Original Research Article

Crude and Chitosan Nano-particles Extracts of Some Maggots as Antioxidant and Anticancer Agents

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6 Abstract: The antioxidant effects besides anticancer activities of Musca domestica, Lucilia 7 sericata and Chrysomya albiceps maggots extracts against human liver carcinoma (HepG-2) 8 and human colon carcinoma (HCT-116) were investigated. Two kinds of extracts, crude 9 and chitosan nano-particles (CNPs) were prepared. The antioxidant activity of different 10 tested extracts was performed by DPPH radical scavenging method, the results obtained 11 revealed that, the highest levels of DPPH scavenging activity were exhibited by the crude 12 extracts of tested maggots with preference to *C. albiceps* extract, which exhibited a much 13 more potent activity followed by *L. sericata* and *M. domestica* in crude and CNPs extracts. 14 Crude extracts have lower anticancer activity than the CNPs extracts; however, the lowest 15 percent of cell viability (6.7±0.7%) was recorded by L. sericata crude extract against 16 HCT-116, followed by *C. albiceps* crude extract (7.57±1.25%) against HepG-2 at the highest 17 used concentration 100 µg/ml. The strongest anticancer activity was observed with CNPs 18 extracts and it was recorded at concentrations of 80, 90 and 100 µg/ml against cell lines 19 tested. Depending on Median inhibitory concentrations (IC₅₀) of maggots crude and CNPs 20 extracts, the IC₅₀ values were in the range of 37.3 to 74.3 μ g/ml and the highest anticancer 21 activity was obtained by C. albiceps CNPs extracts against cell lines tested. In conclusion, 22 both tested extracts have optimistic antioxidant activity. CNPs extracts have great 23 therapeutic potential due to its anticancer inducing activities.

24 **Keywords**: Antitumor; Antioxidant; DPPH; *M. domestica*; *L. sericata*; *C. albiceps*; CNPs.

25 Introduction

Oxidative stress is caused by free radicals inducing many chronic and degenerative diseases including, heart disease, aging, diabetes and cancer [1]. Reduction of unstable and reactive free radicals can be achieved via antioxidants that protect cells from free radical attack. One of the main objectives of this study was to find natural origins antioxidants that replace synthetic antioxidants, which are limited by their carcinogenicity and have been suspected to cause negative health effects.

32 Despite considerable progress in medical research, cancer is still one of the 33 high-ranking causes of death in the world. It is the second most common cause of death 34 according to World Health Organization and by 2020 it will be cause death for more than 10 35 million people. Surgical therapy still promising and widely accepted cancer treatments, 36 much attention also received for nonsurgical cancer treatments that aimed to reduce 37 complications of surgical treatments. Also, cancer chemotherapy and radiations showed 38 serious side effects; therefore, it is important to find new, powerful anticancer agents that 39 are highly effective and biodegradable.

40 Maggot therapy has been traditionally practiced for debridement of necrotic wounds 41 as well as for curing infections at the wounds site; maggots promote wound healing, 42 stimulate granulation and promote the formation of human fibroblasts [2].

43 Insects offer a tremendous potential as a natural resource for chitin production. Even 44 chitosan is a derivative of chitin; it has its own unique functions, chitosan is a natural 45 nontoxic polysaccharide that have been widely used due to its various biological functions 46 such as antioxidant [3] and antitumor activity [4]. However, because of its high molecular 47 weight and water-insolubility, the application of chitosan is severely limited; therefore, 48 nanoparticle formulation enhancing therapeutic efficacy of chitosan [5]. CNPs exhibit more 49 superior activities than chitosan and have been reported to boost anticancer activity than 50 those of chitosan. In addition, nanoparticles possess a stronger surface curvature; this 51 produces more dissolution pressure with a corresponding increase in saturation solubility 52 [6].

53 Since not much data are available concerning the antioxidant and anticancer activities 54 of insects, especially flies' maggots, therefore, the objectives of the present study were to 55 evaluate the antioxidant and anticancer activity of the crude and CNPs maggots extracts of 56 *M. domestica, L. sericata* and *C. albiceps*.

