



Ain Shams University
Faculty of Science
Chemistry Department



Control of mycotoxin using nanopolymers prepared from natural sources

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By

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M.Sc. Science in Organic Chemistry (2013).

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بِسْمِ اللَّهِ الرَّحْمَنِ الرَّحِيمِ

﴿وَقُلْ أَعْمَلُوا فَسِيرَی اللَّهِ عَمَلَكُمْ وَرَسُولِهِ
وَالْمُؤْمِنُونَ وَاسْتَرُدُّونَ إِلَى عَالَمِ الْغِیْبِ
وَالشَّهَادَةِ فَيُنَبِّئُكُمْ بِمَا كُنْتُمْ تَعْمَلُونَ﴾

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Content

<i>Content</i>	<i>Page</i>
Acknowledgment	I
List of figures	III
List of tables	VIII
Abbreviations	IX
Papers discussed	
Abstract	XII
Summary	XIV
I. Introduction	1
II. Review and Literature	5
Aim of the work	30
III. Material and methods	31
1. Chemicals and kits	31
2. Preparation and characterization of Glucan Mannan Lipid Particles (GMLPs)	32
3. The chemical composition of yeast wall	35
4. Development of an <i>in vitro</i> method to measure AFB ₁ binding	35
4.1. Aflatoxin preparation	35

4.2. Gastrointestinal model preparation	36
4.3. Incubation conditions for optimization the adsorption assay of AFB ₁ incubation conditions.	36
4.4. Comparison of yeast cell wall-based particulate materials with different beta-glucan, mannan, lipid and chitin contents on AFB ₁ adsorption.	37
4.5.The binding efficacy GMLP on AFB ₁ adsorption	37
5. GMLP Microencapsulation of mycotoxin binding materials (payload) to enhance AFB ₁ binding.	37
5.1. Polyvinylpyrrolidone (PVP) as mycotoxin binder.	37
5.1.1. Preparation PVP/Tannic acid nanoparticle complexation as a nanoparticulate payload (PVP-TA NPs).	38
5.1.2. <i>In vitro</i> assessment of PVP/ TA complexed NPs as aflatoxin binders.	38
5.1.3. Synthesis of fluorescent TA by using 5-(4, 6-dichlorotriazinyl) aminofluorescein (DTAF) as a trapping agent. 100 mg of tannic acid was dissolved in 10 mL	38
5.1.4. Synthesis of PVP--NPs inside GMLPs (GMLP PVP-NP formulation).	38
5.1.5. Characterization of GMLP/ PVP and GMLP- PVP TA-NP formulations.	40
5.1.6. AFB ₁ binding capacity of GMLP encapsulated	41

DMwt PVP /TA NPs (GMLP/PVP-TA NPs formulation)	
5.2. Humic acid (HA)	42
5.2.1. HA-metal nanoparticle complexation as a nanoparticulate payload HA NPs) using chitosan (CS), Polyethylenimine (PEI) and metal ion.	42
5.2.2. Synthesis of fluorescent Chitosan using fluorescein isothiocyanate (FITC-CS).	43
5.2.3. Synthesis of fluorescent PEI using fluorescein isothiocyanate (FITC-PEI).	43
5.2.4. Synthesis of HA-complexed-NPs inside GMLPs (GMLP HANP formulation)	44
5.2.5. Characterization of GMLP HA and GMLP HA-NP formulations	44
5.3. Diamond nanopowder as mycotoxin sequestering agent (payload).	45
5.3.1. Dispersion Nano-diamond particles (ND)	45
5.3.2. Introducing ND as payload molecules into the GMLP to form a core	45
5.3.3. Synthesis of ND-NPs inside GMLPs (GMLP ND/NP formulation)	46
5.4. Super Activated Porous Carbon NPs (carbon nanopowder) as mycotoxin sequestering agent.	46
5.4.1. Saturated GMLP cell wall with Congo red dye	47

