

Three-Dimensional Nanoscale Morphological Surface Analysis of Polymeric Particles Containing *Allium sativum* Essential Oil

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Abstract: Biodegradable particles were developed using poly-ε-caprolactone and gelatin carriers containing different concentrations of *Allium sativum* essential oil (EO) (360 µg/mL, 420 µg/mL, and 460 µg/mL). Atomic force microscopy was useful to evaluate the particles’ surface based on morphological parameters. The particles’ size varied from 150 nm to 300 nm. The diameter was related to the increase of the particles’ height as a function of the EO concentration, influencing the roughness of the surface core values (from 20 to 30 nm) and surface irregularity. The spatial parameters Str (texture aspect ratio) and Std (texture direction) revealed low spatial frequency morphological parameters. The particles’ size varied from 150 nm to 300 nm. The diameter was related to the increase of the particles’ height as a function of the EO concentration, influencing the roughness of the surface core values (from 20 to 30 nm) and surface irregularity. The spatial parameters Str (texture aspect ratio) and Std (texture direction) revealed low spatial frequency components. The hybrid parameters Sdq (root mean square gradient) and Sdr (interfacial area ratio) also increased as a function of the EO concentration, revealing fewer flat particles. On the other hand, the functional parameters (inverse areal material ratio and peak extreme height) suggested differences in surface irregularities. Higher concentrations of EO resulted in greater microtexture asperity on the particles’ surface, as well as sharper peaks. The nanoscale morphological surface analysis allowed the determination of the most appropriate concentration of encapsulated EO, influencing statistical surface parameters.

Keywords: morphological parameters; atomic force microscopy; biodegradable particles; *Allium sativum*; essential oil

1. Introduction

*Allium sativum* bulbs (Asparagale: Amaryllidaceae) are well known worldwide due to their health benefits, as well as culinary applications [1]. The antimicrobial [2], antifungal [3,4], and antioxidant [1] bioactivity of this essential oil has been extensively reported in the scientific literature. Organosulfur and phenolic compounds are the major essential oil compounds [5,6]. The insertion of nanotechnology in food products such as gelatin
to develop alternative controlling agents represents an interesting area of research [7–9]. Encapsulation techniques increase the stability of EO in processing and storage, as well as reduce their natural volatility, oxidation, and photodegradation [10]. Furthermore, the encapsulation process allows their controlled release [11,12]. The emulsion–diffusion encapsulation technique enables particle size control, good reproducibility, and high encapsulation efficiency [13–19]. In this work, different concentrations of *A. sativum* essential oil were encapsulated in particles constituted of poly-ε-caprolactone (PCL) and gelatin carriers.

The development of controlled release systems requires a systematic understanding of the particles’ surface. The atomic force microscopy (AFM) technique has been extensively applied to evaluate complex surfaces [20–23], providing information on morphological [24–27] and fractal [28–30] parameters, as well as a power spectrum density (PSD) [31–34]. These parameters allow the proposition of specific applications associated with the release mechanism of encapsulated compounds [35]. There are several free and commercial softwares to perform morphological evaluations, representing useful alternative tools for material surface characterization.

The aim of this paper was to evaluate the morphology and spatial frequency (to understand the dynamics of topographic changes) of the PCL/gelatin particles to identify surface patterns of interest for technological purposes.

A novelty discussion of statistical surface parameters related to nanoparticle-based polymeric carriers was reported here based on the AFM technique and the use of image processing. Our results allowed us to understand the influence of various essential oil concentrations on the evaluated statistical surface parameters, such as roughness, volume, peak distribution, height distribution and texture homogeneity, providing many important aspects of polymeric films surface, including the capsules’ shape. All analyses were performed using standards such as the International Standardization Organization (ISO).

2. Materials and Methods

2.1. Materials

All aqueous solutions were prepared using ultrapure deionized water (18.2 M.Ω.cm, Gehaka, Master WFI, São Paulo, SP, Brazil). Dichloromethane and acetone were purchased from LabSynth, Diadema, SP, Brazil. Gelatin, Tween 80®, PCL, sorbitan stearate (Span 60), and caprylic/capric acid triglyceride (TAAC) were purchased from Sigma-Aldrich, Barueri, Brazil. Transglutaminase was purchased from GastronomyLab, Brasilia, Brazil.

