Facile Route for Synthesis of Novel Flame Retardant, Reinforcement and Antibacterial Textile Fabrics Coatings

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Received: 31 May 2020; Accepted: 17 June 2020; Published: 19 June 2020

Abstract: New and innovative textile fabrics coatings were facilely developed. The coating was developed based on synthesis of novel charring and antibacterial organic agent in conjunction with chitosan. N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid was synthesized as organic antibacterial, reinforcement, and charring agent (OA) and then, dispersed in chitosan solution followed by coating on textile fabrics using immersion route forming new flame retardant coating layer. The developed organic molecule structure was elucidated using spectroscopic techniques. The mass loadings of developed organic agent dispersed in chitosan solution were varied between 20–60 wt.%. The coated textile fabrics have special surface morphology of fiber shape aligned on textile fibers surface. The thermal stability and charring residues of the coated textile fabrics were enhanced when compared to blank and organic agent free coated samples. Furthermore, the flammability properties were evaluated using LOI (limiting oxygen index) and UL94 tests. Therefore, the coated textile fabrics record significant enhancement in flame retardancy achieving first class flame retardant textile of zero mm/min rate of burning and 23.8% of LOI value compared to 118 mm/min. rate of burning and 18.2% for blank textile, respectively. The tensile strength of the coated textile fabrics was enhanced, achieving 51% improvement as compared to blank sample. Additionally, the developed coating layer significantly inhibited the bacterial growth, recording 18 mm of clear inhibition zone for coated sample when compared to zero for blank and chitosan coated ones.

Keywords: flame retardants; thermal stability; antibacterial; coatings; textile fabrics

1. Introduction

Textile-based materials have been used extensively in a variety of applications due to their interesting properties [1]. However, the demand of the textile fabrics for some applications require new smart functions to be included, such as high thermal stability, flame retardancy, and inhibition of the bacterial growth [2–4]. Various materials have been used for improving the thermal stability, flame retardancy, and antibacterial behavior of textile fabrics while using different nanoparticles [5–7]. However, the tensile strength of the treated textile fabrics was negatively affected due to the inclusion of nanoparticles inside fibers [8]. Therefore, organic agents were synthesized and used as a capping agent for synthesis of nanoparticles to avoid tensile strength failure [8]. Previously, many organic phosphates compounds have been used for imparting thermal stability and flame
retardancy properties to textile fabrics [8–10]. Therefore, there are several studies that have been reported for the incorporation of flame retardancy and antibacterial functions for textile fabrics via different textile treatments processes [4,5,10–12]. On the other hand, a variety of spherical metal nanoparticles have been reported for enhancing antibacterial properties of textile fabrics, such as silver nanoparticles (AgNPs), TiO₂NP, and ZnO; however, the tensile strength properties of the treated textile fabrics were reduced [7,9,13,14]. This is due to the negative impact of treatment process conditions, such as high acidic medium and higher temperatures processing [7,13]. Thus, the design and synthesis of novel and potential antibacterial, flame retardant, and reinforcement coating materials for textile fabrics are a much appreciated and required research trend. On the other hand, derivatives of 2-substituted-4H-3,1-benzoxazin-4-one have been attracted significant attention from chemists and pharmacologists because of their broad spectrum in biological activities and antibacterial applications [15–18]. One of the most important features of 4H-3,1-benzoxazinones chemistry is their use as key starting materials for further transformations in the design and synthesis of biologically active compounds [15–21]. The synthetic and biological importance of 4H-3,1-benzoxazinones stemmed from their facile ring-opening of the oxazinone ring with different nucleophiles to give acyclic products, which undergo heterocyclization to give other synthetically and biologically useful compounds. Chalcones are also known as an important class of biologically active compounds [22–24]. Hence, the synthesis of new charring and antibacterial molecules for imparting good flame retardancy and antibacterial properties for textile fabrics is necessity. Therefore, here, a new organic charring molecule was developed and exploited for the first time with chitosan as textile fabric coating and exhibited excellent antibacterial, flame retardant, and reinforcement properties for coated textile fabrics that are superior than previous reported materials [7,8,10]. It is noteworthy to note that several methods were used for the coating of textile fabrics using coatings, such as sol-gel [25], layer-by-layer [25], back coating [10], and pad-dry-cure (immersion) [13]. However, the pad-dry-cure method is a much more convenient and cost-effective treatment route when compared to the above-mentioned methods in terms of less reagents being used and short steps acquired for treatment [14,26,27]. It is important to note that our group has long been involved in the development of different antibacterial, flame retardant, and thermal stability materials for different textile fabrics [4,8,10,28,29]. In this study, N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid was developed and elucidated as active antibacterial, flame retardant/charring, and reinforcement agent. The developed material was used with chitosan for coating textile fabrics. The mass loading of developed organic molecule was varied and optimized. The surface morphology of uncoated and coated textile fabrics was studied. The tensile strength, thermal stability, flammability, and antibacterial properties were investigated.

