Encapsulation of N-Nitroso-melatonin with Poly(lactide-co-glycolide)

Michael Kirsch1*, Hans-Gert Korth2, Joachim Fandrey3 and Katja B Ferenz1
1Institut für Physiologische Chemie, Universität Duisburg-Essen, Hufelandstrasse 55, 45122 Essen, Germany
2Institut für Organische Chemie, Universität Essen, 45117 Essen, Germany
3Institut für Physiologie, Universitätsklinikum Essen, Hufelandstrasse 55, 45122 Essen, Germany

Abstract

N-Nitrosomelatonin (NOMela) is well known for its capabilities to transnitrosate nucleophiles such as thiols and ascorbate thereby generating nitric oxide (NO)-releasing compounds. Like molsidomine, NOMela is one of the few NO-releasing substrates not inducing nitrate tolerance and may be therefore highly suitable as NO-therapeutical. As the physical and chemical properties of NOMela do not allow its direct application (oral or intravascular) in animals/humans, the encapsulation with biodegradable poly(lactide-co-glycolide) (PLGA) polymers was performed and NO-releasing kinetics were studied. NOMela could be successfully encapsulated in PLGA (NOMela-PLGA) with an efficiency of 85% thereby prolonging its half-life time in aqueous solution (e.g. in the cytoplasm of endothelial and smooth muscle cells) about 3-fold. In the presence of "activated hydroxy compounds" like vitamin C and thus under physiological conditions, NOMela-PLGA yielded two therapeutically relevant hormones, melatonin and nitric oxide, via reactions only known (until now) for unencapsulated, freely diffusing NOMela. Importantly, in the absence of any activated hydroxy compound the unwanted hydrolysis reaction of NOMela dominated, generating the non-functional nitrite (and not nitric oxide). These findings suggested that PLGA-encapsulated NOMela will be highly attractive as a novel NO-releasing drug lacking common side-effects of classical NO-releasing molecules such as glyceroltrinitrate.

Keywords: Nitric oxide; N-nitrosomelatonin; Vitamin C; Melatonin; poly(lactide-co-glycolide); poly(lactide-co-glycolide) capsules; EPR spectrometry

Introduction

Nitric oxide (NO) fulfills multiple physiological functions such as in neurotransmission, in the immune response by activated macrophages, and in vasodilation through smooth muscle relaxation [1]. In the blood NO is a short-lived intermediate because it is reversibly oxidized to nitrate by oxyhemoglobin [2] and reversibly but strongly bound by deoxyhemoglobin (deoxyhemoglobin binds NO 105 times stronger than molecular oxygen) [3]. Because of these highly effective mechanisms, it has been suggested that in vivo NO is transported by carrier molecules in order to increase its efficacy [4,5].

In 2001, Blanchard-Fillion et al. observed that N-nitrosomelatonin (NOMela) can act as a NO releasing compound [6]. There is now compelling evidence that a shift of the nitroso function from NOMela to "activated hydroxy compounds" (enols or phenols such as vitamin C [7,8], vitamin E [9], catechols [10] or serotonin [11]) generates melatonin and NO via intermediary alkyl or aryl nitrates, respectively.

\[
\text{NOMela + } R - OH \rightarrow \text{melatonin + } R - \text{ONO} \quad \text{(Eq. 1)}
\]

\[
R - \text{ONO} \rightarrow R - O^+ + NO \quad \text{(Eq. 2)}
\]

\[
4\text{NO} + O_2 + 2\text{H}_2\text{O} \rightarrow 4\text{H}^+ + 4\text{NO}_2^- \quad \text{(Eq. 3)}
\]

At physiological pH such reactions can even outcompete the hydrolysis reaction of NOMela.

\[
\text{NOMela + } H_2O \rightarrow \text{melatonin + } H^+ + NO_2^- \quad \text{(Eq. 4)}
\]

In fact, in a complex biological environment of a cell culture system, NOMela simultaneously generates melatonin and biologically active NO [12]. Importantly, all mentioned reactions (Eqs. 1-4) proceed spontaneously. Because of this fact, release of NO from NOMela cannot be hampered by so-called "nitrate tolerance", a common side-effect of other NO-releasing drugs [1].