2.2. Essential Oil Encapsulation

*A. sativum* bulbs were collected in Manaus/AM, Brazil, and dried under controlled humidity at 30 °C until reaching constant weight. Then, 400 g of milled bulbs were subjected to hydrodistillation using a Clevenger-type apparatus (NetLab, Tatuapé, SP, Brazil) for 2 h at 100 °C. The essential oil was stored at −18 °C.

PCL/Gelatin-based particles loaded with *A. sativum* essential oil were synthesized based on previous report [36] with modifications. Solution I was prepared as follows: Gelatin was heated to 50 °C in distilled water under constant stirring. Then, Tween 80® (0.30 g) was added to Solution I when the temperature decreased to 40 °C. Solution II was prepared using PCL (0.05 g), sorbitan stearate (span 60; 0.02 g), and caprylic/capric acid triglyceride (TAAC; 0.1 g) solubilized in acetone (15 mL). Different concentrations of essential oil (360 µg/mL (P₃₆₀), 420 µg/mL (P₄₂₀), or 460 µg/mL (P₄₆₀)) were added to Solution II, which was transferred to Solution I under constant stirring (10,000 rpm) using an ultra-disperser. Transglutaminase (0.19 g) was added to the final solution. Then, the formulation was kept under constant stirring until acetone evaporation. All systems were maintained in bio-oxygen demand (BOD) chamber at 25 °C in sealed vials. P₀ represents the unloaded system.
2.3. AFM Imaging

The systems NP0, NP360, NP420, and NP460 94 (1 µL) were dripped on a glass slide and, to speed up the drying process, we exposed the films to a flow of nitrogen gas with the aid of an air gun to control this flow on the surface. Then, the glass slide containing the formed film was fixed on the AFM sample holder using an adhesive tape. Measurements were performed at room temperature (296 ± 1 K) and (40 ± 1)% R.H. on an Innova (Bruker®, Billerica, MA, USA) equipment operated in tapping mode and equipped with a silicon tip and Al coated cantilever with a spring constant of 42 N/m (Tap190AL-G from Budget SensorsTM, Izgrev, Sofia, Bulgária). Scans were performed using (10 × 10) µm² with (256 × 256) pixels at a scan rate of 1 Hz.

2.4. Surface Analysis

The AFM images were analyzed using the Mountains Map® (Besançon, France), Premium software version 8.4.8872 [37]. Morphological parameters were extracted according to the ISO 25178-2: 2012 [38]. Height, functional, spatial, hybrid, feature, core roughness Sk, and volume parameters were discussed according to their previous definition [24,39–41]. Additionally, the one-dimensional power spectrum density (PSD) was evaluated using the free software WsxM© 5.0 (Nanotec Electrónica S.L., Madrid, Spain) [42] using the height distribution autocorrelation function from Equation (1) [43], where \( L_z \) is the image pixels number per line and \( q_z \) is the vector associated with the z value of the height distribution.

The average PSD was obtained from the topographic maps using Equation (2) [43].

\[
\begin{align*}
\text{h(x, y)} & = L_z^{-1} \sum_{q_z} \text{PSD}^{1D}(q_z) e^{i q_z z}, \\
\text{PSD}^{1D}(q_z) & = L_x^{-1} \left[ \int_{L_x} h(x, y) e^{-i q_x x} \right]^2,
\end{align*}
\]

The linearized graph (PSD x q) was obtained from the spectrum data using Equation (3), where \( C_0 \) is constant.

\[
\text{PSD} = C_0 q^{-2 - 2H},
\]

The Hurst coefficient (H) of the surface nanotexture was estimated from the linearized graph using Equation (4) [43,44], where \( \mu \) is the absolute slope of the average PSD curve.

\[
H_c = \frac{\mu - 2}{2},
\]

2.5. Statistical Analysis

Statistical analysis was performed using the OriginPro® software version 9.0 (OriginLab Corporation, Northampton, MA, USA). The variance analysis (ANOVA) was applied using a 5% significance level (p-value < 0.05) based on Tukey’s test.