2. Experimental Section

2.1. Materials

Commercial final (dyed) furniture textile fabrics composed from polyester/microfilament/fibran/wool 35%/25%/25%/15%, respectively, were purchased from Al Mahalla Co., Algharbia, Egypt. Anthranilic acid and all chemicals and solvents used were purchased from El Gomhoria Company of chemicals, Cairo, Egypt. Deionized water (DI) was used in the synthesis of coating systems.

2.2. Synthesis of 2-[3-(4-methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one

N-[4-(4-methoxy-3-methylphenyl)-4-oxo-2-butenoyl]anthranilic acid (starting material (SM)) was prepared based on reported methods [30,31]. However, 2-[3-(4-methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one was synthesized, as follows: a mixture of (1 g) N-[4-(4-methoxy-3-methylphenyl)-4-oxo-2-butenoyl]anthranilic acid (SM) and 5 mL of acetic anhydride were refluxed for 30 min. The obtained precipitate after cooling was
filtered off, washed with benzene, and then dried. It was then, crystallized from toluene to produce 2-[3-(4-methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one.

2.3. Synthesis of N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid (OA)

A solution of (0.005 mol) bromine in (20 mL) acetic acid was added dropwise in a solution of 2-[3-(4-methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one (0.8 g, 0.0025 mol) in acetic acid (20 mL). The reaction mixture was stirred at room temperature for 2 h then kept overnight. Subsequently, the obtained precipitate was filtered off, washed with diluted acetic acid, and crystallized from acetic acid to give (N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid.

2.4. Synthesis Coating Composite for Furniture Textile Fabrics (CH–OA)

In a glass beaker containing 50 mL of DI water with 2 vol.% acetic acid, 1g of chitosan was dissolved forming chitosan solution of 2 wt.% In the prepared (2 wt.%) chitosan solution, variable mass loadings of OA were dispersed using ultrasonication for 10 min. This process gives rise to 20, 50 and 60 wt.% of OA based on final composite mass, as depicted in Table 1. This is in addition to the pure chitosan solution free of OA.

2.5. Treatment of Furniture Fabrics with Coating Layer (T–CH–OA)

In 100 mL beaker containing chitosan coating dispersion with OA of 0, 20, 50, and 60 wt.% individually, textile fabrics of 20 cm × 20 cm dimension were immersed and squeezed individually and then curried at 130 °C for 10 min. The immersion and curing steps were repeated twice. Finally, different textile composites were obtained, as depicted in Table 1 and denoted as T–CH–OA–X, as T refers to textile fabrics and CH for chitosan, OA refers to developed organic agent, and X refers to the concentration of OA by wt.%.

| Sample Code | Chitosan Mass Loading (CH) (wt.%) | Organic Agent Mass Loadings (OA) (wt.%) | Add On (%) |
|-------------|----------------------------------|----------------------------------------|-----------|
| T-0         | 0                                | 0                                      | 0         |
| CH-T        | 100                              | 0                                      | 10        |
| T-CH-OA-20  | 80                               | 20                                     | 15        |
| T-CH-OA-50  | 50                               | 50                                     | 20        |
| T-CH-OA-60  | 40                               | 60                                     | 20        |