57 Materials and Methods

58 1- Tested species.

59 Common species of medical importance in many parts of the world, including Egypt, 60 used in this study were *Musca domestica* (Diptera: Muscidae), *Lucilia sericata* and *Chrysomya* 61 *albiceps* (Diptera: Calliphoridae) maggots. They were obtained from the susceptible 62 laboratory-reared strains continuously raised in the institute of medical entomology, 63 Dokki, Egypt.

64 2- Tested extracts.

Two kinds of extracts, crude and CNPs extracts from each species with serial concentrations were prepared as the following:

67 2-1- Crude extracts preparation.

The extraction was performed according to [7] as the following: 3rd larval instar (100 larvae) were washed with 70% methanol and sterile double distilled water (ddH₂O) then incubated overnight at 30 °C, excess water was removed by using filter paper. Ten grams of each species was thoroughly homogenized. The homogenate was centrifuged at 13,000 rpm for 30 min. at 4 °C. After centrifugation the supernatants was decanted, filtrated with filter paper, dried in a rotary evaporator at 40 °C for 40 min. The dry extracts were weighed and dissolved in methanol and used as methanol extract.

- 75 2-2- CNPs extracts preparation.
- 76 Extracts *preparation* as CNPs was done as the following.

77 2-2-1- Extraction of chitin.

78 Chitin was isolated from the tested maggots as the following: 3rd larval instar (500 79 larvae) from each species were washed with 70% ethanol and sterile ddH₂O, dried for 48 h

80 and crushed with a mortar to create the maggots powder. The prepared maggots powder

81 were weighed, deproteinized using NaOH 2N with a ratio of 12ml/1g (v/w). The treatment 82 was carried out for 30 min., the residue was collected with filter paper, washed with 83 distilled water and dried in an oven at 50 °C. Deproteinized products were weighed; 84 demineralization was carried out with a diluted HCl solution 1M for 30 min at room 85 temperature [8]. Decolorization was done by treatment of the precipitate with 1% 86 potassium permanganate solution 100 ml for 1 h then, chitin was washed with distilled 87 water and dried at 50 °C.

88 2-2-2- Preparation and characterization of chitosan.

89 The product from decolorization was N-deacetylated using NaOH 12.5N with ratio of 90 1g/20ml (w/v). The residue was then washed with distilled water, collected with filter 91 paper and dried in oven at 50 °C. Infrared radiation by Fourier transform infrared 92 spectroscopy (FT-IR) with different wavelengths released on the sample was used for 93 chitosan characterization. An infrared spectrum represents a fingerprint of a sample with 94 absorption peaks. Spectrum formed showing the absorption and transmission of the 95 sample molecule. The spectrum is unique for the material as it has the unique combination 96 of atoms and no other compound can produce the same spectrum. FT-IR spectrum was 97 recorded on Jasco4100 spectrometer at Egyptian Petroleum Research Institute (EPRT). 98 Samples were prepared as potassium bromide (KBr) pellet and scanned against a blank KBr 99 pellet background at wave number range 4000–400 cm⁻¹ with a resolution of 4.0 cm⁻¹.

100 2-2-3- Preparation of CNPs.

The CNPs was prepared using a ball milling (RETSCH Planetary Ball Mills Type PM
400, Germany) at EPRT. Chitosan powder was charged and dry mixed into 250 ml stainless
steel agar with 8 grinding balls at 400 rpm for 8 hours.

104 2-2-4- Characterization of CNPs

105 *Transmission electron microscopy:* TEM was used to image the CNPs. The CNPs were 106 suspended in water for 3 min sonication to obtain a dilute suspension. A drop of this 107 suspension was deposited onto a glow discharged carbon-coated microscopy grid and 108 allowed to dry. The sample was investigated and imaged using Hitachi H-7000 TEM at 109 EPRT.



Fig. (1): TEM image of the CNPs.

113 The chitosan nanoparticles were spherical in shape and homogenously distributed 114 with a particle size lower than 50 nm (Fig. 1). The CNPs powder later was dissolved in 0.1% 115 acetic acid solution with a concentration ratio of 1g/100ml (w/v) for being applied as CNPs 116 extracts. Fresh solutions were only prepared when required.