3. Results

3.1. Morphological Analysis

The AFM technique allows an accurate surface analysis, providing a significant increase of spatial resolution compared to all other far-field microscopy optical methods. The lateral resolution is determined by the tip radius (~20 nm or less). On the other hand, the vertical resolution reaches nanometer fractions, allowing a significant evaluation of the particle’s height and surface roughness. For this reason, the morphological evaluation of the developed particles as a function of the essential oil concentration was investigated here using this powerful technique.

Figure 1 shows the 2D and 3D AFM topographical maps. Flat regions on the particle’s surface should be preferred due to the intrinsic features of this technique. For this reason, particles presenting curved surfaces can limit the reliability of the morphometric data and restrict the investigation of samples presenting marginal complex three-dimensional shell
configuration [45]. The particles’ topography evaluated here was dependent on the fashion they were dripped on the glass surface, as well as on the drying process, so irregularities were expected. For this reason, smaller scans were performed to overcome this problem. The average particle diameter, height, and height/diameter ratio are shown in Table 1 [46].

The particles developed had spherical cap-like shapes and showed a uniform size distribution. The average size varied from $184 \pm 30$ nm to $261 \pm 47$ nm. The increase in the average diameter was accompanied by the average increase in the height of the particles, but not in the same proportion (since the concentration of essential oil also increased), as shown in Table 1. Due to significant differences in the height values, the height/diameter ratios were obtained: a significant increase was observed as a function of the encapsulated essential oil concentration. Other reports have applied the AFM technique to estimate the particles range for systems formed by gelatin and PCL [35,36,47].

### Table 1. Average particle diameter, height, and height/diameter ratio.

| Parameter             | P$_{360}$       | P$_{420}$       | P$_{460}$       |
|-----------------------|-----------------|-----------------|-----------------|
| Diameter (nm)         | $184 \pm 30$    | $236 \pm 49$    | $261 \pm 47$    |
| Height (nm)           | $18 \pm 5$      | $31 \pm 6$      | $56 \pm 6$      |
| Height/Diameter (nm)  | $0.09 \pm 0.04$ | $0.13 \pm 0.05$ | $0.21 \pm 0.06$ |

Figure 1. Two-dimensional and 3D AFM images of (a) P$_{360}$, (b) P$_{420}$, and (c) P$_{460}$ particle surfaces.
3.2. Stereometric Evaluation

Statistical stereometric evaluation parameters are shown in Table 2. Most of these parameters presented a significant difference ($p$-value < 0.05), except the area material ratio (Smr), texture aspect ratio (Str), and texture direction (Std).

Table 2. Statistical morphological parameters according to ISO 25178-2: 2012.

| Parameter | $P_{360}$ | $P_{420}$ | $P_{460}$ |
|-----------|-----------|-----------|-----------|
| **Functional** | | | |
| Smr * (%) | 100 ± 0 | 100 ± 0 | 100 ± 0 |
| Smc (nm) | 10 ± 1 | 15 ± 1 | 21 ± 1 |
| Sxp (nm) | 15 ± 2 | 20.4 ± 0.3 | 24 ± 3 |
| **Spatial** | | | |
| Sal (nm) | 0.7 ± 0.1 | 0.64 ± 0.07 | 0.49 ± 0.06 |
| Str * | 0.49 ± 0.06 | 0.37 ± 0.07 | 0.6 ± 0.2 |
| Std * ($^\circ$) | 131 ± 74 | 132 ± 74 | 133 ± 74 |
| **Hybrid** | | | |
| Sdq (-) | 0.039 ± 0.001 | 0.064 ± 0.001 | 0.100 ± 0.004 |
| Sdr (%) | 0.076 ± 0.005 | 0.22 ± 0.02 | 0.49 ± 0.04 |
| **Feature** | | | |
| Spd (1/µm²) | 1.2 ± 0.1 | 1.14 ± 0.04 | 0.92 ± 0.06 |
| Spc (1/µm) | 0.67 ± 0.05 | 1.11 ± 0.07 | 2.3 ± 0.3 |

*Samples without significant difference ANOVA One-Way ($p < 0.05$).

From the functional parameter values, only the Smr parameter was similar in all developed systems, exhibiting the highest possible proportion (100%) [40]. However, the inverse areal material ratio (Smc) and the peak extreme height (Sxp) values increased as a function of the essential oil concentration. From the medium plane to the highest peak, the proportion of material increased for higher essential oil concentrations, suggesting differences in surface irregularities.