2.6. Characterization

1H NMR spectra were measured using Bruker (300 MHz, DMSO, San Jose, CA, USA) and TMS was used as the internal standard. A FT-IR spectrum was obtained using a Perkin Elmer 1430 ratio recording infrared spectrophotometer (Perkin Elmer, MA, USA). A mass spectrum was measured by a GC-MRSP 1000EX Shimadzu (Shimadzu, Kyoto, Japan). The SEM images were taken using a scanning electron microscope (Quanta FEG-250, Thermo Scientific, OR, USA). The tensile strength was tested using tensile testing machine model H1-5KT/S. Thermogravimetric (TGA) analysis was evaluated using TGA 50 (TA Shimadzu Inc., Kyoto, Japan) from room temperature to 550 °C using nitrogen flow of 100 mL·min⁻¹ and the heating rate used was 10 °C·min⁻¹. The antibacterial activity of samples against staphylococcus aureus bacteria was investigated using the AATCC test method 147-2004 [32]. The antibacterial inhibition zone was measured by the average clear inhibition zone of five replicates based on equation W = (T − D)/2. As, W is the width of the clear inhibition zone in mm and T is the total diameter of both test specimen and clear zone in mm and D is the width of the test specimen itself in mm [29,32]. The flammability properties of the blank and different coated textile fabrics were studied while using a Fire Testing Technology UL94 flame chamber according to
modified ISO 3795 [4,8,10,28,33]. The flame retardant performance were evaluated in terms of rate of burning B in mm/min. Additionally, the flammability properties were further investigated using a Fire Testing Technology limiting oxygen index instrument (LOI) (%).

3. Results and Discussion

3.1. Structural Characterization of Developed Organic Agent (OA)

3.1.1. 2-[3-(4-methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one (2)

The key starting material, 2-[3-(4-methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one (2) was synthesized in a good yield by the cyclization reaction of N-[4-(4-methoxy-3-methylphenyl)-4-oxo-2-butenoyl]anthranilic acid (1) with acetic anhydride, as depicted in Scheme 1.

![Scheme 1](image)

The structure of compound (2) was elucidated while using spectral and elemental analysis. The FT-IR spectrum of this compound exhibited the characteristic absorption bands of compound (2), as indicated in Figure S1a. Hence, the peak observed at 1660 cm\(^{-1}\) corresponds to vibration of C=O of \(\alpha,\beta\) unsaturated carbonyl, and the peak situated at 1769 cm\(^{-1}\) is attributed to C=O of benzoxazinone ring. Furthermore, the peaks located at 2850–2919 cm\(^{-1}\) are assigned to stretching vibration of C–H aliphatic. The \(^1\)H-NMR spectrum showed the appearance of two singlets in the aliphatic regions at 2.23 ppm and 3.90 ppm, which are characteristic for Ar–CH\(_3\) and Ar–OCH\(_3\), respectively. However, the observed multiplets are attributed to the olefinic and aromatic protons appeared between 7.05 and 8.18 ppm, respectively (Figure S1b).

3.1.2. N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid (Organic Agent: OA)

Actually, attempts to synthesize the 2-[2,3-dibromo-3-(4-methoxy-3-methylphenyl)-3-oxopropyl]-4H-3,1-benzoxazin-4-one through the reaction of the benzoxazinone (2) with bromine in different solvent failed to give the desired product. However, the treatment of benzoxazinone (2) with bromine in acetic acid or methylene chloride afforded the N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid (3) (OA), as shown in Figure 1.
Figure 1. Chemical structure illustration representing the synthesis of N-[2,3-Dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid (3) (OA).