117 *3- DPPH scavenging activity.*

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118 Free radical scavenging activity of different treatments were measured by 1,1-119 diphenyl-2-picryl hydrazyl (DPPH), 0.1 mM solution of DPPH was prepared. This solution 120 was added to 3 ml of each treatment. The mixture was shaken vigorously and allowed to 121 stand at room temperature for 30 min. then, absorbance was measured at 517 nm, using 122 spectrophotometer [9]. The IC_{50} value of the sample, which is the concentration of sample 123 required to inhibit 50% of the DPPH free radical, was calculated using Log dose inhibition 124 curve. Therefore serial concentrations were prepared for each treatment. Lower absorbance 125 of the reaction indicates higher free radical activity [10]. The percent of DPPH scavenging 126 effect calculated using the following equation: DPPH scavenging (%)= (A0-A1)/A0×100. 127 Where A_0 control absorbance and A_1 sample absorbance, the control used in this study was 128 the antioxidant and free radical scavengers Eugenol [11].

129 4-Cytotoxicity.

Human Liver Carcinoma cell line HepG-2, Human Colon Carcinoma cell line HCT-116 and skin normal human cell line (BJ-1) were obtained from VACSERA-Cell Culture Unit, Cairo, Egypt. These cell lines originally obtained from the American Type Culture Collection, and cultured in RPMI medium-1640 supplemented with 10 % inactivated fetal bovine serum (FBS). The reagents RPMI-1640 medium, SulphoRhodamine-B (SRB), dimethyl sulfoxide and fluorouracil (5-FU) were purchased from (Sigma Co., St. Louis, USA). Fetal bovine serum was obtained from (GIBCO, UK). The cell lines were used to

determine the inhibitory effects of different extracts on cell growth using the SRB assay.This colorimetric assay is based on the ability of SRB to bind to protein components of cells

139 that have been fixed in tissue culture plates by trichloroacetic acid (TCA).

140 5- Cytotoxicity screening.

141 The cancer cells were cultured in RPMI-1640 medium with 10% FBS. Antibiotics were 142 added 100 units/ml penicillin and 100µg/ml streptomycin at 37 °C in a 5% CO₂ incubator. 143 The cells were seeded in a 96-well plate at a density of 1.0×10^4 cells/well at 37 °C for 48 h in 144 incubator. After incubation, the cells were treated with crude and CNPs extracts and 145 incubated for 48 h, medium discard, fixed with 10% TCA 150 µl/well for 1 h at 4 °C and 146 washed 3 times by water. Wells were stained by SRB 70 µl/well for 10 min at room 147 temperature in dark place then washed with acetic acid 1% to remove unbound dye. The 148 plates were air-dried for 24 h. The dye solubilized with 50 µl/well of 10 mM tris-base (PH 149 7.4) for 5 min. The optical density of each well measured at 570 nm with an ELISA 150 microplate reader (EXL 800 USA). The relative cell viability in percentage was calculated as 151 (A570 of treated samples/A570 of untreated sample) X100 [12]. The IC₅₀ values were 152 calculated using multiple linear regressions [13]. The BJ-1 cells were used as a normal cell 153 model to compare HepG-2 and HCT-116 cells. The IC50 value or the 50% cytotoxicity was 154 determined from the linear equation obtained from the relation between the cell 155 cytotoxicity % and the concentrations tested.

156 *6- Statistical analysis.*

The statistical analysis of the obtained data was done according to [14, 15]. The analysis
was revised and graphics were drawn by SigmaPlot. The obtained data were assessed by
calculation of the mean (M), standard deviation (SD) and student t-test.

160 Results

161 1- Antioxidant activities

162 1-1- Crude extracts.

