	Original Research Article
Helium-Neon Lase Blood by Spe	r Effects on Human Whole ctroscopy in Vitro Study
ABSTRACT	
Low-power helium-neon laser recently ha FTIR and UV-Vis spectroscopic technique the serum of whole blood samples. Aims: To study (He-Ne) laser (λ = 632nm, irradiated to different times from 10 min to Study design: Human Whole Blood Irradi Place and Duration of Study: Institut technology (SUST), Soba Hospital, Kharto Methodology: Blood samples were co exposed to(H-N) laser and control comp used to study the effect of laser radiation. Results: Absorption spectrum and FTIR after He-Ne laser radiation shows, a signif exposed blood showed the peaks due to 0 group), and C-H (aromatic group). N-H (A transmittance in FTIR spectra for C=C transmittance were increased. The most He-Ne laser radiation for 10 and 20 min ar to denaturation of the protein. Conclusion: Photodegradation of blood causes changes in the structure and of	is been used numerously in medical applications. is employed to study the spectral differences in power=2mW) effect on human whole blood, after 50 min. iated to (He-Ne) laser(λ = 632nm, power=2mW) te of Laser, Sudan University of science and num-Sudan, February 2018. bllected from healthy volunteers; blood sample bared; UV-Vis spectrophotometer and FTIR were spectra of whole blood are compared before and ficant decrease in intensity. FTIR spectrum of non D-H (free group), C=O (amide I group), N=O (nitro Amino acid (amide II) Laser radiation changes in D group and O-H, N=O, the percentage of effects are found when whole blood irradiated to nd transmittance decreases for C-H, and N-H, due components due to absorption of laser radiation conformational changes in the polypeptide and

- Keywords: Laser, blood, UV-Vis, FTIR, spectroscopic

1. INTRODUCTION

Low-intensity helium-neon laser has been used extensively in medical applications.

- Interaction of lasers with biological materials such as blood, skin, and tissues is
- important to be understood. The study of blood change by spectroscopic techniques
- can be used for understanding the biological nature of the disease, and also for the
- diagnosis of the disease [1, 2].
- Photobiomodulations involves exposing tissues to low-level light. This type of
- therapy called Low-level laser therapy (LLLT), also known as cold laser therapy as

the power densities used produces no heating effect on the tissues. LLLT has a
photochemical effect which means the light is absorbed and cause a chemical
change. [3, 4, 5]

FTIR and UV-Vis spectroscopic technique are employed to study the spectral 26 27 differences in the serum of normal blood samples[2], blood samples were irradiated in that study by He-Ne laser (Wavelength λ = 632.8 nm, Power = 3mW). The FTIR 28 29 spectra for irradiated blood samples showed significant changes [1]. He Ne laser $(\lambda = 632$ nm, power=2mW) was used to irradiate human red blood cells and 30 31 investigated by absorption spectrum, FTIR and fluorescence spectra of RBC. The absorption spectrum of RBC after exposure to He-Ne laser shows a significant 32 decrease in absorbance. The FTIR spectrum of irradiated RBC clearly showed 33 34 changes in transmittance [6]. Some rheological factors of the human blood, such as complete blood count (CBC) parameters and blood sedimentation rate (BSR) 35 36 affected by low-level laser radiation (LLLR) laser blood biostimulation investigated the effect of LLLT on rheological parameters of human blood, they noticed a change 37 38 in both viscosity and size of erythrocytes [7,8]. Human blood exposed to low-39 intensity He-Ne-laser radiation causes clearly defined changes in the IR and visible 40 absorption spectra of the blood and erythrocytes. These spectral changes arise as a 41 result of partial photodissociation of haemoglobin-ligand [9]. 42 This paper investigates the effect of He-Ne laser (Wavelength λ = 632.8 nm, Power

43 = 2mW) with different exposure times using UV-Vis spectrophotometer and FTIR

44 spectrometer.

45 2. MATERIAL AND METHODS

46 **2.1 Samples Collection**

Blood samples were taken from healthy volunteers; 3 ml of each volunteer by medical standard laboratory conditions and blood samples were saved in a tube to prevent from coagulation to (EDTA) and each sample was divided into two samples one sample was control and other exposed to the helium-neon laser with different exposure times.