The proposed structure of this compound was supported by the identity of its melting point with that of an authentic sample prepared by the treatment of the N-[4-(4-methoxy-3-methylphenyl)-4-oxo-2-butenoyl] anthranilic acid (1) with bromine in acetic acid. The structure of compound (3) was elucidated by spectroscopic techniques as shown in Figure S2. Figure S2a represents the FT-IR spectra of compound (3) revealed the absence of the absorption bands corresponding to the carbonyl group of the benzoxazinone ring. In addition, the absorption band of carbonyl group of the side chain was observed at 1659 cm\(^{-1}\). Moreover, the absorption bands characteristic for O–H and N–H groups were observed at 3287 and 3206 cm\(^{-1}\), respectively. The \(^1\)H-NMR spectra (DMSO-d\(_6\)) showed the appearance of two singlets in the aliphatic regions observed at 2.23 ppm and 3.92 ppm which is characteristic for Ar–CH\(_3\) and Ar–OCH\(_3\), respectively (Figure S2b). The signals assigned to the CO–C–H–Br–C–H–Br–CO protons were observed as two doublets at 5.22–5.25 and 5.98–6.01 ppm. The multiplets attributed for the aromatic protons appeared between 7.11 and 8.19 ppm as shown in Figure S2b. The signal for N–H group was observed as singlet at 11.33 ppm. Furthermore, a broad signal over 12 ppm was attributed to the O–H group. The mass spectrum depicted in Figure S2c showed the molecular ion peak at \(m/z\) 499 (0.9%) which is consistent with its molecular formula C\(_{26}\)H\(_{23}\)N\(_5\)O\(_2\) (499.15). Furthermore, the molecular ion underwent suggested fragmentation as shown in the following pattern as shown in Figure S3.

3.2. Characterization of Developed Coated Textile Fabrics

Furniture textile fabrics were coated with chitosan layer of 2 wt.% producing T–CH. Subsequently, variable mass loadings of developed OA were individually dispersed via ultrasonication, producing 20, 50, and 60 wt.% of CH–OA based on the final mass of final composite (Table 1). Afterwards, the produced composites were coated on textile fabrics surface, denoted generally as T–CH–OA. The surface morphology of uncoated and coated textile fabrics was investigated using SEM. Figure 2a represents the blank uncoated surface of textile fabrics, which reflects smooth surface of untreated fibers. In contrast, when chitosan chains were coated on the fabrics, the surface becomes rough, reflecting the coating layer of chitosan chains of T–CH composite (Figure 2b). This coating layer was clarified in high magnification SEM image (Figure 2c). Interestingly, when OA was incorporated on chitosan coating layer through dispersion with aid of ultrasonication process, the coating layer was found to be in fiber shape, as indicated on SEM images shown in Figure 2d,e of the T–CH–OA–50 composite. These fibers were aligned on textile fiber
surface, as indicated by arrows (Figure 2d,e). This fiber shape of coating might be formed during the ultrasonication process for dispersion OA in chitosan solution.

3.3. Thermal Stability and Flammability Properties

On the other hand, thermogravimetric analysis of untreated and treated textile fabrics gives a clear indication of the effect of OA on thermal stability and charring tendency of coated textile fabrics. Figure 3a shows the thermogram of uncoated sample, which has an initial mass loss that started at 280 °C (onset decomposition temperature) due to the decomposition of cellulose chains of fibran and wool chains. However, the main mass loss due to the decomposition of polyester and microfilament chains started at 391 °C [4]; moreover, the temperatures at 20% and 50% mass losses were found to be 325 and 414 °C, respectively, and leaving a char residues of 14% at 550 °C, as indicated in Table 2. However, for coated samples, similar thermal behavior was noticed; however,
the temperature of main mass loss was improved from 393 °C in T–CH and reached 397 °C in T–CH–OA–50, as indicated in Figure 3b–d and Table 2. It was noteworthy to note that the thermal stability was enhanced as OA mass loading incorporated in coating layer was increased (Figure 3c,d and Table 2), achieving higher decomposition temperature at 50% mass loss for T–CH–OA–50 when compared to T–CH–OA–20, T–CH, and blank samples. Interestingly, the char residue was also enhanced and the char residues recording 17.2 and 18% in T–CH–OA–20 and T–CHOA–50, respectively, as compared to 14.7% and 14% in T–CH and blank samples respectively (Figure 3 and Table 2). This is indicated that the inclusion of OA in coating layer induces the textile fabrics for forming char barrier during combustion process yielding good charring effect. This behavior was due to the nitrogen contained structure of OA. Additionally, the influence of formed char barrier was clearly noticed at the main decomposition step as indicated in point A in Figure 3 which shifts the decomposition temperature to higher one (397 °C) for T–CH–OA–50. Additionally, the ability of T–CH–OA–50 for forming high char residue was also noticed after decomposition of textile fabrics as indicated in point B on Figure 3. On the other hand, the flame retardancy performance for blank and coated textile fabrics samples were evaluated using rate of burning test (UL94) according to modified ISO 3795 standard [4,8,10,28] and also further evaluated using LOI test (Table 3). The flame retardancy behavior was significantly improved when considerable mass loading of new charring molecule was incorporated in the textile coatings. The rate of burning of virgin textile was found to be 118 mm/min; however, once 20 wt.% of OA was incorporated in the coating layer, the rate of burning was reduced to 45 mm/min. in T–CH–OA–20. Interestingly, when 50 wt.% was introduced in textile coating, zero rate of burning was achieved for T–CH–OA–50 sample achieving first class flame retardant textile fabrics [28]. This flame retardancy behavior was corroborated by achieving LOI value of 23.8% for T–CH–OA–50 as compared to 18.2%, 19.3%, and 21% for blank, free OA, and 20 wt.% contained coated samples, respectively. This affirms the char layer formed was strong enough to afford protective layer upon burning of coated textile fabrics retarding the heat and mass transfer process and, hence, isolating textile flaming layer from burning ones [8,10]. Noteworthy to note that, the flame retardancy behavior of developed textile coated sample T–CH–OA–50 was found to be superior than previous reported coatings for this purpose [9,34].

