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STUDY OF DYNAMIC ABSORPTIVITY AT 10.6 μm (CO_2) AND 1.06 μm (Nd-YAG) WAVELENGTHS AS A FUNCTION OF TEMPERATURE

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Abstract

One of the main advantages of Nd-YAG lasers compared to CO_2 lasers consists of easy beam handling using optical silica fibers. However, up to now Nd-YAG applications for material processing were limited due to a rather low available mean output power (less than about 500 W). A Eureka project (termed "EU-226") which consequently aimed at developing the so called "High Power Solid State Laser (HPSSL)" capable of 1 to 3 kW, involved absorptivity studies results of which are given in this contribution. Metallic surfaces exhibit a higher absorptivity at 1.06 μ m (Nd-YAG) than at 10.6 μ m which improves the energy transfer with a Nd-YAG laser and decreases the thermal runaway. This was acertained by measurements of absorptivity as a function of temperature and by thermal cycle calculations. This permits laser surface hardening of steels without any coating or gas shielding, which prevents in particular pollution. Various materials, i. e. pure nickel, Inconel 718 and AISI 1045 steel, were studied, featuring in particular the influence of surface roughness, oxidation and initial absorptivity on laser-material interaction.

Keywords

Laser, CO₂, Nd-YAG, Absorptivity, Surface Treatment, Thermal runaway, Steel, Nickel, Inconel 718, Eurêka.

Introduction

In laser surface treatments, the beam absorption can vary considerably depending on surface quality, temperature and environment. It is well known that metal surfaces show a great absorptivity increase when irradiated with a focused laser light. However, the exact mechanisms involved are not clear yet [1]. These would however be of a great interest to characterize the potential of new Nd-YAG lasers in high power systems capable of 1 to 3 kW and called "High Power Solid State Laser (HPSSL)" studied in Eureka project EU-226. Reliable data on absorption are therefore missing to accurately modelize the heat flow and field of temperature during for example laser hardening.

Two different methods for measuring the absorptivity respectively at 10.6 μ m (CO₂) and 1.06 μ m (Nd-YAG) were used. The first one, already used at I.S.L. (Institut Saint-Louis) [2] and E.T.C.A. [3], is based on temperature measurements; the second one, for 1.06 μ m, is optical using an integrating sphere [4].

Although laser processing is generally carried out with gas shielding, this preliminar study does not involve any gas protection and all measurements are done in air environment because careful control of the surrounding atmosphere is very difficult when using a mere nozzle.

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Materials and apparatus

Materials

Three materials (table I) with various surface roughnesses (table II) :

- AISI 1045 steel (0.46 wt % C) in the annealed state,

- Pure cast nickel,
- Conventinally heat treated Inconel 718.

Table I: C	Chemical	composition	(weight	percent).
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	Fe	Ni	Cr	Nb	C	Mn	Si	S	Р	Al	Mo	Ti	Co
Steel 1045	Bal.	-	-	-	0.466	0.33	0.19	0.01	0.01	-	-	-	-
Nickel	0.02	99.9	-	-	0.003	-	0.001	0.001	0.001	-	-	-	-
Inco. 718	18.1	Bal.	18.0	5.15	0.03	0.08	0.12	0.01	0.01	0.41	2.95	0.94	0.47

Table II : Surface roughnesses

 $(R_a : arithmetic average roughness, R_z : peak-to-peak roughness depth and R_m the maximal depth).$

		R _a (μm)	R _z (μm)	R _m (μm)
Steel 1045	ground	0.36 ± 0.06	2.25 ± 0.5	2.6 ± 0.5
	sandblasted	0.95 ± 0.04	7.80 ± 0.7	9.4 ± 0.9
	polished	0.02 ± 0.03	0.30 ± 0.08	0.3 ± 0.1
Nickel	ground	0.53 ± 0.08	3.60 ± 0.7	4.2 ± 0.8
	polished	0.01 ± 0.01	0.2 ± 0.1	0.3 ± 0.1
	ground	0.37 ± 0.05	2.65 ± 0.3	3.6 ± 0.4
Inconel 718	milled	2.80 ± 0.22	10.1 ± 2.2	10.5 ± 3.0
	polished	0.01 ± 0.01	0.15 ± 0.1	0.2 ± 0.1

For handling, steel is immersed in glycerine to prevent oxidation, which is not required for nickel and Inconel 718. All surfaces are carefuly cleaned before being irradiated.

Experimental

Thermal method at 10.6 μm (CO₂)



Figure 1 : Experimental setup used at 10.6 µm.

