Obtaining and Characterization of a Polyurethane Carrier Used for Eugenol as a Possible Remedy in Oral Therapies

ZORAN POPA1*, LAURA CRISTINA RUSU2#, RAZVAN SUSAN3, IULIA PINZARU4, ELENA ARDELEAN5, FLORINBORCAN6, MIRELA VOICU7, IOANA TUTA SAS8, RAMONA AMINA POPOVICI2#, VOICHITA LAZUREANU1

1 Victor Babes University of Medicine and Pharmacy Timisoara, Faculty of Medicine, 2 Eftimie Murgu Sq., 300041, Timisoara, Romania
2 Victor Babes University of Medicine and Pharmacy Timisoara, Faculty of Dentistry, 2 Eftimie Murgu Sq., 300041, Timisoara, Romania
3 Victor Babes University of Medicine and Pharmacy Timisoara, Faculty of Pharmacy, 2 Eftimie Murgu Sq., 300041, Timisoara, Romania

The cloves are antiseptic, antiparasitic, antibacterial, antifungal, antiviral, anesthetic, analgesic, anti-inflammatory, tonic, carminative, anti-ulcer, antithrombotic, antioxidant and anti-cancerous. They contain eugenol, tannins and flavonoids that also help to strengthen the vein wall. This paper presents the obtaining and the characterization of a polyurethane drug delivery system which can be used for the transmembrane transport of eugenol in oral therapies. The products were analyzed by pH and solubility measurements, thermal decomposition and zetasizer tests and they were applied on mice skin to evaluate their harmlessness. The results suggest that were obtained neutral pH structures with low solubility and a good thermal stability, with sizes between 241 and 289 nm and no toxicity effect was found in the case of studied samples.

Keywords: drug carrier, DSC, eugenol, polyurethane, zetasizer

Experimental part

The reagents

Hexamethylene-diisocyanate (HMDI), polyethylene glycol, MH=200 (PEG), the solvent (acetone) and surfactant (TWEEN®20) were obtained from Merck (Germany); ethylene glycol (EG) was purchased from Lach-Ner (Czech Rep.), and eugenol from Sigma-Aldrich. The reagents were used without any previous purification.

The obtaining of samples

The polyurethane structures were obtained following a protocol which was already described in scientific literature in our previous papers [7-17]:

The 1st step - The preparations of the two phases

The aqueous or hydroxy-phase comprises a mixture of ether and/or ester polyols (e.g. polyethylene-glycol, poly- epsilon-caprolactone) as the main component, diols or diamines with low molecular weight (e.g. ethylene glycol, 1,4-butanediol, 1,6-hexanediol or ethylene diamine) used as chain extenders and a surfactant (e.g. Tween or Span) which are dissolved in distilled water, heated at 35°C and homogenized at 300 rpm for at least 20 min.

Separately, the organic-phase comprising one or two aliphatic di-isocyanates (e.g. isophorone-diisocyanate, hexamethylene-diisocyanate, lysine-diisocyanate) dissolved in acetone, heated at 35°C and homogenized at 300 rpm for at least 20 min.

The biological active substance is introduced in a ratio of 5-15% in one of the two phases depending on the reactivity of its functional groups and/or its solubility in water and acetone. Eugenol was introduced into the aqueous hydroxy-phase.

The 2nd step - The mixing of the two phases

The organic-phase is rapidly injected into the hydroxy-phase, heated at 40°C and magnetically stirred at 350 rpm. This is the moment when the polyurethane structures begin to form and the obtained solution begins to turbid. The stirring will continue, with heating for approx. 4 h; the finishing of the polyurethane structures’ walls (maturing of the finished product) is a lasting process because no catalyst was used at all. In the polyurethane industry, the synthesis involves the use of a catalyst such as tertiary amine compounds (e.g. 1,4-diazabicyclo [2.2.2] octane) or an organometallic derivative of tin.

* email: ramona.popovici@unit.ro

# Authors with equal contribution
The 3rd step - The purification of obtained products

A ratio between the hydroxyl- and organic-phase of 1.2:1, with an excess of alcohol compounds, was used in relation to the isocyanate stoichiometric requirement in order to prevent the formation of secondary products. However, it is possible to have different amines through final products, resulting from the reaction of isocyanates with water. The synthesized products were washed at least 3 times with a 1:3 (v/v) water-acetone mixture in order to remove these amines or some unreacted precursors.