Unfortunately, the physical and chemical characteristics of NOMela are unfavorable for its application in animals and humans. An oral administration is obviously senseless because conditions, as present in the stomach, are well known to accelerate the degradation of N-nitroso-indole compounds in general [13]. The alternative intravenous application with a NOMela-enriched salt solution seems to be unpracticable because the solubility in water is too low. NOMela is slightly soluble in ethanol, but the use of this vehicle in animals would alter both metabolism and systemic parameters. In order to confect a pharmacological acceptable approach with a prolonged life-time of NOMela, its encapsulation seems to be a promising alternative. In fact, the encapsulation of sparingly water-soluble drugs into biodegradable poly(lactide-co-glycolide) (PLGA) polymers is an established procedure [14-16].

In the present study, we report on the encapsulation of NOMela with PLGA polymers (NOMela-PLGA capsules) and discuss the releasing kinetics of both NO and melatonin from this system with the idea of placing a useful application form of those two hormones at the disposal of clinicians.

Materials and Methods

Reagents

Poly (DL-lactide-co-glycolide) (50:50) produced by LACTEL (B6013-2) was purchased from NRC Nordmann Rassmann GmbH (Hamburg, Germany). Vitamin C, Trolox, Nile-red and hemoglobin came from Sigma-Aldrich (Deisenhofen, Germany).

Synthesis of N-nitrosomelatonin

Melatonin (276 mg) was dissolved in acetone (3 mL) and H_2O (9
mL) was added by vortexing. NaNO₂ (208 mg) was dissolved in H₂O (3 mL) and was then added to the melatonin solution by vortexing for about 5 min. Afterwards HCl (1.5 mL, 1 M) was added under rigorous stirring and this mixture was dried cooled down to 4°C. The generated product N-nitrosomelatonin was extracted by shaking with dichloromethane (20 mL, 4°C) and the separated, water insoluble organic phase was dried by adding Na₂SO₄ (750 mg). After removing Na₂SO₄ this mixture was cooled down to −20°C and taken to evaporation under reduced pressure (18 Torr) to yield about 250 mg N-nitrosotryptophan (Ɛ₃₄₆=7070 M⁻¹ cm⁻¹ [17]).

**Capsule preparation**

The preparation of the PLGA-NOMela microcapsules basically followed the emulsion-evaporation procedure reviewed by Pisani et al. for capsules with sizes in the low µm range [18]. An organic solution of PLGA (100 mg) in the absence or in the presence of NOMela (26.1 mg) in dichloromethane (4 ml) at 4°C was emulsified into an aqueous solution (20 ml) of 35 mM sodium cholate by using an Ultraturrax T25 (IKA Werke, Staufen, Germany) operating with a S25KV-25G-IL dispersing tool at a velocity of 3200 rpm. For laser-scanning microscopy, 100 µL of a stock solution of Nile-red (0.057 mg/ml in dichloromethane) was added to the organic solution prior to emulsification [19]. The dichloromethane was then removed by bubbling with synthetic air for 20 min at 4°C. Such a procedure yielded polydisperse dispersions containing capsules with mean sizes of about 2 µm which were used for all experiments.

**Laser-scanning microscopy (LSM)**

A laser-scanning microscope equipped with a helium/neon laser was used to study the formation of microcapsules. Imaging of nanocapsules (d<0.5 µm) was not possible due to the resolution limit of the optical system. Nile red-stained NOMela-PLGA capsules were diluted 50-200-fold with 0.9% NaCl solution and placed on an object slide or within a modified Pentz chamber. The objective lens was a 63x NA 1.40 plan-apochromat. Red fluorescence of Nile red excited at 543 nm was collected through a 585 nm long-pass filter. Image processing and evaluation were performed using the software of the LSM 510 imaging system. The diameter of the capsules was analyzed by evaluating the LSM pictures with the CEEXO 2006 CellExplorer software (BioSciTec GmbH, Frankfurt a.M., Germany).

