Modulating the Antioxidant Activity of Thin Layer-by-Layer Films with Polyphenols

Vincent Ball*
Institute National de la Santé et de la Recherche Médicale, France

Abstract
Layer-by-layer deposition of a polycation and of a cheap polyphenol, tannic acid, allows to produce thin coatings having an antioxidant activity proportional to the amount of deposited polyphenol. This means that the used probe, 2,2-diphenyl-1-picrylhydrazyl (DPPH), is able to reach all tannic acid molecules present in the film whatever their location. However, when the tannic acid containing film is capped with a few nanometers thick capping layer made of poly (allylamine hydrochloride) and poly (sodium 4-styrene sulfonate), the DPPH has no access anymore to the embedded tannic acid. On this basis, an application is proposed for the production of a packaging film containing tannic acid as a probe able to sense if the packaging has undergone some mechanical damage.

Keywords: Tannic acid; Layer-by-layer deposition; Antioxidant activity

Introduction
Plants and fruits contain many antioxidant molecules among them polyphenols [1]. Such natural antioxidants can find applications in food science [2] and in particular for food packaging when the molecules of interest are integrated in a film or a membrane. Many polyphenols, among them tannic acid (Figure 1) deposit spontaneously at solid liquid interfaces in a versatile manner. The deposition of TA and other polyphenols occurring spontaneously on glasses from wine or from tea has been exploited recently in materials science [3,4]. Polyphenols contain redox active groups which can also act as coordination centers for metal cations like Fe$^{3+}$. This chemical modality allows the deposition from polyphenol-metal cation mixtures to yield conformal coatings [5] or to obtain films by alternating adsorption steps of the polyphenol and metal cations according to a Layer-by-Layer (LBL) deposition method [6]. The same deposition strategy can be used to deposit polyphenol containing films by alternating adsorption steps of the polyphenol and metal cations according to a Layer-by-Layer (LBL) deposition method [6]. The same deposition strategy can be used to deposit polyphenol containing films by alternating adsorption steps of the polyphenol and metal cations according to a Layer-by-Layer (LBL) deposition method [6]. The same deposition strategy can be used to deposit polyphenol containing films by alternating adsorption steps of the polyphenol and metal cations according to a Layer-by-Layer (LBL) deposition method [6]. The same deposition strategy can be used to deposit polyphenol containing films by alternating adsorption steps of the polyphenol and metal cations according to a Layer-by-Layer (LBL) deposition method [6]. The same deposition strategy can be used to deposit polyphenol containing films by alternating adsorption steps of the polyphenol and metal cations according to a Layer-by-Layer (LBL) deposition method [6].

Material and Methods
TA (ref. 403040, Mw=1701 g mol$^{-1}$), PAH (ref. 283218, Mw=15000 g mol$^{-1}$ obtained by means of gel permeation chromatography as given by the furnishers) and PSS (ref. 24,305-104 Crimson Publishers)
Mw=70000 g mol⁻¹ obtained by means of gel permeation chromatography as given by the furnisher were purchased from Sigma Aldrich and used without further purification. All solutions were prepared before use in 50 mM sodium acetate buffer. The buffer was made from double distilled and deionized water (Millipore D8 system, ρ=18.2 MΩ cm). TA, PAH and PSS were dissolved in the sodium acetate buffer at a concentration of 1 mg mL⁻¹. DPPH (ref. D9132, Sigma-Aldrich) was dissolved at 10⁻⁴ M in absolute ethanol and stored at 4 °C in the absence of light to ensure its stability. The chemical structure of TA and DDPH, as well as the reaction of DDPH with an oxidant are shown in Figure 1. The adsorption substrates were 4 cm x 1 cm quartz slides (Thuet, Blodelsheim, France) cleaned with ethanol, dried under a stream of filtered air and subjected to 5 min of plasma cleaning in an air plasma (PDC-32G-2, Harrick Scientific, USA). The LBL deposition was performed as described in ref. [12]. Briefly: the cleaned quartz slide was dipped during 5 min in the PAH solution, in sodium acetate buffer, in the TA solution and again in sodium acetate buffer. The contact with buffer allowed to remove weakly bound molecules from the surface. The successive deposition of a PAH and TA led to the deposition of one-layer pair. This deposition step was repeated n times. The as obtained (PAH-TA)n films were then rinsed with distilled water and dried under a stream of filtered air, in order to avoid crystallization of salt from the buffer.

![chemical structure](image)

**Figure 1:** Chemical structure of TA (A) and of 2,2-diphenyl-1-picrylhydrazyl (DPPH) as well as the reaction undergone in the presence of an oxidant with the concomitant color change.

The absorbance of the quartz slide was then measured between 250 and 700 nm using a double beam spectrophotometer (UVmc², Safas, Monaco) against a cleaned quartz slide taken as the reference. The films were then immersed in a 10⁻⁴ M DPPH solution dissolved in absolute ethanol which constitutes time t=0 of the DPPH quenching kinetics. To follow that kinetics as a function of the number of PAH-TA layer pairs, the absorbance of the solution with the immersed quartz slide+film was measured every 15 s during 15 min at λ=517 nm with the mc² spectrophotometer. In these kinetic experiments, the reference cuvette contained the ethanolic DPPH solution and an uncoated quartz slide. Hence, the spontaneous quenching of DDPH by oxygen is taken into account. The absorbance will appear as negative values since the films induce a faster quenching than dissolved oxygen. To characterize the film deposition, water contact angles were measured with 5 μL water droplets using the sessile drop method with an attention theta goniometer (Biolin Scientific, Sweden). The obtained contact angle values correspond to the average of 5 regularly spaced locations along the quartz slide and the error bar corresponds to one standard deviation. The water contact angle of LBL films characterizes their surface composition and a last layer effect is often observed for relatively stratified films [13].

