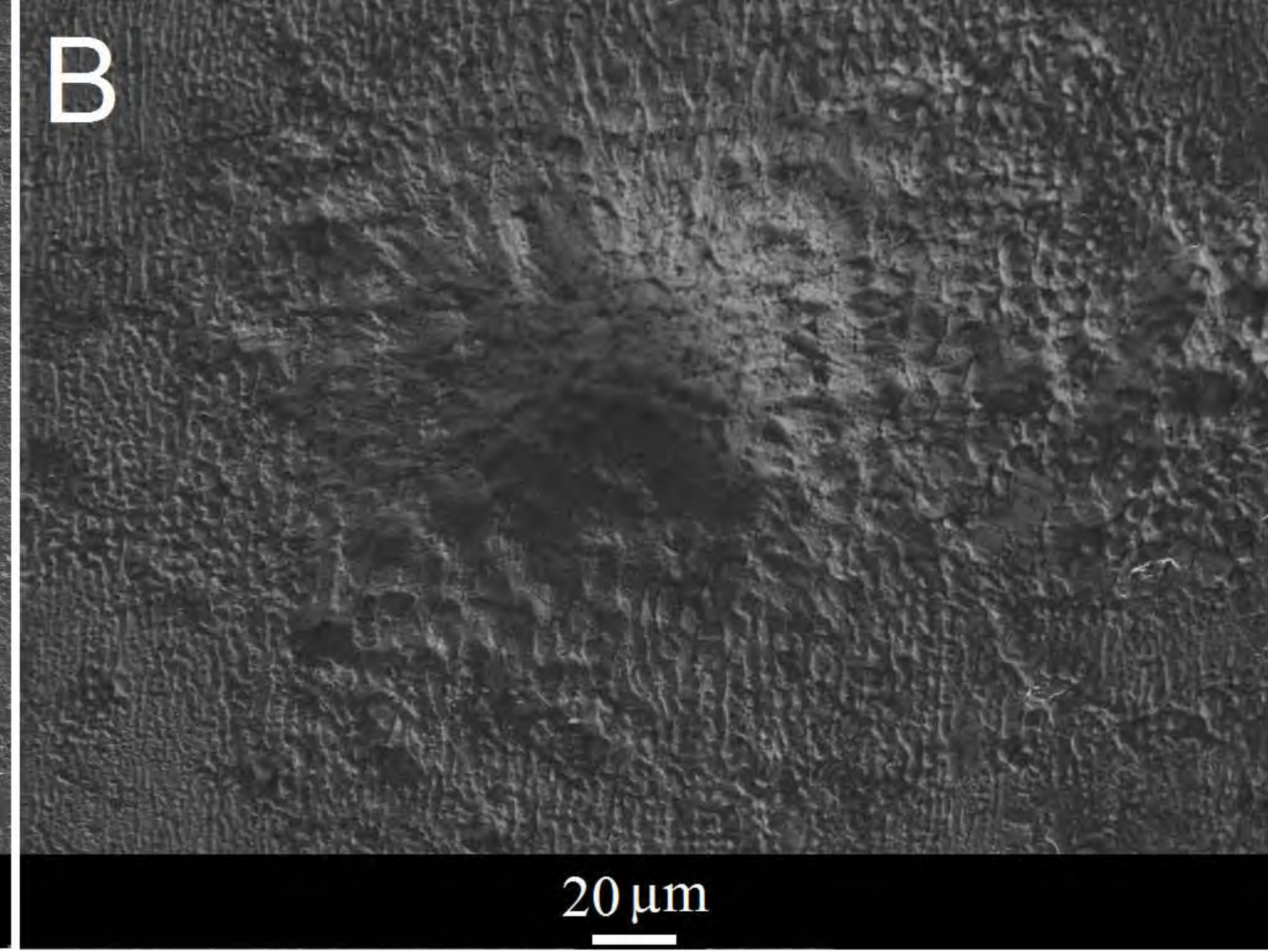
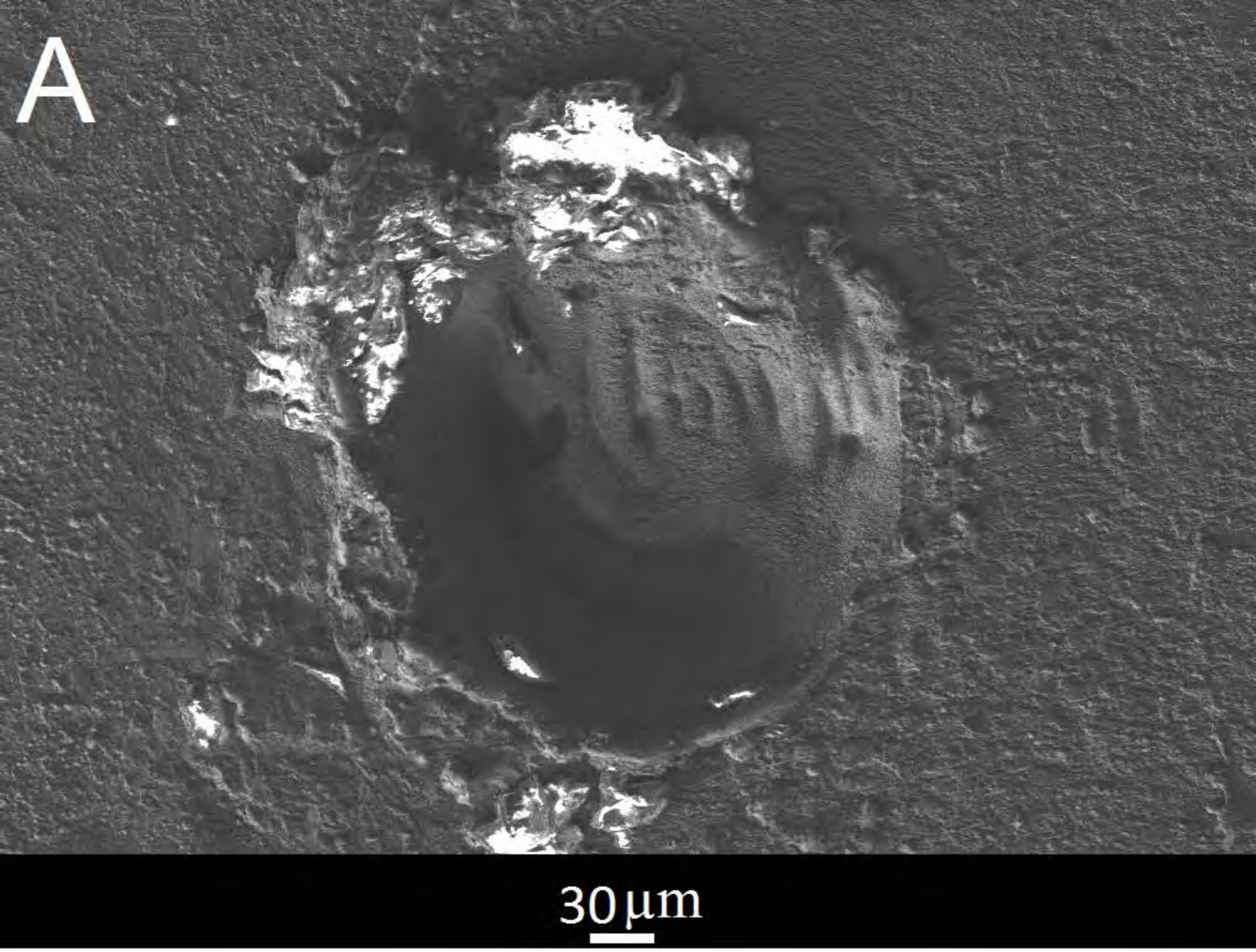
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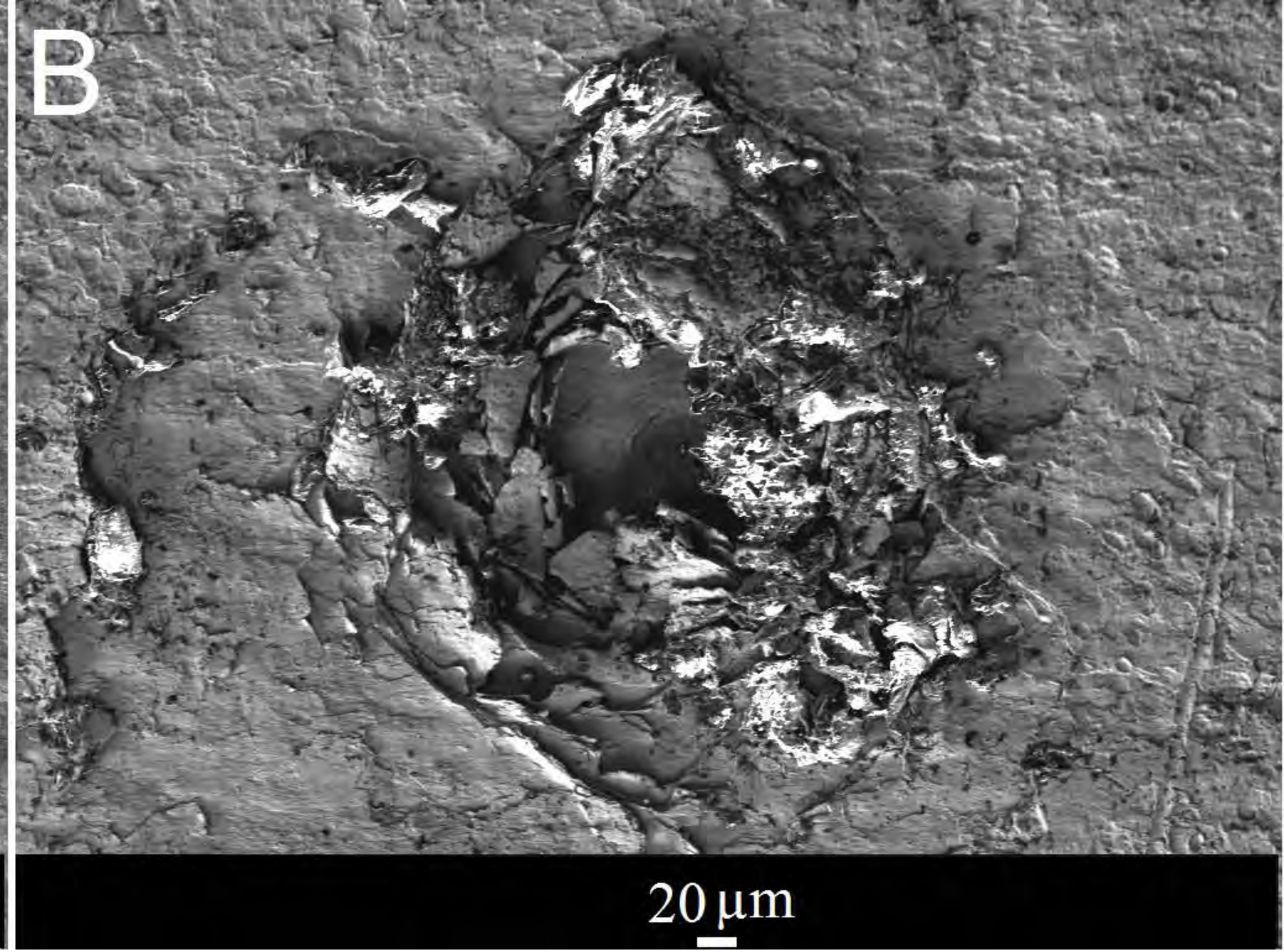
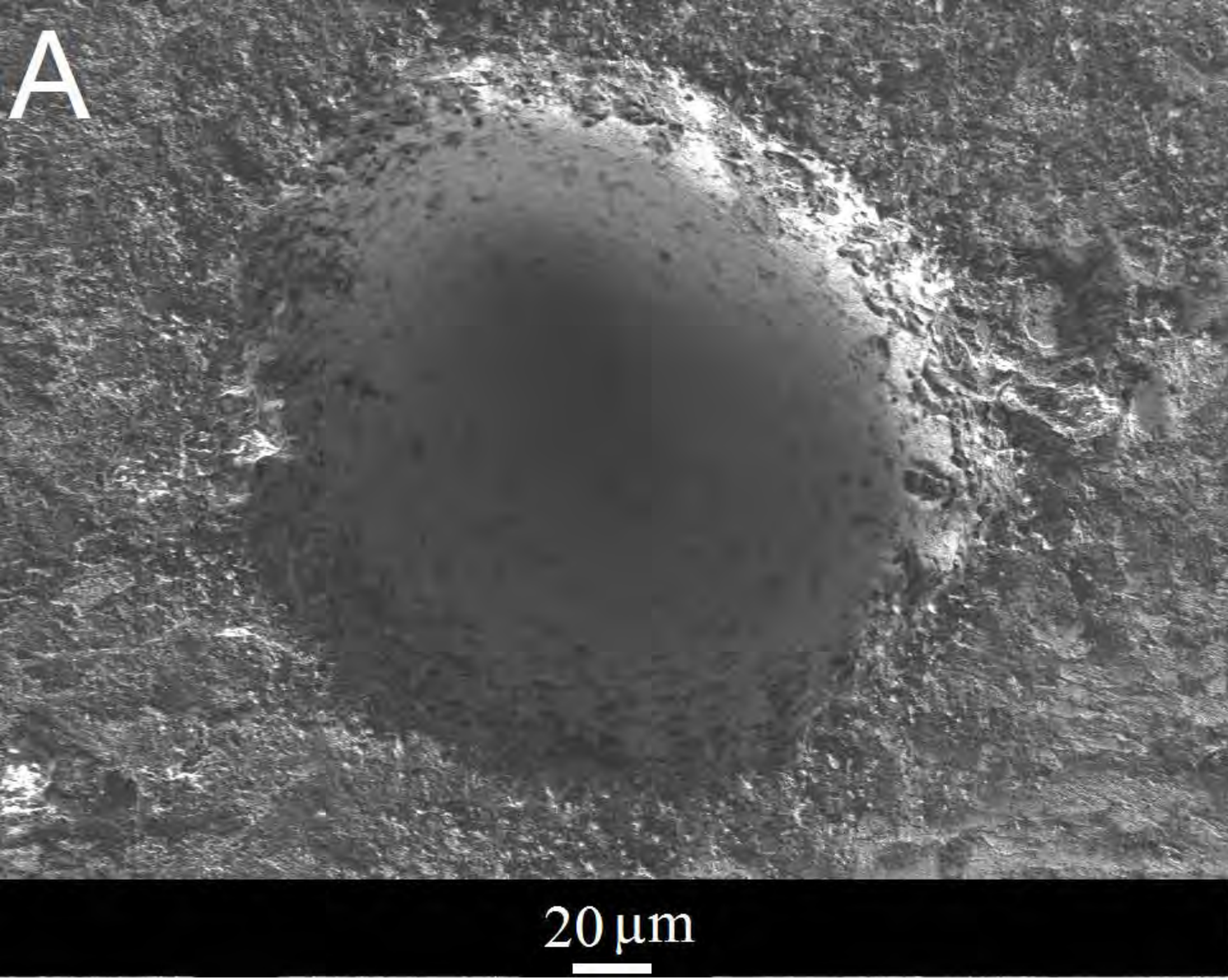