51 2.2 Laser irradiated

52 Samples were exposed to a Helium-Neon laser beam, operating in continuous wave mode, 53 as a radiation source (632.8 nm, 2 mW), for (10, 20, 30, 40 and 50) minutes The distance 54 between the laser source and the samples was set to be 10 cm and the diameter of a laser 55 spot was chosen to be 1.5 cm. To studied the effect of laser radiation were used UV-Vis 56 spectrophotometer (Jasco-670) and Fourier Transform Infra Red Spectra (FTIR) were 57 obtained used FTIR spectrophotometer (Shimadzu) for control, and He-Ne laser irradiated 58 blood serum samples. FTIR spectra of sera samples were recorded in the frequency range 4000 – 450 cm-1 on using Shimadzu at Central laboratory University of Khartoum,. IR transparent Thallium Bromide material without the serum was scanned as the background for each spectrum and 16 scans were co-added at a spectral resolution of 1 cm-1. FTIR spectra were obtained by spreading a small volume of serum on a Thallium Bromide plate (IR transparent material) and allowed to dry for few minutes to remove the water bands. To minimize problems from avoidable baseline shifts, the spectra were baseline corrected and normalized

66 UV -Vis spectra. Blood were diluted with normal saline and placed in Kartell disposable
 67 polystyrene cuvette of 10 mm path length. The cuvette is placed in Jasco-670)UV -Vis
 68 spectrophotometer for analysis The spectra were scanned in the region between 300nm to
 69 800nm using Jasco-670) at Laser institute laboratory, SUST,Khartoum

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71 3. RESULTS AND DISCUSSION

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3.1 UV-Vis spectra



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Figure .1 Spectrum of non- irradiated blood sample (control).

76 Figure 1 shows the spectrum of non- irradiated blood sample (control). This spectrum

referred to non- irradiated blood sample which specified by peaks at (576.0, 542.0, 416.0

78 and 340.0) nm with intensities 0.793, 0.755, 2.604 and 1.253 respectively.



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Figure2. Relation between Absorbance (a) and wavelength (λ) for whole blood before and after irradiated to (He-Ne) laser power 2 mW

The absorption spectra of the whole blood recorded in the range of 300–800 nm Figure 2. Contain absorption bands with λ_{max} = 340, 416 nm, a doublet band with λ_{max} = 542 and 576 nm. We investigated only those changes in the absorption spectra of the whole blood exposed to the (He-Ne laser) radiation that was detected for all of the samples studied.

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Table.1 The intensity of normal and irradiated samples

Wavelength (nm)	<mark>:h</mark>	Absorbance (a.u.)				
	control	10 min	20 min	30 min	40 min	50 min
340	1.253	1.01	0.933	1.065	1.12	0.868
416	2.604	2.49	2.391	2.501	2.538	2.347
542	0.755	0.633	0.536	0.614	0.699	0.492
576	0.793	0.653	0.547	0.633	0.718	0.525

Different serum samples are analyzed quantitatively by calculating the intensities among the absorption peaks which is show decrease intensity, all irradiated serum sample less than control serum sample. These results indicate to that there is photodegradation happened to the blood components. Laser radiation interacts with blood at the molecular level. Hemoglobin is a blood photoreceptor that selectively absorbed.

93 Absorption intensity slightly decreases for all peaks at, due to increasing ligand
 94 electronegativity [9].

In the UV-visible absorption spectrum of the irradiated blood, (figure.2 and table1) the
 most intense absorption band at 416 nm, the light with this wavelength that strikes this

97 biological tissues will be highly absorbed. This phenomenon is the key for the desired effect 98 on the tissues [10]. Figure 2 compared the light absorption at 340nm, 414nm, 542nm and 99 576nm for different irradiation time. The minimum light absorption occurred at 50 minutes of 100 irradiation with the fewer intensities recorded. The concentration of absorbing centers is 101 decreasing. This fluctuation of light absorption is known as biphasic response. The 102 mechanism of LLLT at cellular level has been associated with the absorption of 103 monochromatic visible and near infrared radiation. Effective tissue penetration is maximized 104 at specific optical window [11].

- 3.2 FTIR spectra 105
- 106 Table .2 FTIR spectral data (wave number, function group and transmission) for 107 normal blood control

FTIR spectral data for normal blood (control)					
Sr. No	Wave number	1/cm Group	% T		
1	3444.63	0-н	0.48		
2	1650.95	C=O	1.19		
		\sim			
3	1548.73	N=O	6.36		
4	1452.30	С-Н	14.26		
5	1317.29	N-H	15.3		
6	1168.78	C-0	17.12		

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109 Table .3 FTIR spectral data (wave number, function group and transmission) for

110 irradiated blood sample blood control

FTIR spectru	m of blood irradiate	d with he-ne laser fo	r duration 10, 2	0, 30,40 and 50
min	\sim			
Sr. No	Irradiated Time	Wave number	Group	Т%
	(minute)	1/CM		
1		3396.77	O-H	0.77
2	2 3 10 4	1650.96	C=O	1.78
3		1545.10	N=O	4.49
4		1450.73	C-H	15.20
5		1312.59	N-H	16.12
6		1161.74	C-O	18.70