(CR).	
5.4.2. Loading activated carbon onto GMLP /CR surface	47
6. Cytotoxicity assay of GMLP-HA and GMLP-HA-AFB1 formulations.	48
7. <i>IN VIVO</i> studies.	49
7.1. Experimental animals.	49
7.2. Experimental Design.	49
7.3. Serum biochemical assay.	50
7.4. Histopathological study.	51
8. Statistical analysis.	51
IV. Results and discussions	52
1. Optimization of AFB ₁ adsorption assay	52
2. The effect of yeast cell wall composition on AFB ₁ adsorption	53
3. The binding efficacy of GMLP on AFB ₁ adsorption	55
4. GMLP Microencapsulation of mycotoxin binding materials to enhance AFB ₁ binding	56
4.1. Polyvinylpyrrolidone (PVP)	57
4.1.1. PVP/TA NPs as a nano-particulate payload (PVP-TA NPs)	57
4.1.2. GMLP encapsulated PVP/TA NPs as a nanoparticulate payload (GMLP/ PVP-TA NPs).	59

4.1.3. AFB ₁ -binding properties of free and encapsulated (PVP 360KD-TA NPs)	64
4.1.4. AFB ₁ -binding properties of free and encapsulated (PVP 1300KD-TA NPs)	66
4.2. Humic acid (HA)	70
4.2.1. Characterization of aflatoxin-binding properties of unencapsulated free HA and HA complexes nanoparticles (NPs)	73
4.2.2. GMLP encapsulated HA NPs (GMLP HA-NPs)	74
4.2.3. AFB ₁ binding capacity of GMLP encapsulated HA complexed NPs.	77
4.3. Nanaodiamond (ND)	80
4.3.1. Nano-diamond particles Dispersion (ND	80
4.3.2. GMLP encapsulated ND-PEI complexed NPs (GMLP/ND-PEI).	80
4.3.3. AFB ₁ binding capacity of GMLP encapsulated ND.	83
II. Incorporation large NPs (MB complexes) onto GMLPs Surface	85
4.4. Super Activated Porous Carbon NPs (carbon nanopowder) as mycotoxin sequestering agent.	
4.4. 1. Saturated GMLP cell wall with Congo red dye (CR).	85
4.4. 2. Incorporation NC on GMLPs Surface (CR-NC	85

CONTENT

GMLP).	
4.4.3. AFB ₁ binding capacity of GMLP/CR-NC	88
5. <i>In vitro</i> cytotoxicity of GMLP HA-NP-AFB ₁ complexes	90
II. <i>IN VIVO</i> studies	93
a. Body weight and food intake	93
b. Serum biochemical parameters	97
c. Histopathological Examination	109
1. The histological examination of the liver	109
2. The histological examination of the kidney	112
V. References	117
Arabic Summary	

List of figures

Figure	Title	Page
Figure 1	Chemical structure of the most common aflatoxins	9
Figure 2	Aflatoxin B ₁ metabolism in liver	10
Figure 3	Poly vinyl pyrrolidone (PVP) structure	
Figure 4	Example of a typical humic acid, having a variety of components including quinone, phenol, catechol and sugar moieties.	23
Figure 5	Schematic representation showing the different types of glucan particles	32
Figure 6	Production of Glucan Mannan Lipid Particles	33
Figure 7	a) Sealing of a small pipette tip (1–20 μ L) with a Bunsen burner, (b) sealed pipette tip for mixing of GMLP samples, (c) image of a mixing stick device, and (d) mixing of GMLP samples and solution into a thick paste using the mixing device inserted into the sealed pipette tip.	40
Figure 8	High content imaging was acquired from Image Xpress Micro (on the left). Fluorescent photomicrographs were captured by Axiovert 200 M (on the right).	41
Figure 9	AFB ₁ degradation in SGF and SIF.	55
Figure 10	The calibration curve of AFB ₁ in SGF after 10 min and SIF after 1h using HPLC. The values are expressed as mean ($\pm$ standard deviation)	56
Figure 11	The absorption efficacy of AFB ₁ by different yeast cell wall particulate materials in SGF after 10 min and in SIF after 1h.	57
Figure 12	The binding of AFB ₁ by GMLP in SGF after 10 min and in SIF after 1h.	58
Figure 13	The mass ratio required to achieve efficient complexation of 1 mg TA using (DMwt of PVP 10, 360 and 1300 kDa), the concentration of the former complex depends on the turbidity values at 600nm.	
Figure 14	Schematic representation of PVP (10, 360 and	62