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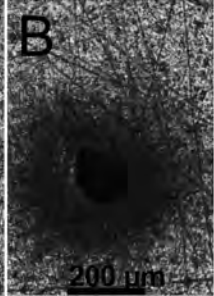
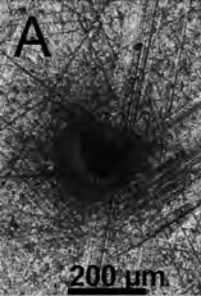
Figure Captions

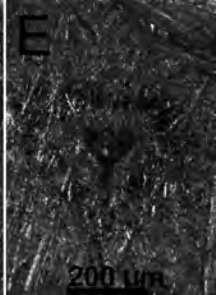
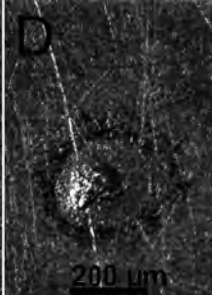
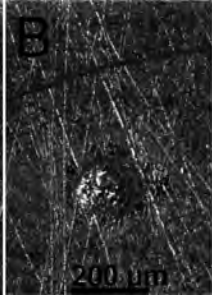
- Figure 1** SEM images of Ni ablated craters in various media; A: air ($N=500$, $F = 310 \text{ Jcm}^{-2}$); B: Water ($N=500$, $F = 400 \text{ Jcm}^{-2}$); C: Ethanol ($N=500$, $F = 400 \text{ Jcm}^{-2}$).