**Result and Discussion**

The deposition of (PAH-TA)n films (5 min per adsorption and rinsing step as in ref. [12]) was characterized by means of UV-vis spectroscopy (Figure 2) and by contact angle goniometry (Figure 3). The film growth is of linear type, meaning that the absorbance increases at λ=300 nm, due to TA (PAH does not absorb light above λ=200 nm), scales in proportion to the number of deposited layer pairs (Figure 2). In addition, the interaction of the film with tiny water droplets reflects the nature of the last deposited layer (Figure 3). The films are more hydrophilic (static water contact angle close to 8°) when they are made from TA as the last deposited layer from PAH as the last layer (contact angle close to 12°). The behavior found for (PAH-TA)n films is typical of linearly growing LBL films [6, 11, 13]. The morphology of the (PAH-TA)n films has also been characterized by means of Atomic Force Microscopy and some typical images acquired in the contact mode and in the dry state can be found in Figure 4 of ref. [12].
Figure 2: Absorbance spectra of (PAH-TA)_n films with n = 1 (_____), n = 2 (_____), n = 4 (_____), n = 6 (_____) and n = 8 (_____) deposited layer pairs. The inset displays the absorbance increase at \( \lambda = 300 \text{nm} \) as a function of the number of deposited layer pairs. The full and dotted lines represent the linear regression to the data and the limit of the 95% confidence interval respectively. The (PAH-TA)_n films were deposited on quartz slides and their spectra were measured against a plasma cleaned quartz slide.

Figure 3: Evolution of the static water contact angles (obtained by the sessile drop method from 5µL drops) with the deposition of the (PAH-TA)_n films. n=2.5 and 3.5 correspond to (PAH-TA)_{2.5*-PAH} and (PAH-TA)_{3.5*-PAH} films, hence to coatings ending with the deposition of the polycation whereas n=2 and 3 correspond to films with TA as the last deposition step. All points correspond to independently prepared films.

Figure 4: A: Antioxidant activity of (PAH-TA)_n films as a function of the number of deposited layer pairs. The inset represents the reduction in absorbance at \( \lambda = 517 \text{nm} \) after 15min of reaction between the quartz slide covered with a (PAH-TA)_n film and the 10-4M DPPH solution. Full line and dashed lines represent the linear regression to the data and the limits of the 95% confidence interval respectively. B: Picture of the DPPH solution (10-4M in absolute ethanol) and the DPPH solution containing a quartz slide with a (PAH-TA)_{4 film on both its faces, after 15min of reaction.
The (PAH-TA)ₙ films display some antioxidant activity as quantified through a decrease in absorbance at λ=517nm, typical of the transformation of the purple DPPH into the yellowish DPPH-H (Figure 1). From a qualitative point of view, one observes indeed a reduction in the purple color of a 10⁻⁴M DPPH solution when a quartz slide covered with a (PAH-TA)₄ film on both its faces is immersed in it during 15min (Figure 4A). After 15min the reservoir of active TA in the films seems not yet exhausted as shown by the kinetic runs in Figure 4B. But for convenience we stopped the assays after 15min, and all films were compared for their antioxidant activity after this given reaction time with DPPH. It appears that the antioxidant activity of the (PAH-TA)ₙ films is proportional to the number of deposited layer pairs (Figure 4A) and hence to the amount of incorporated TA (Figure 2). This means that the films are permeable to DPPH which can reach all TA molecules in the film irrespective of their position, either close to the film/solution or to the quartz/film interface. This result is in perfect agreement with the data reported in ref. [11].

There is however a single way to suppress access of DPPH to the TA present in the (PAH-TA)ₙ films by capping them with (PAH-PSS) layer pairs (Figure 5A&5B). Such layer pairs are known to be of compact nature and to suppress the diffusion of a polyelectrolyte like poly-L-lysine (PLL) from a superior (PLL-HA)ₙ stratum (HA represents hyaluronic acid) to a previously deposited (PLL-HA)ₙ stratum [14]. Hence (PAH-PSS)ₙ layer pairs are able to stratify an otherwise un-stratified (exponentially growing) (PLL-HA)ₙ film. In the present investigation we show that an extremely thin (PAH-PSS)₁ film (about 5-6nm in thickness [15]) is able to almost quantitatively suppress the diffusion of a small molecule like DPPH in the underlying (PAH-TA)ₙ film. This is the major finding of the present investigation. It could have some applications in packaging: when a (PAH-TA)ₙ-(PAH-PSS)₁ film is deposited on a package (for instance by alternated spray deposition to speed up the process with respect to alternated dipping) no color change of a drop of deposited DDPH is found as long as the coating has not undergone damage. But as soon as the package has been damaged, the (PAH-PSS)₁ capping layer contains some cracks and becomes permeable to DPPH which then undergoes a visible and fast color change.

**Figure 5:** A: Reduction of the antioxidant activity of (PAH-TA)₃ films after deposition of PAH-PSS “capping layers” as indicated in the inset. Each kinetic curve corresponds to an independently prepared film. B: Evolution of the relative antioxidant activity after 15min of contact with the 10⁻⁴M DPPH solution as a function of the number of (PAH-PSS) capping layer pairs. The relative activity of 1 corresponds to that of the (PAH-TA) film.

**Conclusion**

It has been shown that the antioxidant activity of (PAH-TA)ₙ films can be modulated through an increase in the number of deposited layer pairs and hence with an increase in the amount of TA, the antioxidant polyphenol, incorporated in the film. The addition of only two (PAH-PSS) capping layer pairs totally suppresses the access of DPPH to the underlying TA highlighting the compact and impermeable nature of those very thin capping layers. A practical application in possible sensing of packaging integrity is proposed on the basis of those findings.

**References**

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