The source is a 5 kW Spectra-Physics cw-CO₂ laser with an unpolarized and multiple mode beam. An homogeneized top-hat beam is obtained using a kaleidoscope (figure 1). The parallelipedic sample ($10 \times 10 \times 1 \text{ mm}^3$) is put onto three tungstene pits, which minimize thermal losses. The absence of gas shielding for all experiments induces oxidation. The laser beam irradiates the whole square surface of the sample with a mean power density ranging from 5.10^6 W/m^2 to 9.10^6 W/m^2 . Temperature is monitored by a chromel-alumel thermocouple welded to the sample bottom.

Since A_i is used to calculate $T_c(t_i,e)$, iterations are required to obtain the best result. Consequently, absorption results are obtained from iterative solving of the onedimensional thermal problem using finite element calculations [3] which consist in minimizes the following functional :

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$$J(A_1,...,A_N) = \frac{1}{2} \sum_{i=1}^{N} [T_m(t_i,e) - T_c(t_i,e)]^2$$
(1)

with A_i absorptivity for [t_{i-1}, t_i], e sample thickness, T_m and T_c measured and calculated temperatures. In these calculations, variable thermal properties (i. e. thermal conductivty and specific heat) of the material are involved although these values are equilibrium one, which reduces the accuracy. Absorptivity results could be given as a function of temperature or time.

Reflectivity measurements at 1.06 µm (Nd-YAG)

A 250 W mean output power "Lasag" Nd-YAG laser with an unpolarized beam was used. The previouslydescribed thermal method cannot be used because of the low power and low repetition rate. Reflectivity measurements are therefore carried out with a sphere (figure 2) which integrates the reflected part of the beam. A single Nd-YAG laser pulse is focused on the sample. The pulse duration is 15 ms and the incident mean power density of about 5.10⁸ W/m².



Figure 2 : Experimental arrangement for the reflectance measurements at 1.06 µm.

The photodiode number 1 signal (reflected intensity) over the signal number 2 (incident intensity) gives, after previous calibration with a gilded molybdenum mirror, the reflectivity. Then, the absorptivity results from substracting the reflectivity to 1 because the tested materials are not transparent. However, the as measured absorptivity can given as a function of pulse duration since only be temperature is heterogeneous at the irradiated surface. Temperature increase has been calculated at the center and at the periphery of the impact using finite elements. This can be applied because the power density is known, the spatial intensity is multimode (as monitored with a CCD camera), the intensity versus time is recorded and finally the absorptivity is measured as a function of time. Figure 3 shows the results for sand blasted steel and give some indications about temperatures attained during these experiments which will be used in a further discussion.

Consequently, absorptivity could be estimated as a function of temperature ; first, by eliminating the time dependence between the experimental result A(t) and calculated T(t) ; second, by extrapolating this new dependence A(T) to the



Figure 3 : Calculated temperatures at center and periphery of the impact for sandblasted steel.

whole time range (outside [0,15 ms]). This will be used in the discussion.

Results [5]

AISI 1045 steel

Figure 4 shows the absorptivity measured at 10.6 μ m as a function of temperature and at 1.06 μ m as a function of pulse duration for a given power density noted π and mesured in W/m². The scatterband bars for 10.6 μ m measurements are obtained from the envelope of all curves corresponding to three or five experiments in the same conditions. At 1.06 μ m, the reproducibility always ranges in ± 0.03.



At room temperature, absorptivity is lower at $10.6 \mu m$ (0.08 to 0.20) than at $1.06 \mu m$ (0.30 to 0.52). Moreover, in both cases it greatly depends on surface roughness. It also increases with temperature due to surface oxidation. Roughness plays no part at 10.6 μm when temperature increases, contrary to what observed at 1.06 μm . This is probably due to the surface oxidation kinetics and optical differences between the two wavelengths.



Absorptivity measurements at $10.6 \,\mu\text{m}$ as a function of temperature and at $1.06 \,\mu\text{m}$ as a function of pulse duration are exhibited in figure 5. At room temperature, absorptivity is lower for $10.6 \,\mu\text{m}$ (0.04 to 0.10) than for $1.06 \,\mu\text{m}$ (0.30). When temperature increases, absorptivity does not vary very much for the two wavelengths, the surface oxide is probably transparent or very thin, which prevents any influence on the result.

Inconel 718

Figure 6 shows the absorptivity results for Inconel 718. At 10.6 μ m. Whatever the temperature, absorptivity remains almost the same, i.e. about 0.25 as mesured at a high temperature (1500 K). For 1.06 μ m, absorptivity is higher (0.30) and increases with temperature up to 0.55. The oscillations, systematically occured and are reproducible which is probably due to interference phenomenons in the developing oxide.



Discussion

Absorptivity

The differences between the methods used for the two wavelengths prevents easy direct comparison. Hoever, a significant comparison can be done using the extreme values, i.e. initial (at room temperature) and final (at elevated temperature) absorptivities (table III). One must also keep in mind that the duration of the irradiations, power density, thermal cycle and oxidation rate are very different for the two wavelengths.