- Figure 2** SEM images of Fe ablated craters in various media; A: air ($N=500$, $F = 310 \text{ Jcm}^{-2}$); B: Water ($N=500$, $F = 400 \text{ Jcm}^{-2}$); C: Ethanol ($N=500$, $F = 400 \text{ Jcm}^{-2}$).
- Figure 3** SEM images of W ablated craters in various media; A: air ($N=100$, $F = 310 \text{ Jcm}^{-2}$); B: Water ($N=100$, $F = 700 \text{ Jcm}^{-2}$).
- Figure 4** Representative optical micrographs of laser-treated nickel. $N = 500$. a: in water. b: in ethanol. A: $F = 100 \text{ Jcm}^{-2}$, B: $F = 200 \text{ Jcm}^{-2}$, C: $F = 400 \text{ Jcm}^{-2}$, D: $F = 700 \text{ Jcm}^{-2}$, E: $F = 1100 \text{ Jcm}^{-2}$.
- Figure 5** Representative optical micrographs of laser-treated iron. a: in water. b: in ethanol. $N = 500$. A: $F = 100 \text{ Jcm}^{-2}$, B: $F = 200 \text{ Jcm}^{-2}$, C: $F = 400 \text{ Jcm}^{-2}$, D: $F = 700 \text{ Jcm}^{-2}$, E: $F = 1100 \text{ Jcm}^{-2}$.
- Figure 6** Representative optical micrographs of laser-treated tungsten in water. $N = 100$. A: $F = 100 \text{ Jcm}^{-2}$, B: $F = 200 \text{ Jcm}^{-2}$, C: $F = 400 \text{ Jcm}^{-2}$, D: $F = 700 \text{ Jcm}^{-2}$, E: $F = 1100 \text{ Jcm}^{-2}$.
- Figure 7** Squared diameter of the ablation area versus pulse fluence (D^2 vs F ; Eq. 2). Fe in ethanol. \diamond : $N = 50$, \blacksquare : $N = 100$, \circ : $N = 200$, \blacktriangle : $N = 500$, \bullet : $N = 1000$.
- Figure 8** Incubation behaviour of iron in ethanol (acc. Eq. 3).
- Figure 9** Incubation coefficients in air and various liquids. Ni data from [7].
- Figure 10** Incubation coefficients versus heat of vaporization L_v (from [65]). \bullet : in air; \circ : in water. The arrows indicate the change of the incubation coefficient upon immersion into water.
- Figure 11** Incubation coefficients versus ultimate tensile stress σ_{TS} (from [66]). \bullet : in air; \circ : in water. The arrows indicate the change of the incubation coefficient upon immersion into water.