**Deoxyhemoglobin solutions**

Commercially available methemoglobin (4 mM) from bovine was reduced with vitamin C (20 mM) under aerobic conditions in phosphate buffered solution (pH 7.0) containing either NOMela (2 mM) or NOMela-PLGA capsules (100 mg/mL) and deoxyhemoglobin (1.5 mM) in the absence and in the presence of vitamin C (20 mM). The reaction solutions were quickly transferred to an aqueous solution quartz cell (Willmad, Buena, NY). Recording conditions were as follows: microwave frequency, 9.635 GHz; modulation, 0.2 mT; signal gain, 1 x 10⁶; sweep range, 20 mT; microwave power, 20 mW; sweep time, 6 min. Spectrum simulation was performed using the WinSim program [22].

**Determination of the entrapment efficiency**

All experiments were performed in phosphate buffer (50 mM, pH 7.4, 37°C). Time-dependent adsorption of either melanotin or NOMela (220 µM each) by empty PLGA capsules (capsule material=100 mg/mL) was analyzed. An aliquot of 1 mL sample was taken from the supernatant after centrifugation (5000 rpm) and the concentration of either free melanotin or NOMela was determined by fluorescence (λₑₓ=276 nm/λₑₘₓ ≥ 354 nm) or UV-vis absorption (Ɛ₃₄₆=7070 M⁻¹ cm⁻¹ [23]), respectively. The % entrapment efficiency was determined indirectly by using the following formula:

\[
\text{Entrapment efficiency (\%) } = \left( \frac{\text{Total amount of drug used} - \text{Amount of free drug}}{\text{Total amount of drug used}} \right) \times 100
\]

**Results**

The modified oil-in-water (o/w) emulsion/solvent evaporation technique encapsulated NOMela with an entrapment efficiency of 85 ± 8%. Capsules with a mean size between 1 and 3 µm were obtained at the employed velocity of 3200 rpm (Figure 1).

In order to demonstrate that the encapsulated NOMela is accessible for solvated, “activated” hydroxy compounds, the release of melanotin was studied in the absence and in the presence of Trolox or vitamin C, respectively, as additives (Figure 2). In general, the presence of the selected additives facilitated the release of melanotin from NOMela-PLGA capsules. Within a reaction period of 5 h, the amount of melanotin

![Figure 1: Formation of NOMela-PLGA capsules.](image-url)
released from NOMela had been increased two- to three-fold. Although both hydroxy compounds react with solvated nitrosoindole derivatives with nearly the same rate constant [8,9,17,24], Trolox was a bit more effective than vitamin C in releasing melatonin.

In order to demonstrate that the decay of encapsulated NOMela proceeds via reactions outlined in equations 1-4, the formation of nitrite was monitored (Figures 3A and 3B). Since vitamin C interferes with the formation of the nitrite reporter dye formation reaction, these measurements were only performed with Trolox. In the absence of Trolox (Figure 3A), nitrite was simultaneously generated with melatonin over a reaction period of 3 h, but the yield of nitrite was always about 21% higher than the yield of melatonin. For instance, after 5 h of reaction 1490 ± 71 µM nitrite and 1229 ± 19 µM melatonin were detected. The presence of Trolox increased the yield of both products but did not change the enhanced yield of nitrite by about 21% (Figure 3A). For instance, after a reaction period of 5 h 2780 ± 62 µM nitrite and 2300 ± 53 µM melatonin were detected, respectively. Although melatonin and nitrite were simultaneously formed in the absence as well as in the presence of Trolox, they were obviously not generated in the expected 1:1 stoichiometric ratio. In contrast, melatonin and nitrite were released in a strict 1:1 stoichiometric ratio from uncapsulated NOMela in the absence as well as in the presence of Trolox (Figure 3B). The data of Figures 3A and 3B also demonstrate that encapsulation significantly increased the half-life time of NOMela from the known value of about 140 min (Figure 3B, [17]) to >5 h (Figure 3A).