163 The antioxidant activity of maggots crude extracts of *M. domestica, L. sericata* and *C.* 164 *albiceps* and eugenol as a control were examined in the context of DPPH scavenging as a 165 representative of antioxidant activity. Data obtained in (Table 1) showed that, *C. albiceps* 166 extract exhibited the highest levels of DPPH scavenging activity; followed by *L. sericata* and 167 *M. domestica,* the IC₅₀ recorded 37.18, 72.28 and 81.5 µg/ml; respectively, compared to 4.05 168 µg/ml for eugenol.

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Table (1): IC₅₀ of DPPH scavenging activity of maggots crude extracts.

Samples	Linear equation	R ²	IC₅₀ µg/ml
M. domestica	Y= 0.5157× -3.1846	0.93503	81.5
L. sericata	Y= 0.7061× -3.1561	0.97068	72.28
C. albiceps	Y= 0.9267×+15.557	0.80864	37.18
Eugenol	Y=10.921× +5.5	0.9758	4.05

170 1-2- CNPs extracts.

171 The CNPs extracts of *M. domestica, L. sericata* and *C. albiceps* maggots and eugenol as a 172 control were examined for their antioxidant activity. Data obtained in (Table 2) revealed

173 that, *C. albiceps* extract exhibited the highest levels of DPPH scavenging activity; followed

174 by L. sericata and M. domestica, the IC50 recorded 103.13, 75.9 and 60.02 µg/ml for M.

175 *domestica, L. sericata* and *C. albiceps*; respectively, compared to 4.05 µg/ml for eugenol.

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Table (2): IC₅₀ of DPPH scavenging activity of maggots CNPs extracts.

Samples	Linear equation	R ²	IC50 µg/ml
M. domestica	Y= 0.6522× -3.1648	0.96485	103.13
L. sericata	Y= 0.6962× -2.864	0.97068	75.9
C. albiceps	Y= 0.7988× +2.054	0.80864	60.02
Eugenol	Y=10.921× +5.5	0.9758	4.05

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178 Comparing the results, on the basis of IC_{50} , *C. albiceps* extract exhibited the highest 179 levels of DPPH scavenging activity; followed by *L. sericata* and *M. domestica* for both crude 180 and CNPs extracts. The highest levels of DPPH scavenging activity were exhibited by the 181 crude extracts of tested maggots with preference of *C. albiceps*, which exhibited a much 182 more potent antioxidant activity than other tested species.

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184 *2- Anticancer activities*

185 2-1- Crude extracts.

186 The anticancer activity of *M. domestica*, *L. sericata* and *C. albiceps* maggots crude extracts 187 were examined against two human tumor cell lines. The anticancer activity was evaluated 188 by determining the cell viability average percentages of cancer cells in the test cultures. The 189 cell viability was variable among tumor cells tested. Data given in (Table 3, Figs. 2,3) 190 showed that, at the highest used concentration 100 µg/ml the lowest percent of cell viability 191 (6.7±0.7%) was recorded by L. sericata against HCT-116, followed by (7.57±1.25%) for C. 192 albiceps against HepG-2. The cell viability percent was decreased as the concentration used 193 increased. At the lowest concentration used 10 µg/ml, cell viability percentages recorded 194 (88.9±0.78; 95.8±0.43%) for L. sericata and M. domestica; respectively against HCT-116, and it 195 was 93.3±0.58% for C. albiceps against HepG-2 tumor cell line. BJ-1 was almost inactive at 196 the highest concentration tested.

197 198 Table (3): Cytotoxicity effect *M. domestica, L. sericata* and *C. albiceps* maggots *crude extracts* against liver and colon carcinoma cell lines.