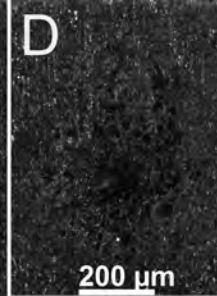
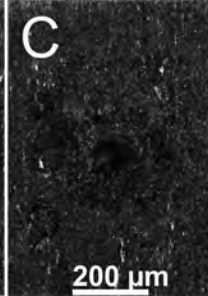


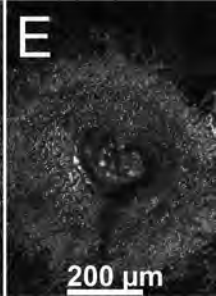
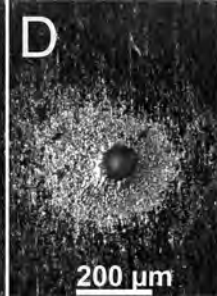
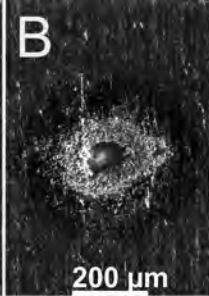


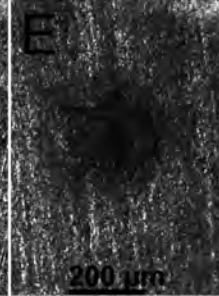