A reasonable explanation for the apparent non-stoichiometric production of melatonin and nitrite in the aqueous phase is a slightly preferred accumulation of melatonin but not of nitrite in the PLGA capsule material. In order to clarify this possibility, experiments with PLGA capsules and melatonin were performed (Figures 4A and 4B). In the first set of experiments melatonin was encapsulated in PLGA with an entrapment efficiency of 25 ± 4% and the kinetic of melatonin release in buffer was monitored (Figure 4A). Melatonin was released in an
exponential manner but leveled off to about 500 µM after an incubation period of 3 h. This time dependence reflected the onset of absorption-desorption equilibrium with a steady-state concentration of melatonin in the capsules somewhat (about 25%) higher than in the bulk aqueous phase. This was further confirmed by the effect which unloaded PLGA capsules exerted on the concentration of both melatonin and nitrite (220 µM each) in the surrounding aqueous phase (Figure 4B). In the presence of unloaded PLGA capsules (capsule material 100 mg/ml) the melatonin concentration decreased exponentially by about 27% from 220 µM to ca. 160 µM to level off after 3 h of reaction, whereas the nitrite concentration remained essentially unaffected over the whole reaction period (Figure 4B). Noticeably, unloaded PLGA capsules decreased the concentration of 220 µM NOMela by about 90% within 15 min (data not shown). Both decreases (27% for melatonin and 90% for NOMela) corresponded very well with the observed entrapment efficiencies (25% for melatonin and 85% for NOMela) in PLGA.

So far the proof is missing that according to equations 1 and 2, NO is released from NOMela-loaded PLGA capsules. Unfortunately, it was not possible to monitor NO production electrochemically because the NO-sensitive electrodes were impaired by adsorption of PLGA. In order to exclude any artefacts driven by PLGA and to unequivocally prove the intermediacy of nitric oxide, deoxyhemoglobin was used as NO scavenger and the corresponding nitrosylhemoglobin was detected by EPR spectrometry. Experiments were performed in the absence and in the presence of vitamin C (in order to maintain iron ions in the ferrous state) as an additive (Figure 5). In the absence of vitamin C formation of nitrosylhemoglobin from decaying NOMela-PLGA capsules could not be detected (data not shown). This failure is in agreement with observations from the literature that hydrolytic decomposition of NOMela does not produce any NO (Eq. 4) [7-9]. However, in the presence of vitamin C, the characteristic three-line spectrum of nitrosylhemoglobin (g=2.011; a (14N)=1.67 mT) was recorded from encapsulated NOMela (lines a and b) as well as from an aqueous solution of neat NOMela (line c). The observation that NO is in fact released in the presence of an "activated hydroxy compound" (Eq. 2) is in agreement with data from the literature [7-9]. The prolonged half-life time of PLGA-encapsulated NOMela is also reflected by the EPR data (compare line a with line c; from the relative ESR intensities an about ten-fold faster production of NO from uncapsulated NOMela is estimated).

**Discussion and Conclusion**

Here we demonstrated for the first time that NOMela can be encapsulated in PLGA capsules and that such a procedure prolongs the life-time of NOMela in aqueous solution. PLGA-encapsulated NOMela yielded in the presence of either vitamin C or Trolox two hormones, melatonin and nitric oxide, according to reactions (Eqs. 1-3). In the absence of such hydroxy compounds nitrite, but not nitrit oxide, was generated by hydrolysis of NOMela (Eq. 4). The apparent excess production (by about 21%) of nitrite over melatonin can satisfactorily be explained by a slight accumulation of the formed melatonin in the capsules’ material (i.e., [melatonin]_aq/[melatonin]_caps>1), whereas nitrite does not accumulate in the capsules, i.e., its concentration is the same in both phases ([nitrite]_aq/[nitrite]_caps=1). Noticeably, PLA is highly effective in transporting hormones [14,25]. Since NOMela-PLGA capsules can be separated from the synthesis solution and thus can be dispersed in any other aqueous solution, the low solubility of NOMela in an aqueous environment is, of course only in principle, no longer a hindrance for its in vivo application.