Figure 3. Thermogravimetric analysis (TGA) graphs of (a) Blank textile, (b) T–CH, (c) T–CH–OA–20, and (d) T–CH–OA–50.
Table 2. Thermal characteristic and flammability data for uncoated and coated samples.

| Sample Code     | (a) \(T_{20}\) °C | (b) \(T_{on}\) °C | (c) \(T_{50}\) °C | (d) Char Residue wt.% |
|-----------------|--------------------|-------------------|-------------------|----------------------|
| Blank           | 325                | 391               | 414               | 14                   |
| T–CH            | 339                | 393               | 420               | 14.7                 |
| T–CH–OA–20      | 325                | 395               | 415               | 17.2                 |
| T–CH–OA–50      | 330                | 397               | 423               | 18                   |

(a) Temperature °C at 20% mass loss, (b) onset decomposition temperature of main peak, (c) temperature at 50% mass loss and (d) char residue at 550 °C.

Table 3. Flammability data for uncoated and coated samples.

| Sample Code   | Rate of Burning (mm/min) | LOI (%) |
|---------------|--------------------------|---------|
| Blank         | 118                      | 18.2    |
| T–CH          | 90                       | 19.3    |
| T–CH–OA–20    | 45                       | 21      |
| T–CH–OA–50    | 0                        | 23.8    |

3.4. Mechanical and Antibacterial Properties of Developed Textile Fabrics

The effect of coating on the tensile strength properties of the developed fabrics was evaluated, as tabulated in Table 4. The tensile strength of the untreated sample was found to be 535 N with elongation of 29%. While, when chitosan layer was coated, the tensile strength of T–CH was increased to 759 N, with a reduction in elongation (Table 4). However, when OA was introduced in the coating the tensile strength was also increased recording 626 N in T–CH–OA–20. Interestingly, when OA mass loading was further increased in T–CH–OA–50, the tensile strength was further enhanced, achieving 51% and 7% improvement as compared to blank and T–CH samples, respectively (Table 4). It is noted that the tensile strength was found to be 782 N, along with 15.7% elongation for T–CH–OA–60, which is superior when compared to T–CH and T. This enhancement in tensile strength was attributed to the special fiber shape effect of CH–OA coating, which aligned on the fabrics fiber surface strengthen their yearn and, hence, reinforced their strength.