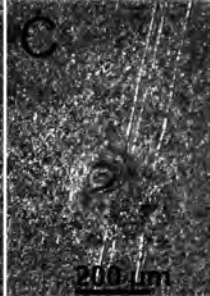
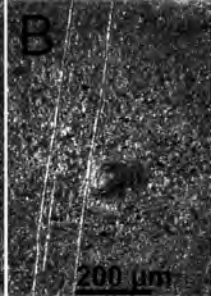


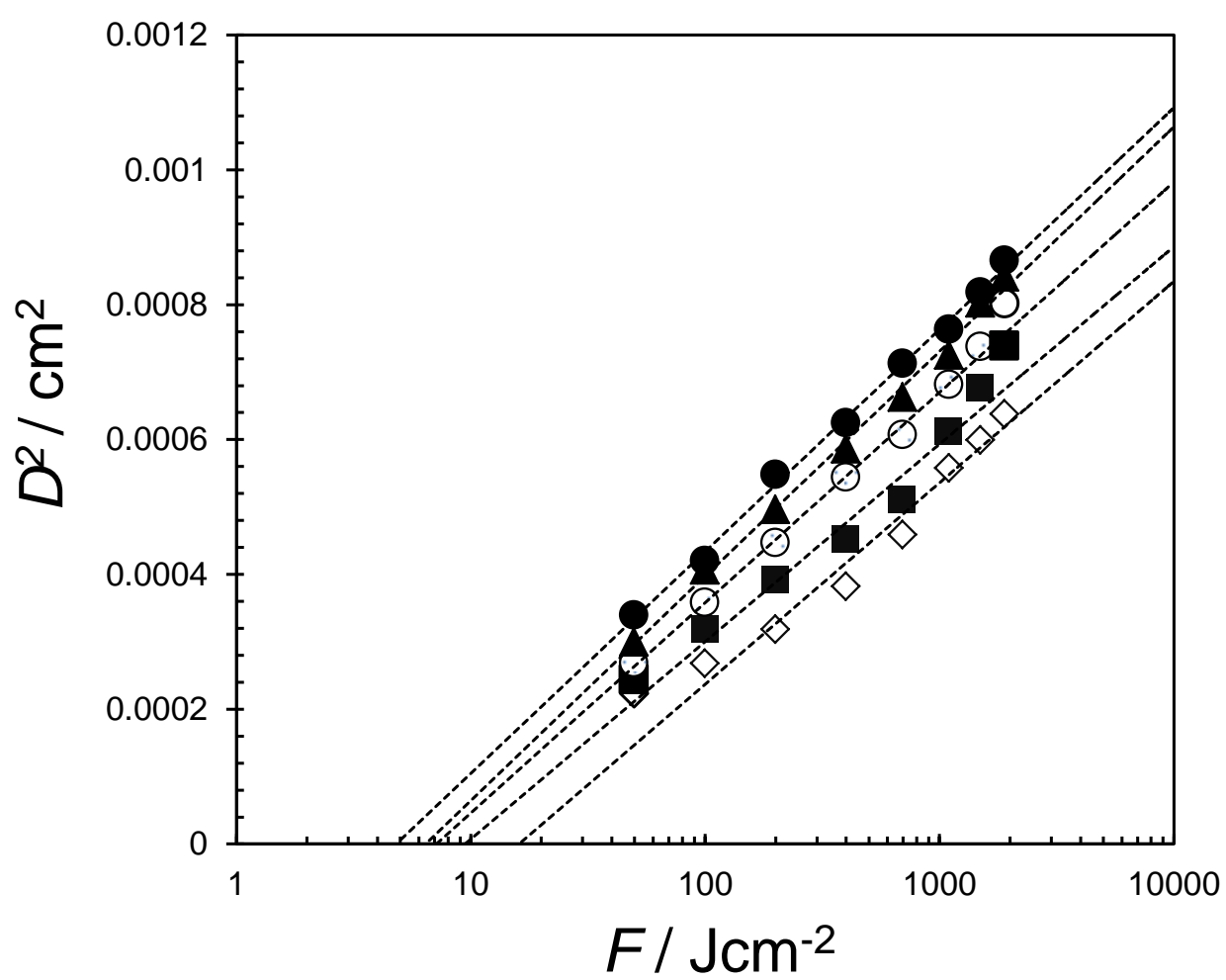


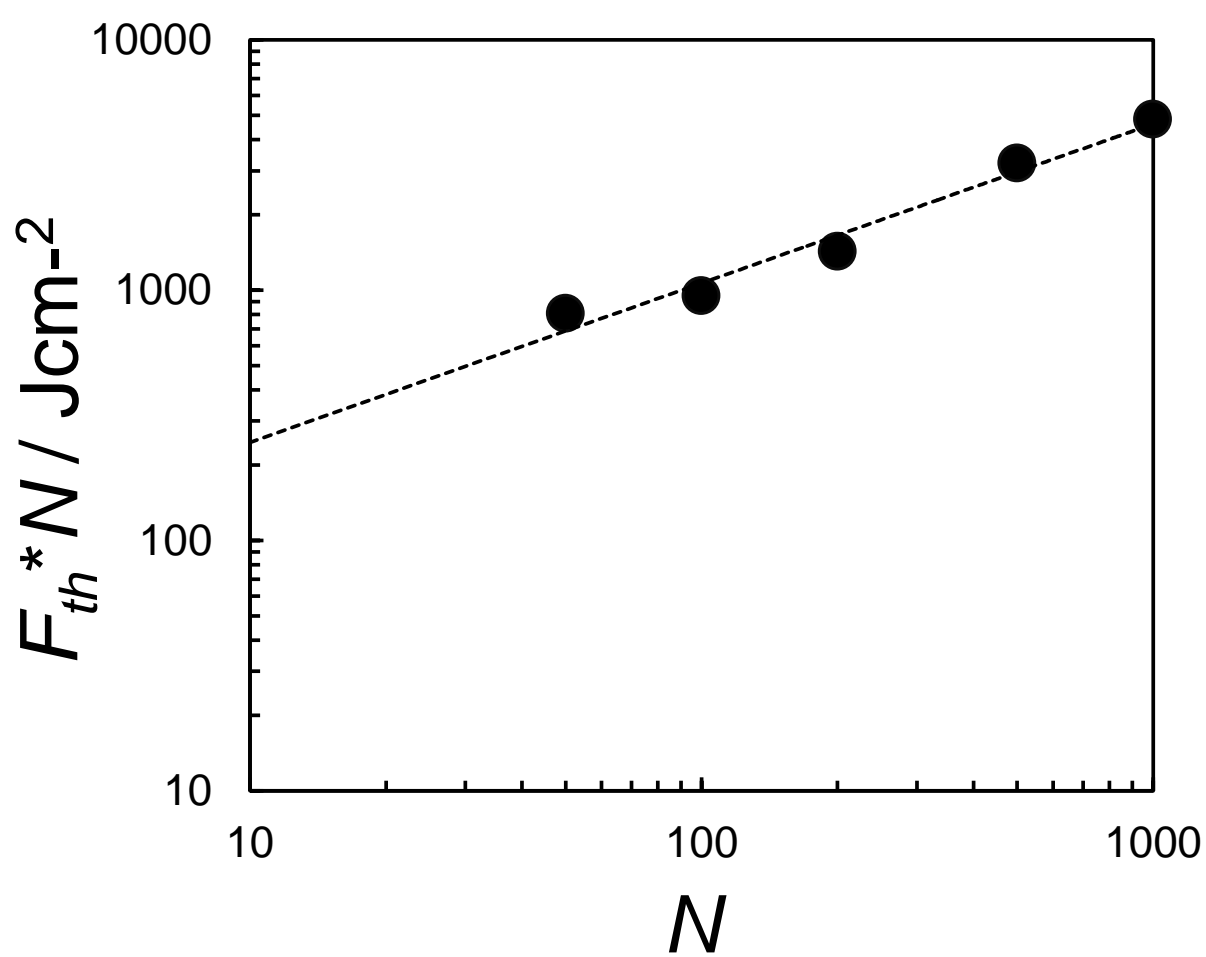


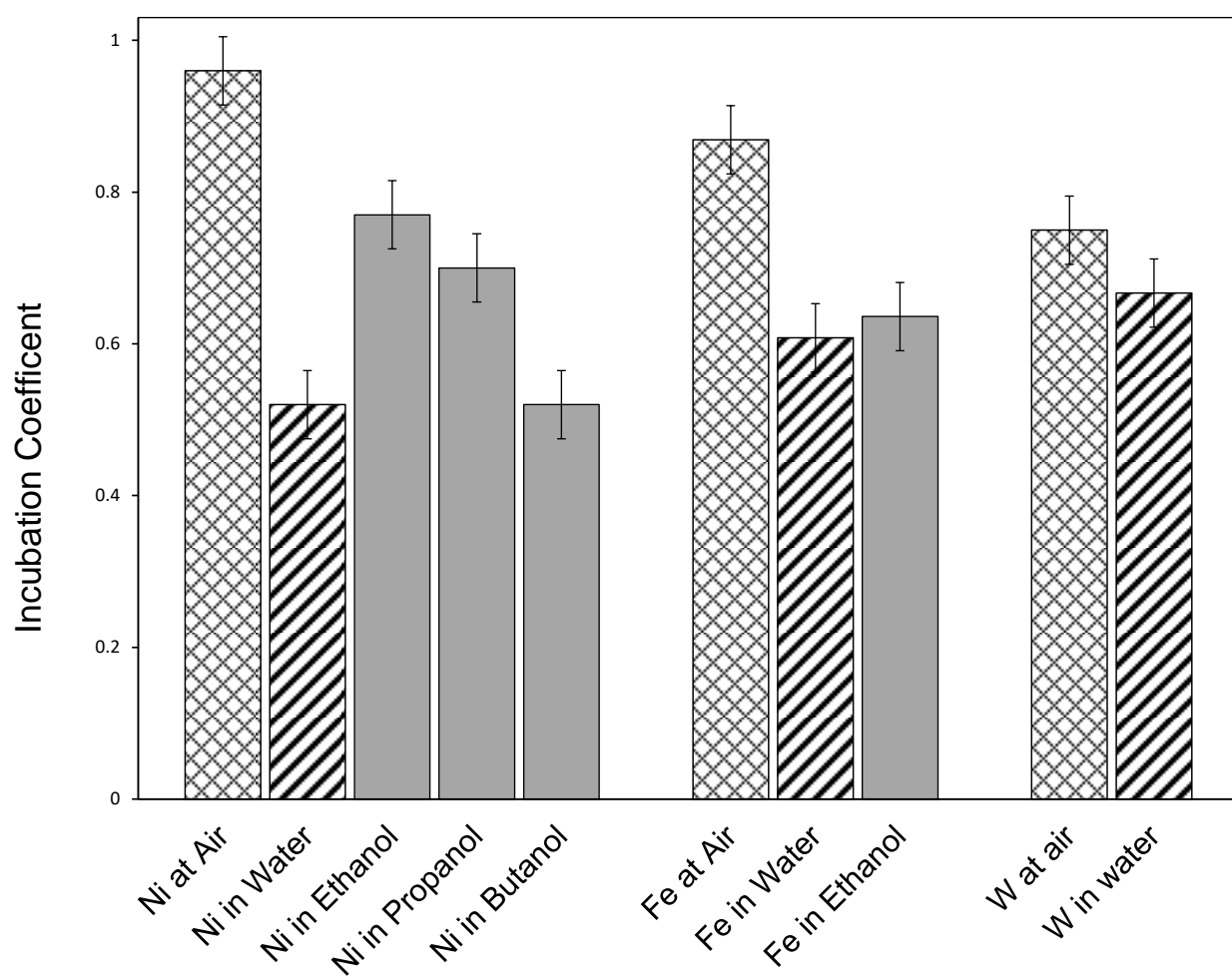


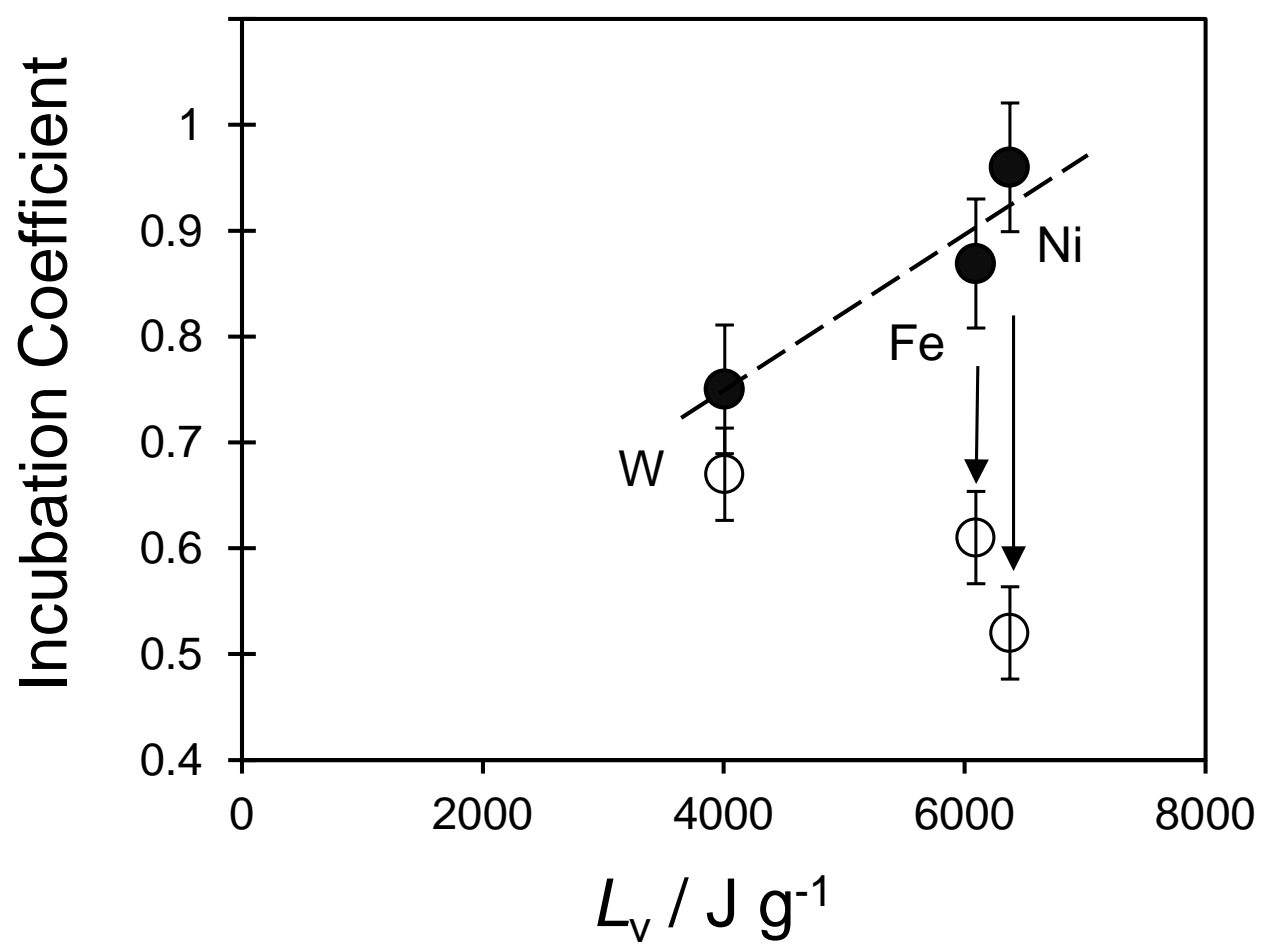