Considering the spatial parameters, the Str and Std average values were not significant, revealing similar isotropy patterns. Furthermore, the autocorrelation length (Sal) parameter decreased from $P_{360}$ to $P_{460}$. This result shows that the developed systems are dominated by low spatial frequency components [24], providing patterns with relatively irregular texture, as more regular textures present $S_{al} \leq 0.2$ [40].

Considering the hybrid parameters, the root mean square gradient (Sdq) and developed interfacial area ratio (Sdr) increased when the concentration of EO was increased. Less flat topographical spatial patterns were observed when the concentration of EO was increased (full flat surface present Sdq = 0 and Sdr = 0% [24,40]).

However, considering the feature parameters, a slight decrease of the peak density (Spd) was observed because this parameter decreased when the concentration of EO was increased. This result indicated greater spacing between rough peaks for high EO concentrations. Moreover, the peak shape also became more pointed when the concentration of EO was increased (as the arithmetic mean peak curvature (Spc) increased from $P_{360}$ to $P_{460}$), suggesting a growth of peaks.

3.3. Watershed Segmentation of Particles’ Microtexture

The thickness and volume parameters related to the core of the surface’s microtexture are shown in Table 3, and their representations are shown in Figure 2. All parameters presented a significant difference ($p$-value < 0.05), except the valley material portion (Smr2) parameter. The roughness of the surface’s core increased because the parameters core
roughness \( S_k \), reduced peak height (\( S_{pk} \)), and reduced valley depth (\( S_{vk} \)) increased as a function of the EO concentration. For this reason, the roughness of the surface’s core was increased, resulting in a significant surface irregularity. Furthermore, the peak material portion (\( S_{mr1} \)) parameter decreased from \( P_{360} \) to \( P_{460} \). This result was consistent, although the \( S_{mr2} \) parameter presented a similar value in all developed particles. Similarly, the parameters \( S_k \), peak material volume (\( V_{mp} \)), core material volume (\( V_{mc} \)), dale void volume (\( V_{vc} \)), and core void volume (\( V_{vv} \)) increased from \( P_{360} \) to \( P_{460} \). This result reveals that the volume of the material that can be deposited on the peaks and valleys of the surface increased as a function of the essential oil concentration. For this reason, physical properties such as friction, wear, and adhesion can be affected. This result is a consequence of the increase in global and local roughness as the increase of particles (diameter and height, as shown in Table 1) promoted an increase of roughness in all surface regions.

| Table 3. \( S_k \) and volume parameters according to ISO 25178-2:2012. |
|-----------------|-------|------|------|
| Parameter       | \( P_{360} \) | \( P_{420} \) | \( P_{460} \) |
| \( S_k \) (nm)  | 20 ± 2     | 29.013 ± 3 | 30.484 ± 4 |
| \( S_{pk} \) (nm) | 8.9 ± 0.9  | 13.397 ± 0.828 | 26.031 ± 4 |
| \( S_{vk} \) (nm) | 7.2 ± 0.8  | 9 ± 1     | 14 ± 2     |
| \( S_{mr1} \) (%) | 13 ± 1    | 12.0 ± 0.7 | 17 ± 1  |
| \( S_{mr2} \) * (%) | 89.7 ± 0.8 | 92 ± 1    | 91.0 ± 0.9 |
| \( V_{mp} \) (\( \mu m^3/\mu m^2 \)) | \( 4 \times 10^{-4} \pm 4 \times 10^{-5} \) | \( 6 \times 10^{-4} \pm 4 \times 10^{-5} \) | \( 1 \times 10^{-3} \pm 1 \times 10^{-4} \) |
| \( V_{mc} \) (\( \mu m^3/\mu m^2 \)) | \( 7 \times 10^{-3} \pm 9 \times 10^{-4} \) | \( 1.02 \times 10^{-2} \pm 8.18 \times 10^{-4} \) | \( 1.19 \times 10^{-2} \pm 1.04 \times 10^{-3} \) |
| \( V_{vc} \) (\( \mu m^3/\mu m^2 \)) | \( 1 \times 10^{-2} \pm 1 \times 10^{-3} \) | \( 2 \times 10^{-2} \pm 1 \times 10^{-4} \) | \( 2 \times 10^{-2} \pm 1 \times 10^{-3} \) |
| \( V_{vv} \) (\( \mu m^3/\mu m^2 \)) | \( 9 \times 10^{-4} \pm 7 \times 10^{-5} \) | \( 1 \times 10^{-3} \pm 7 \times 10^{-4} \) | \( 1 \times 10^{-3} \pm 2 \times 10^{-4} \) |

* Samples without significant difference ANOVA One-Way (\( p < 0.05 \)).