7		3442.45	O-H	0.65
8		1651.63	C=O	1.68
9		1545.10	N=O	4.68
10	20	1451.01	C-H	11.43
11		1312.59	N-H	12.58
12		1161.74	C-0	13.76
13		3410.57	O-H	4.92
14		1651.63	C=O	6.50
15		1551.23	N=O	12.82
16	30	1451.01	C-H	22.14
17		1312.59	C-H / N-H	24.29
18		1167.96	C-0	26.31
19		3304.04	O-H	12.12
20		1645.41	C=O	13.21
21	40	1545.10	N=O	16.46
22		1447.23	C-H	25.43
23		1312.59	N-H	27.11

24		1161.74	C-0	28.50
25		3442.45	O-H	2.49
26		1651.63	N-H	6.46
27	50	1545.10	N=O	12.54
28	50	1451.01	C-H	22.28
29		1318.81	N-H	23.44
30		1167.96	C-0	25.59

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112 An FTIR spectrum of whole blood in vitro without laser radiation is shown in (Figure) 3. 113 Table2. Shows the groups OH, C=O, N=O, C-O and C-H in the region between the wave number 4000 cm⁻¹ to 500 cm⁻¹. The most intense absorption band in proteins is the amide I 114 peak, which is observed at 1650.95 cm⁻¹. Amide I is mainly associated with C=O symmetric 115 stretching and or C-O stretching vibrations. There are another very strong prominent amide 116 absorptions one at 1545 cm⁻¹ due to strong N-H in-plane bending and termed as an Amide II 117 118 band. The strong characteristic band at 3295 cm⁻¹ due to N-H symmetric stretching 119 confirmed the existence of amino acid group [2] The medium band at 2873 cm⁻¹ due to C-H asymmetric and symmetric stretching of CH3 group established the presence of lipids and 120 the medium bands at 2854 cm⁻¹ due to C-H symmetric stretching of CH2 group established 121 the presence of lipids, fatty acids[12,13,15,15]. The FTIR spectra of blood showed clear 122 123 bands at1080, and 12451 cm⁻¹, are composed of mononuclear cells containing nucleic acids such as DNA and RNA. The nucleic acid components found in WBCs [9]. The bands at 1170 124 125 cm^{-1} is associated with triglycerides of human blood. The band at 2936 cm^{-1} is related to 126 platelets due to -C-H symmetric stretching of -CH2[16]. 127 The whole blood sample is irradiated to He-Ne laser radiation for 10, 20, 30, and 40min. and 128 50 min duration respectively, figure (4 to 8) table 3. Shows the groups associated with 129 spectral peaks whole sample irradiated to He-Ne laser radiation for 10 min duration shows 130 an increase in transmittance for all groups except for C-H decreases due to the denaturation of the protein. FTIR spectra of whole blood irradiated with He-Ne laser for 20 minute show 131 132 decreases in transmission for group, C-H, and N-H, to denaturation of protein i.e. it breaks 133 the polypeptide bonds due to conformational changes of proteins, but in 30, 40, and 50 134 minutes show an increase in transmittance for all groups is observed the separate 135 chromophore and auxochrome groups (C=O, N=O, C-H, N-H, C-O, O-H) show also 136 significant changes and indicates a significant inecreasing in their concentration. Laser 137 irradiation of blood causes changes in absorption band in stretching and bending Vibrations

138 of peptide group.



139

140 Figure 3. FTIR spectra of irradiated blood by He-Ne laser for (0, 10, 20,30, 40 and 50)

minutes

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144 4. CONCLUSION

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This work had shown that laser radiation interacts with blood at the molecular level. 146 Hemoglobin is a blood photoreceptor that selectively absorbed He-Ne Laser beam with 147 output power 2mW, (632.8 nm). The absorption of laser beam by blood leads to partial 148 photodissociation. The results showed a decrease in intensity, all irradiated serum sample 149 intensity was less than control serum sample; this result indicates that there is 150 photodegradation happened to the blood components, this causes changes in the structure 151 and conformational changes in the polypeptide of N-H and CO and COO- groups in the 152 regions 1500-1700 and 3000-3500 cm-1 of the IR spectrum. Sample irradiated for 30, 40, 153 154 and 50 minutes show an increase in transmittance for all groups is observed the separate 155 chromophore and auxochrome groups (C=O, N=O, C-H, N-H, C-O, O-H) show also 156 significant changes and indicates a significant inecreasing in their concentration.

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