The 4th step - The storage of products

The obtained emulsions were layered in Petri dishes and dried at room temperature for approx. 10 days until it is noticed that the mass of the Petri dish is no longer changing. The resulting powder is stored in Eppendorf tubes at room temperature for further evaluation.

Two different samples (EU_0 and EU_1) were synthesized using the procedure previously described; the precursors used for the synthesis of every sample are presented in table 1.

<table>
<thead>
<tr>
<th>Sample code</th>
<th>Aqueous phase</th>
<th>Organic phase</th>
<th>Active substance</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>EG</td>
<td>PEG</td>
<td>Tween®20</td>
</tr>
<tr>
<td>EU_0</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>EU_1</td>
<td>x</td>
<td>x</td>
<td>x</td>
</tr>
</tbody>
</table>

Yield of encapsulation

The entrapment efficiency was calculated based on a method described by C.P. Dora et al. [18]: the free, entrapped drug can be measured by Beer-Lambert law and it is related to the quantity of drug added to the synthesis.

The formula is:

Entrapment efficiency (EE) = \( 1 - \frac{\text{Quantity of free drug}}{\text{Quantity of drug added}} \) · 100

Animals

C57 female mice (8 weeks old at the beginning of tests) were acquired from Charles River (Budapest, Hungary). The work protocol was analyzed and approved by our ethics committee and it followed the National Institute of Animal Health rules: animals were maintained inside the university Biobase in standard conditions: 12 h light-dark cycle, food and water ad libidum, temperature 24±2°C, humidity above 55%.

Polyurethane structures without and with eugenol (EU_0 and EU_1) and blank solution (solvent without any active compound) were applied on mice skin for 21 days (50 µL solution / application). 12 mice were used in this experiment: group blank (4 mice treated with solvent), group EU_0 (4 mice, treated with empty polyurethanes structures) and group EU_1 (4 mice, treated with polyurethane structures with eugenol). The mice were shaved in the first, the 6th and the 15th day and the applications and measurements were done every three days.

The determinations were performed within 30 min after each application. All measurements on the mice skin were carried out with a Multiprobe Adapter System (MPAS) from Courage-Khazaka, Germany: the measurements of transepidermal water loss (TEWL) were carried out with a Tewameter®TM 300 probe, the levels of melanin and erythematere were measured with a Mexameter®MX 18 probe and the moisture of cornaceous layer with a Corneometer®CM 825 probe.

Statistics

All statistical analyses were performed using a trial version of IBM SPSS. Numerical data were presented as mean ± standard errors. Student t test was used to determine the statistical difference between various experimental groups. Statistical significance was considered at a p-value less than 0.05; ** and *** indicate p<0.01 and <0.001.

Results and discussions

Eugenol is a phenylpropanoid compound, liquid at room temperature (m.p. = -7.5°C) with a good solubility in organic solvents as ethanol and ether, a normal solubility in glacial acetic acid, chloroform, and aqueous sodium hydroxide and it is almost insoluble in water [19]. On the other hand, the values of the solubility and pH of samples based on polyurethane structures with and without eugenol are presented in table 2.
The DSC curves of samples EU_0 and EU_1 indicate that were obtained very stable structures between -10 and 270°C; it is known that polyurethane decomposition start at just 280°C [20]; on the other hand, the endothermic process of pure eugenol sample between 240 and 260°C can be associated with the boiling point of the active compound (in the scientific literature eugenol b.p. = 254°C).

The size measurements of polyurethane structures with and without eugenol were done using the same parameters of the instrument and the same concentration of samples. The results of structures’ sizes are shown in figure 3 and it can be observed that the average diameter of empty structures is 241 nm, while the diameter of structures with eugenol is 289 nm. This increase can be due to the fact that the active substance can play the role of a chain extender.

The particles’ surface charge, analyzed by a Wallis Zetapotential instrument, depends on their electrophoretic mobility.