3.4. Power Spectrum Density (PSD) of the Surface Nanotexture

The power spectrum density (PSD) evaluation has been extensively used to define different patterns in several systems [31–34]. The average PSD spectrum of the developed particles is shown in Figure 3, revealing a small difference in the spatial complexity of the particles, mainly in the \( P_{360} \) and \( P_{420} \) systems. A statistical difference (\( p \)-value < 0.05) was observed.

According to the Hurst (H) results, the \( P_{360} \) system was controlled by higher spatial frequencies, resulting in \( (0.59 \pm 0.04) \), \( (0.70 \pm 0.04) \), and \( (0.065 \pm 0.02) \), respectively, for the systems \( P_{360} \), \( P_{420} \), and \( P_{460} \). Higher concentrations of essential oil promoted a decrease of the dominant spatial frequencies of the particle surfaces. These results agree with the reported Sal values (\( P_{360} \) presented spatial dominant frequencies greater than \( P_{420} \) and \( P_{460} \)). Furthermore, as the Tukey test revealed that the \( P_{420} \) and \( P_{460} \) systems are similar, a significant morphology modification was better noted in \( P_{360} \) to \( P_{420} \) (also in agreement with the obtained height parameters).
presented a significant difference (p-value < 0.05), except the valley material portion (Smr2) parameter. The roughness of the surface's core increased because the parameters core roughness (Sk), reduced peak height (Spk), and reduced valley depth (Svk) increased as a function of the EO concentration. For this reason, the roughness of the surface's core was increased, resulting in a significant surface irregularity. Furthermore, the peak material portion (Smr1) parameter decreased from P360 to P460. This result was consistent, although the Smr2 parameter presented a similar value in all developed particles. Similarly, the parameters Sk, peak material volume (Vmp), core material volume (Vmc), dale void volume (Vvc), and core void volume (Vvv) increased from P360 to P460. This result reveals that the volume of the material that can be deposited on the peaks and valleys of the surface increased as a function of the essential oil concentration. For this reason, physical properties such as friction, wear, and adhesion can be affected. This result is a consequence of the increase in global and local roughness as the increase of particles (diameter and height, as shown in Table 1) promoted an increase of roughness in all surface regions.

Figure 2. Graphical evaluation of Sk parameters (left) and volume parameters (right) based on the Abbott curve calculated for (a,d) P360, (b,e) P420, and (c,f) P460.
4. Conclusions

A systematic 3D nanoscale morphological surface analysis of polymeric particles containing *Allium sativum* essential oil was performed. The developed particles presented spherical cap-like shapes and, depending on the EO concentration, their average sizes ranged from $184 \pm 30$ nm to $261 \pm 47$ nm. The increase of the average diameter occurred as the concentration of EO was also increased. Based on the Hurst (H) results, the values of the H parameter were found around $(0.59 \pm 0.04)$, $(0.70 \pm 0.04)$, and $(0.065 \pm 0.02)$, respectively.
for the systems P_{360}, P_{420} and P_{460}, indicating the P_{360} system was controlled by higher spatial frequencies. Therefore, higher concentrations of EO promoted a decrease of the dominant spatial frequencies of the particle surfaces, which agrees with the Sal-reported values. The evaluation of those stereometric parameters, especially functional, spatial, hybrid, and feature parameters, can influence the development of particles with suitable surface properties for technological application. According to our results, we can suggest that the P_{420} and P_{460} systems presented better conditions for application as biodefensive, since the morphological analysis revealed greater microtexture asperity, lower density of rough peaks, and sharper peaks, which can have an influence on the suitable anchoring of the particle onto another surface and EO release.

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