	Cell Viability average percentages %					
Concentrations µg/ml	HepG-2 cell line			HCT-116 cell line		
	M. domestica	L. sericata	C. albiceps	M. domestica	L. sericata	C. albiceps
100	25.9±0.9	14.43±1.5	7.57±1.25	30.1±0.11	6.7±0.7	14.1±1.0
90	37.5±0.62	19.2±1.06	11.17±1.25	33.37±0.4	10.4±0.5	19.87±0.2
80	42.17±1.12	25.2±1.59	19.07±1.16	42.03±1.82	13.93±1.0	23.87±0.8
70	51.9±1.97	36.1±1.04	29.4±0.58	52.8±1.4	26.1±1.2	36.8±0.2
60	66.07±0.2	49.57±1.4	44.7±1.4	60.13±0.23	38.5±0.6	51.23±1.6
50	70.9±1.8	61.93±1.7	53.6±1.7	69.63±0.4	51.46±1.4	63.27±0.64
40	81.97±2.0	77.0±2.6	62.67±3.8	81.93±1.95	62.7±3.8	79.0±1.0
30	87.8±0.72	95.7±2.2	69.07±1.2	86.63±1.6	69.63±1.58	94.3±1.2
20	100.0±0.0	98.78±0.0	89.67±1.5	91.43±0.47	75.3±2.0	97.8±0.3
10	100.0±0.0	100.0±0.0	93.3±0.58	95.8±0.43	88.9±0.78	100±0.0
0.0	100	100.0	100.0	100.0	100.0	100.0
BJ-1	92.9±2.01	91.8±2.5	89.3±1.3	88.9±4.7	88.9±1.1	98.0±0.8





Fig. (2): Cytotoxicity curve of *M. domestica, L. sericata* and *C. albiceps* maggots crude extracts against liver carcinoma cell line.



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Fig. (3): Cytotoxicity curve of *M. domestica, L. sericata* and *C. albiceps* maggots crude extracts
 against colon carcinoma cell line.

205 *2-2- CNPs extracts.*

206 The anticancer activity of maggots CNPs extracts of M. domestica, L. sericata and C. 207 albiceps were examined against HepG-2 and HCT-116 cell lines. Data given in (Table 4, Figs. 208 4,5) showed that, the cell viability percent was decreased as the concentration increased, the 209 CNPs extracts revealed optimistic results in both tested cell lines, and it was highly 210 effective. The cell viability ranged from (77.18±0.3) to (93.0±1.5) at the lowest concentration 211 used 10 µg/ml and it was concentration dependent. Meanwhile, the highest anticancer 212 activity was recorded at the concentrations of 80, 90 and 100 µg/ml of the different tested 213 CNPs extracts against HepG-2 and HCT-116 cell lines tested, where the cell viability was at 214 its lowest recorded values. BJ-1 was almost inactive at the highest concentration tested. 215

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222Table (4): Cytotoxicity effect *M. domestica, L. sericata* and *C. albiceps* maggots CNPs extracts223against liver and colon carcinoma cell lines.

	Cell Viability average percentages %					
Concentrations µg/ml	HepG-2 cell line			HCT-116 cell line		
	M. domestica	L. sericata	C. albiceps	M. domestica	L. sericata	C. albiceps
100	1.87±0.35	1.99±0.3	1.6±0.36	1.87±0.11	1.4±0.23	1.61±0.26
90	4.3±0.36	4.3±0.47	4.03±0.15	7.3±0.2	7.5±0.18	7.9±0.3
80	9.07±0.45	9.0±0.26	9.6±0.67	10.9±0.3	11.3±0.3	11.1±0.11
70	14.4±0.47	14.8±0.26	14.1±0.15	15.53±0.5	14.7±0.3	13.2±0.41
60	21.6±0.49	19.1±0.95	17.7±1.1	22.6±0.5	20±0.7	18.87±0.5
50	28.73±0.95	31.6±1.3	27.2±1.4	32.7±0.4	30.4±0.8	28.4±1
40	43.7±1.15	45.0±0.57	39.6±0.58	43.9±1.1	41.3±0.5	40.3±0.2
30	52.0±0.46	55.0±1	49.8±0.46	56.35±0.6	55.9±0.9	53.27±0.8
20	73.6±1.8	75.0±0.2	73.0±1.2	70.9±1.1	69.9±0.3	66.8±0.43
10	89.43±0.5	93.0±1.5	86.0±0.58	88.6±0.5	85.9±0.5	77.18±0.3
0.0	100.0	100	100	100	100	100
BJ-1	96.9±2.02	95.8±1.1	96.3±1.3	98.9 ± 1.7	98.9 ± 3.6	98.0±5.0









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231 Comparing the results, the cell viability was concentration dependent and it was 232 highly affected by the CNPs treatments. The highest anticancer activity was recorded at the 233 concentrations of 80, 90 and 100 μ g/ml of different tested CNPs extracts against HepG-2 234 and HCT-116 cell lines.