<table>
<thead>
<tr>
<th>Parameter vs. Sample</th>
<th>EU_0</th>
<th>EU_1</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH</td>
<td>6.71±0.11</td>
<td>6.82±0.17</td>
</tr>
<tr>
<td>Solubility in water, mg/mL</td>
<td>0.36</td>
<td>0.32</td>
</tr>
<tr>
<td>Solubility in ethanol, mg/mL</td>
<td>0.93</td>
<td>1.01</td>
</tr>
<tr>
<td>Solubility in acetone, mg/mL</td>
<td>1.10</td>
<td>1.11</td>
</tr>
<tr>
<td>Solubility in DMSO, mg/mL</td>
<td>1.07</td>
<td>1.09</td>
</tr>
</tbody>
</table>

No important differences between the mobility of samples EU_0 and EU_1 were obtained (fig. 4a and b). The average results of samples are: the mobility $\mu_{EU_0} = -4.85 \mu$m . cm/V . s and Zeta potential $\xi_{EU_0} = -62.33$ mV, while the mobility $\mu_{EU_1} = -4.34 \mu$m . cm/V . s and Zeta potential $\xi_{EU_1} = -55.76$ mV. These results indicate that both samples present a very good stability against the tendency to form clusters; the literature indicate that the specific range of Zeta potential of colloidal structures unstable to agglomeration is between -30 and +30 mV [21].

The polyurethane structures present an UV-Vis maximum absorption around 300 nm with a small shift due to the addition of eugenol (fig. 5). Eugenol did not present a sharp peak, but its absorbance increase around 210 and 285 nm. The first wavelength value was chosen to evaluate to quantity of free drug in order to establish the entrapment efficiency. There were used 5 standard solutions and the calibration curve. The results indicate an entrapment efficiency equal with 67%.

Figure 6 present the evolution of some skin parameters often used in the prediction of new synthesized compounds’ toxicity. It is known that an irritative compound will increase the transepidermal water loss and the erythema (redness of skin) and will decrease the level of stratum corneum hydration.

There are two main factors that are considered in all toxicological assays: (1) how the measured parameter is changed (if it increase / decrease / remain constant during the experiment) and (2) how much the measured parameter is changed (important or minimum change of values). In our experiment, all four skin parameters were modified: the transepidermal water loss, the melanin and erythema index increase; on the other hand, the hydration...
Fig. 4. The mobility of polyurethane structures from samples: (a) EU_0 and (b) EU_1.

Fig. 5. UV-Vis spectra for EU_0, EU_1 and eugenol.

Fig. 6. The evolution of skin parameters: (a) TEWL, (b) melanin.
of stratum corneum decrease. So, the first factor is respected, and this is a condition for a positive test.

The changes of values are important and they are very different between the exposed samples and the blank in the case of toxic or irritative compounds. In our study, the modifications of skin parameters are not important (e.g.: the melanin scale is between 0-999 arbitrary units and we obtained changes just between 405 and 470, so the difference is $\Delta = 65$ units related to a 1000 scale) and these modifications are comparable with those obtained for blank. This is the reason why, we consider that these samples do not present any toxic or irritative effect.

Conclusions

Polyurethane structures with and without eugenol were synthesized using a polyaddition process combined with a simultaneous emulsification. Structures with sizes between 241 and 289 nm were obtained, and they present a good thermal stability and an average pH of aqueous solution between 6.71 and 6.82. The evaluation of encapsulation efficiency, based on UV-Vis absorption of free drug related to the quantity of eugenol added to synthesis, revealed a 67% of active agent entrapped inside the polyurethane structures. The bioevaluation of products, based on different assays on mice skin, suggest that these products are safe to use in oral therapies.

Acknowledgements: This work was financially supported by UEFISCDI (research contract PN-III-P2-2.1-BG-2016-0455) / 122BG and by Victor Babes University of Medicine and Pharmacy Timisoara, Romania (research contract no. 15250/2012) and Development of existing infrastructure and creation of new infrastructure.POSCCE-A2- O2.2.1- 2013- 1,Center of Genomic Medicine University of Medicine and Pharmacy Victor Babes Timisoara.

References

21. SALOPEK, B., KRAŠI, D., FILIPOVIĆ, S. Rudarsko-geoloiko-
aftnizbornik, 4, 1992, p. 147

Manuscript received: 7.10.2017

Fig. 6. The evolution of skin parameters: (c) erythema, (d) hydration