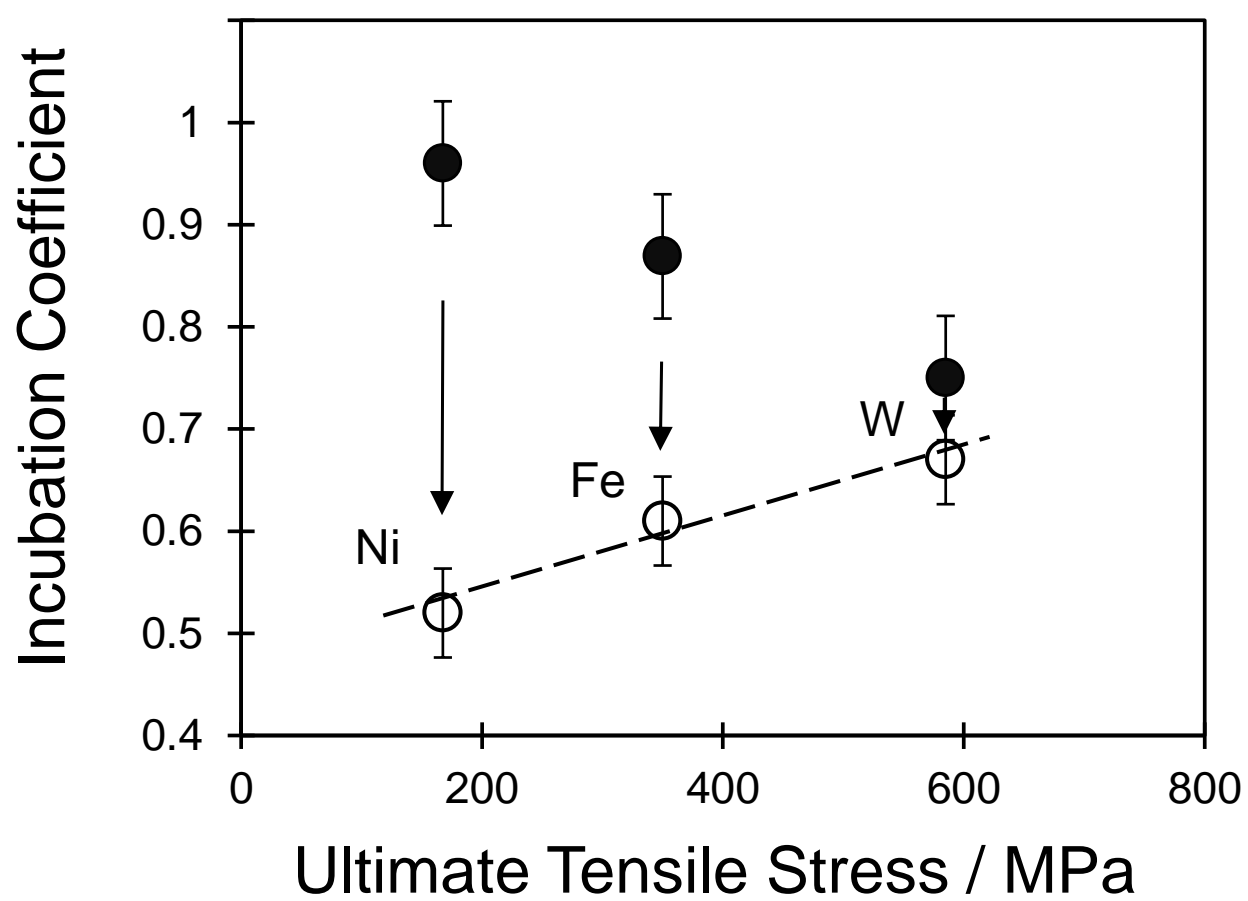


Table 1 Thermal and physical constants of various metals [65]. T_v : boiling point; T_m : melting point; c_p : heat capacity; κ : thermal conductivity; ρ : density; L_v : heat of vaporization; L_m : heat of melting; δ : heat diffusivity; R : reflectivity.

	T_v /K	T_m /K	c_p /JK ⁻¹ kg ⁻¹	ρ /g cm ⁻³	L_v /Jg ⁻¹	L_m /Jg ⁻¹	κ /Wm ⁻¹ K ⁻¹	δ /cm ² s ⁻¹	R_{532nm}
Ni	3005	1726	444	8.90	6378	292	91	0.23	0.60
Fe	3023	1808	444	7.87	6095	272	80	0.23	0.56
W	5933	3683	133	19.30	4009	192	173	0.67	0.49