Generally, there are two possibilities by which activated hydroxy compounds would liberate melatonin and nitric oxide from NOMela-PLGA. On the one side, the activated hydroxy compound may to some degree enter the PLGA capsule in order to react in its interior with NOMela, and the reaction products are then released to the surrounding aqueous phase. This mode would in part explain why the more lipophilic Trolox released melatonin somewhat faster than did the hydrophilic vitamin C (Figure 2). Since both activated hydroxy compounds are known to react with nitrosoindoles with similar rate constant [8,9], an equal distribution of vitamin C and Trolox between PLGA capsules and the water phase would result in an identical rate of melatonin production. If this is not the case, the relative steady-state concentration of Trolox in the capsules must be expected to be higher than that of vitamin C. On the other hand, PLGA-capsules may donate NOMela to the bulk aqueous phase where a rapid reaction with the solvated hydroxy compound takes place. The fact that addition of unloaded PLGA capsules to an aqueous solution of NOMela decreased the NOMela concentration by about 90% indicates that PLGA-encapsulated NOMela is in a favourable equilibrium with free, solvated NOMela ([NOMela]_aq/[NOMela]_caps≈9/1). Of course, both processes must be expected to operate simultaneously, however, the strong enhancement of the half-life time of NOMela in the presence of PLGA capsules indicates that the former mode is of minor importance, i.e., NO release from PLGA-NOMela by reaction with activated hydroxy compounds is primarily controlled by the reduced release of NOMela into the bulk aqueous phase. Such a behavior is desirable for a good transporter system (Figure 6).

In any case, melatonin is expected to support the physiological action of NO because melatonin counteracts the vasospastic effect induced by lysophosphatidylcholine as well as enhances the endothelium-dependent vasodilatation in mesenteric artery and aorta observed in spontaneously hypertensive rats [17].

Initially, PLGA polymers have been developed for use as degradable surgical sutures [26]. Characteristics such as biocompatibility and established regulatory approval for clinical use have attracted interest in PLGA for controlled-release applications [27]. A typical implementation was the injection of hormone-loaded PLGA microcapsules into the
musculature [14]. Particle size was shown to be an important effector. At optimized conditions, i.e., particle size >10 µm, non-circulating PLGA microcapsules remained active for several months [14,27,28]. Such a strategy might be favorable for the encapsulation of melatonin but not for the chemically labile NOMela.

Therefore, due to the relative short half-life time of NOMela, circulating entities (capsules) are obviously a matter of choice. Since circulating particles with a mean size of >5 µm are lethal in animals [29-31], the preferable particle size must be lower than that limiting value. Additionally, the circulatory half-life time of intravenously administered particles is generally limited by phagocytic ingestion. Evidence has been presented that maximal uptake of microparticles into macrophages occurred with particles between 1-3 µm [32-34]. Antibodies and complement proteins which can associate to foreign particles render microparticles more attractive to phagocytes, thereby largely enhancing their clearance from the circulation system [35,36]. Since the mean size of our model NOMela-PLGA capsules presently lies between 1 and 3 µm (Figure 1), it must be expected that they would rapidly be cleared (within a few minutes [37]) from the blood by the reticuloendothelial system. According to Eqs. 1-4 NOMela is a NO-donor generating compound but it is not a direct NO releasing entity. Because of this an expected circulation time of a few minutes would be too short to effectively generate NO-releasing entities. In order to avoid such undesirable effects, the properties of the capsules have to be improved for later in vivo use. So to solve the two problems (i) fast recognition by the RES and (ii) too large size, at first, removal of the capsules by macrophages of the reticuloendothelial system can be minimized by covering their surface with hydrophilic polymers such as covalently bound polyethyleneglycols [37-39]. Although principally, non-covalently bound polymers like Tetronic-908 have been shown to operate similarly in experimental systems [40] because of the replacement of the synthetic macromolecules by plasma proteins, such polymers are not suitable for in vivo applications due to their high toxicity. At second, particles with a lower size (<1 µm, so-called nanoparticles) are removed with a significantly lower rate from the circulation, because they are favorably internalized by endothelial and smooth muscle cells [41]. This is why we are currently working on the preparation of NOMela-PLGA nanocapsules. Covering the surface of those nanoparticles with polyethyleneglycol will additionally down regulate the opsonization process [42]. As we have already successfully developed long circulating polyethyleneglycol-coated perfluorodecalin-filled PLGA-capsules as artificial oxygen carriers [43], we are sure that the preparation of NOMela-PLGA nanocapsules with a modified surface will result in a clinically relevant application form of NOMela.

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