	IC ₅₀ Concentrations µg/ml ± SD					
Tested species	Crude e	extracts	CNPs extracts			
	HepG-2	HCT-116	HepG-2	HCT-116		
M. domestica	74.3±4.2	73.2±3.2	40.1±4.6	40.95±2.6		
L. sericata	61.4±3.1	49.4±3.2	41.3±2.1	39.7±4.1		
C. albiceps	52.8±4.8	61.8±4.1	38.5±4.8	37.3±2.1		
Fluorouracil (5-FU)	28.3±2.1	19.8±2.6	28.3±2.1	19.8±2.6		

235Table (5): IC50 values of *M. domestica, L. sericata* and *C. albiceps* maggots crude and CNPs236extracts against human Liver and Colon carcinoma cell lines.

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238 Median inhibitory concentrations of *M. domestica, L. sericata* and *C. albiceps* maggots 239 crude and CNPs extracts against tumor cells tested are summarized in Table (5). The IC₅₀ 240 values were in the range of 37.3 to 74.3µg/ml. The highest anticancer activity was obtained 241 by *C. albiceps* CNPs extracts against cell lines tested when compared to the anticancer agent 242 fluorouracil (5-FU).

243 Discussion

A primary component of insect cuticle is chitin; therefore, insects are an alternative chitin and consequently chitosan source. The production of chitin from insect has drawn increased attention because insects possess enormous biodiversity and represent 95% of the animal kingdom. Furthermore, insect cuticles have lower levels of inorganic material compared to crustacean shells, which makes their demineralization treatment more convenient [16].

Chitosan and CNPs are biopolymers that have unique structural possibilities for chemical and mechanical modifications to generate novel properties and functions. These biopolymers are biocompatible, biodegradable and nontoxic, and their chemical properties allow them to be easily processed, due to these unique properties, they are excellent candidates for cancer cure or cancer diagnosis [17].

255 The antioxidant effect of chitosan has been documented in several reports; the 256 protective role of chitosan nanoparticle against oxidative stress in rat model was studied 257 [17]; the antioxidative effect of chitosan on chronic hepatic injury in rats was also 258 investigated [18] and the authors found that chitosan has strong antioxidative effects. 259 Results obtained in this study may be in harmony with the previous findings, where high 260 levels of DPPH scavenging activity were exhibited by both crude and CNPs extracts with 261 preference to crude extracts which exhibited a much more potent antioxidant activity; 262 followed by the CNPs extracts, indicating the overall antioxidant activity maybe due to 263 antagonistic effect of free radicals by its antioxidant nature.

264 CNPs have been widely used due to its biological functions and antitumor activity; in 265 this study the CNPs extracts exhibited much more anticancer activity against cell lines 266 tested than crude extracts, that might be due to a difference in the mechanism of 267 cytotoxicity. The stronger cytotoxic effect of CNPs on tested cell lines might be related to 268 the highly positive charged amino group that attracted to the cancer cell membrane that 269 had a greater negative charge than normal cells. These results may be in harmony with [19] 270 who found that chitosan display notable antitumor activity against sarcoma tumors in 271 BALB/C mice, and [20] against Meth-A solid tumor in BALB/C mice.

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273 Conclusion

The crude and CNPs extracts induced antioxidant and anticancer activities; the highest antioxidant activity was induces by *C. albiceps* extract as demonstrated by DPPH scavenging activity. The cell viability was concentration dependent, crude extracts have lower anticancer activity than CNPs extracts. The highest anticancer activity was recorded at concentrations of 80, 90 and 100 μ g/ml of different CNPs extracts tested against HepG-2 and HCT-116 cell lines. Both tested extracts have optimistic antioxidant activity. CNPs extracts have great therapeutic potential due to its anticancer inducing activities.

281 Conflict of Interest

The author declares that there is no conflict of interests regarding the publication of thisarticle.

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