Table 4. Mechanical properties of uncoated and coated textile samples.

| Sample Code     | Tensile Strength (N) | Elongation (%) |
|-----------------|----------------------|----------------|
| Blank           | 535 ± 4              | 29 ± 1.4       |
| T–CH            | 759 ± 3              | 6 ± 0.4        |
| T–CH–OA–20      | 625 ± 44             | 8 ± 1.6        |
| T–CH–OA–50      | 811 ± 44             | 12 ± 4         |
| T–CH–OA–60      | 782 ± 23             | 16 ± 4         |

On the other hand, the developed OA exhibited good antibacterial behavior for treated textile fabrics. The antibacterial properties of the treated and untreated textile fabrics have been tested against staphylococcus aureus bacteria. The untreated, T–CH, and T–CH–OA–20 samples exhibited no antibacterial effect, as indicated in Figure 4. However, when the mass loading of OA incorporated in coating layer was increased in T–CH–OA–50 the antibacterial effect was observed recording 12.3 mm of clear inhibition zone. Interestingly, a further increase of OA mass loading attributed to positive enhancement in the antibacterial effect achieving 18 mm of clear inhibition zone (Figure 4 and Table 5). This clarified that OA has a potential effect for inhibition of the growth of bacterial cells (Table 5). It is important to note that the OA was found to be not only good charring agent, but also good reinforcement, flame retardant, and potential antibacterial material. Thus, the treated textile fabrics can be used in various applications, achieving the required comfortability demanded by users. It is noteworthy to note that the antibacterial property of the developed T–CH–OA–50 was found to be superior to various reported textile fabrics coatings \[7,8,34,35\]. Interestingly, the coated...
textile fabrics were found to be stable against washing process for more than few tens washing cycles affirms the durability properties.

Table 5. Antibacterial zone of untreated and treated textile fabrics.

| Sample Code       | Average Width of Clear Inhibition Zone (mm) |
|-------------------|--------------------------------------------|
| T                 | 0                                          |
| T–CH              | 0                                          |
| T–CH–OA–20        | 0                                          |
| T–CH–OA–50        | 12.3                                       |
| T–CH–OA–60        | 18                                         |

![Digital photos of bacterial inhibition zone of (a) T, (b) T–CH and T–CH–OA–20, and (c) T–CH–OA–50 and T–CH–OA–60.](image)

4. Conclusions

Smart organic antibacterial and charring agent was developed. N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid (OA) was facilely synthesized and used as flame retardant, charring, and antibacterial organic agent for textile fabrics. N-[2,3-dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid was used with chitosan for coating furniture textile fabrics. The mass loadings of developed organic agent was varied and optimized. The coating composite coated on the textile fabrics surface was found in the fiber shape. The thermal stability and char residues of coated textile fabrics were enhanced when compared to blank and chitosan coated samples. Additionally, the flame retardancy was improved achieving 23.8% LOI and zero rate of burning for developed coated sample as compared to 18.2% and 118 mm/min. for blank untreated sample. The tensile strength of the coated textile fabrics was improved by 51% and 7% when compared to the uncoated blank and chitosan coated samples, respectively. The antibacterial behavior for the coated textile fabrics included organic agent was positively enhanced recording 18 mm of clear inhibition zone as compared to negative behavior for blank and organic agent free coated samples.
Supplementary Materials: The following are available online at www.mdpi.com/2079-6412/10/6/576/s1, Figure S1: FTIR spectrum (a) and $^1$H NMR (300 MHz, DMSO) spectrum (b) of 2-[3-(4-Methoxy-3-methylphenyl)-3-oxoprop-1-enyl]-4H-3,1-benzoxazin-4-one (2), Figure S2: FTIR spectrum (a), $^1$H NMR (300 MHz, DMSO) spectrum (b) and mass spectrum (c) of N-[2,3-Dibromo-4-(4-methoxy-3-methylphenyl)-4-oxobutanoyl]anthranilic acid (3) (OA), Figure S3: Suggested fragmentation pattern of compound (OA).

Author Contributions: Conceptualization, N.F.A. and S.S.E.S.; methodology, N.F.A.; software, M.H.S.; validation, N.F.A.; S.S.E.S. and M.H.S.; formal analysis, M.H.S.; investigation, N.F.A.; resources, S.S.E.S.; data curation, M.H.S.; writing—original draft preparation, N.F.A.; writing—review and editing, N.F.A.; visualization, N.F.A.; supervision, S.S.E.S.; project administration, M.H.S.; funding acquisition, S.S.E.S. All authors have read and agreed to the published version of the manuscript.

Funding: This research received no external funding.

Conflicts of Interest: The authors declare no conflict of interest.

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