			CC	Ο ₂ (10.6 μ	m)		Nd-YAG (1.06 μm)					
		Absorptivity		Temperature			Absorptivity		Temperature			
		(%)		(K)			(%)		(K)			
		initial	final	initial	final	$\Delta A/A_i$	initial	final	initial	final	$\Delta A/A_i$	
Steel	sand	17	80	400	1600	371	52	78	300	1240	50	
	grou.	8	80	350	1500	900 [°]	32	67	300	1090	109	
	polish.	7	75	350	1500	971	31	57 ·	300	1100	84	
Nickel	polish.	4	20	350	1550	400	32	52	300	980	62	
Inco.	polish.	22	20	500	1500	- 9	30	55	300	1180	83	

Table III : Absorptivity comparison at 10.6 and 1.06 µm.

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 $\Delta A/A_i$ is expressed as follows :

$$\frac{\Delta A}{A_{i}}(\%) = \frac{A_{\text{final}} - A_{\text{initial}}}{A_{\text{initial}}} \times 100$$
(2)

This ratio features the increase of absorptivity, which therfore gives an idea of the heat flow variation during irradiation. In the second part of the discussion, an illustration of the so called "thermal runaway" will be given for thermal cycles calculated using the previous absorptivity results.

For all surfaces (table III), absorptivity at room temperature is higher for the $1.06 \mu m$ light than for the $10.6 \mu m$ light. The difference seems to be promoted by a roughness decrease. Among the studied metals, Inconel 718 shows the smallest absorptivity increase when irradiating at $1.06 \mu m$. Regarding this property, the material order is then : nickel and steel.

As already mentioned, the absorptivity increase, quantified by $\Delta A/A_i$, can be used to feature the influence on thermal runaway. Roughness is generally a limiting factor, mainly due to a higher initial absorptivity as experimentaly ascertained. $\Delta A/A_i$ is much higher at 10.6 µm (CO₂) than at 1.06 µm (Nd-YAG) except for Inconel 718. It has to be related with the initial absorptivity because the maximum of $\Delta A/\Delta T$ is approximately the same for 1.06 and 10.6 µm (0.10 %/K).

Thermal cycles

Absorptivity measurements are then used to calculate thermal cycles for various heat flow inputs. Finite element calculations ("Zébulon" code) involve thermal properties as a function of temperature (thermal conductivity and specific heat) and a variable heat flow (i. e. the initial power density multiplied by the measured absorptivity). The motion of the laser beam is not taken in account. Figures 7-a and 7-b give respectively the thermal cycle at 10.6 and 1.06 μ m for the sand blasted steel with a pulse duration of one second. Thermal cycles for 1.06 μ m are calculated as mentioned earlier.



Figure 7: Thermal cycles for different deeps calculated with variable absorptivity versus temperature.

Thermal runaway (i. e. the heating rate suddenly increases) clearly appears above 900 K for 10.6 μ m, it is due to surface oxidation which induces a higher absorptivity level. For 1.06 μ m, this phenomenon does not occur, which suggests that surface treatment of steel without any coating (e. g. graphite) is easy to control, this has been experimentally ascertained. The shape of the thermal cycles obtained at 1.06 μ m (figure 7-b) is also appropriate from a metallurgical point of view because the holding time at an elevated temperature is higher than at 10.6 μ m. Cooling at the surface clearly shows a recalescence phenomenon which is not real but only due to the fact that the calculations used do not involve the martensitic transformation.

Conclusion

Absorptivities at two wavelengths 10.6 μ m (CO₂) and 1.06 μ m (Nd-YAG) for three materials (i. e. AISI 1045 steel, nickel and Inconel 718) were determined between room temperature and more than 1000 K in fairly realistic dynamic conditions. All experiments have been carried out in air without any gas shielding. Oxide layer formation increases the energy absorbed by the metal in particular for steel at 10.6 μ m. This induce a sudden growth of temperature, i.e. the so called "thermal runaway", and gives experimental difficulties to control the treatment. For surface hardening of steel with CO₂ laser, this phenomenon can be generally prevented by previous graphite spraying before processing.

Typical thermal cycles calculated for steel at the two laser wavelengths clearly show a difference regarding the thermal runaway. The Nd-YAG wavelength allows treatment without any coating or gas shielding, which prevents pollution and improves the reliability of the process compared to CO_2 treatments. This has been experimentally ascertained. In spite of this advantage and of the beam handling, Nd-YAG lasers generally operate in the pulsed mode which is not suitable for surface treatments.

This study clearly shows the prominent role of adequate measuring of absorptivity. A standard method remains to be defined. A combination of the thermal and optical methods used can be suggested.

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