

QbD based control strategy of loratadine nanosuspensions and dry nanoparticles stabilized by Soluplus®

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Abstract:

The preparation of nanosuspensions has been introduced as a well-defined method to enhance the solubility and dissolution of poorly water-soluble drugs. The aim of this study was to evaluate the feasibility of using Soluplus® as a stabilizer for loratadine nanosuspensions. The concept of Quality by design (QbD) was followed particularly to link **the critical material parameters (CMPs)** and the critical process parameters (CPPs) with the required critical quality attributes (CQAs) and risk assessment (RA) to select the optimized critical **material** and process parameters. The ultrasonic-assisted precipitation method was selected to prepare the nanosuspensions with different concentrations of Soluplus®. Particle size, polydispersity index (PDI), solubility and dissolution were set as the main CQAs. Soluplus® successfully produced loratadine nanosuspensions with particle size ranging between 168.3-245.35 nm and PDI in the range of 0.12 and 0.25. The freeze dried sample with 0.6% Soluplus® (DLNS3) showed an amorphous status of loratadine with particle size and PDI in the range of 220 ± 6.23 and 0.21 ± 0.02 , respectively. Contact angles, surface free energy, and polarity measurements showed an enhancement of the hydrophilic properties of DLNS3. DLNS3 displayed 121-fold

saturation solubility and released approximately 57% of loratadine within 15 min. The effects of **CMPs and** CPPs on the CQA were expected by the QbD approach.

Key words: Loratadine nanosuspension, quality-by-design, risk assessment, precipitation

20 **Introduction:**

Recently, particle size reduction to the submicron level has been proved as one of the most efficient methods to enhance solubility and dissolution, hence the bioavailability of poorly water-soluble drugs. Nanosuspension (NS) is an essential part of nanotechnology that produces particles at the
25 submicron level stabilized by a suitable type and amount of stabilizer(s). Generally, two methods can be applied for producing NS; the top-down and the bottom-up method with the possibility of combining both methods. On the contrary to the top-down, the bottom-up method is based on building up the particles from the molecular state of the drug [1,2].

30 Precipitation assisted by ultrasonication is a commonly used as bottom-up method. The preparation of NS is usually followed by drying procedures, such as spray drying and freeze drying, to ensure long-term stability. All the parameters related to these processes could have significant effects on the properties of NS, such as particle size, particle size distribution, and stability

35 in addition to the properties of the dry particles, such as re-dispersibility, particle size, solubility, etc [3–6].

Loratadine (LOR), a second-generation histamine H₁ receptor antagonist, is the most frequently prescribed antihistamine drug for the treatment of allergic conditions. LOR belongs to class II of the biopharmaceutical
40 classification system and has a pH-dependent solubility, as a consequence, it shows low and variable bioavailability. Many techniques have been adopted to enhance the solubility and dissolution of LOR, including solid dispersion, inclusion with β -cyclodextrin derivatives, and micellar solubilization [7–11].

On the other hand, various drug delivery system such as microparticulated
45 and nanoparticulated systems has been introduced to overcome the inconvenience of the currently used systems [12].

In a multivariate production process, all the parameters of the different operations should be cautiously selected and their effects on the final product must be assessed. In the case of preparing nanosuspension by
50 precipitation ultrasonication, all the parameters related to these processes must be evaluated in addition to the drying procedure. The Quality by Design (QbD) approach supports the development of products with a predefined quality based on knowledge and risk assessment (RA). For QbD-

based development, it is necessary to identify the critical quality attributes
55 (CQAs) which critically influence the predefined quality target product
profile (QTPP). Moreover, the critical material and critical process
parameters (**CMPs and CPPs, respectively**) with high impacts on CQAs
must be defined [13,14].

In practice, the identification of CQAs, **CMPs** and CPPs is based on the
60 previous practice, and literature knowledge and experience. In a recent study
of our team, we evaluated the preparation of loratadine (LOR)
nanosuspension by the precipitation ultrasonication method, with the use of
the most commonly applied stabilizers, including polymers
(hydroxypropylmethylcellulose (HPMC), polyvinylpyrrolidone (PVP-
65 K25)), nonionic surfactant (Tween 80, Pluronic F68) and ionic surfactant
(sodium lauryl sulfate (SLS)) as single or combined stabilizers. In the
present paper, the authors emphasize the impacts of **CMPs**, CPPs and the
effect of using a new material as a stabilizer, e.g. Soluplus[®], on the
production of NS for loratadine. The aim was to demonstrate the efficiency
70 of applying the QbD concept in reducing the experimental trials and
predicting the results based on previously determined the **CMPs** and CPPs.
Moreover, this study aimed to explore further possibilities for LNS

stabilization with Soluplus[®] and evaluate its effect on the CQAs of LNS and DLNS.

2. Materials and methods

75 2.1 Materials

Loratadine was purchased from Teva Ltd. (Budapest, Hungary). Soluplus[®] was purchased from BASF (Ludwigshafen, Germany). Ethanol was supplied by Spectrum-3D (Debrecen, Hungary) and trehalose dihydrate was supplied by Sigma-Aldrich (New York, USA). Water was purified by double
80 distillation.

2.2 Methods

2.2.1 Determination of QbD elements (CQAs, CPPs, and RA)

Based on prior knowledge, previous studies, preliminary experiments, and data from relevant literature, CQAs, CPPs were determined for producing
85 LNS. Previous studies led to the selection of particle size, polydispersity, and zeta potential as CQAs. In the case of DLNS, particle size, polydispersity index, solubility, and dissolution properties were determined as CQAs.

The RA was performed with Lean QbD Software® (2014QbD Works LLC.,
90 Fremont, USA). According to this software, the connections between CQAs,
CMPs and CPPs were evaluated and rated on a three-level scale. This scale
reflects the impact of their interaction on the product as high (H), medium
(M) or low (L). Further, Pareto charts were generated by the software,
presenting the numeric data and the ranking of CQAs, **CMPs** and CPPs.

95 **2.2.2 Preparation of loratadine nanosuspension and dried nanoparticles**

LNSs were prepared with the precipitation-ultrasonication method. LOR was
dissolved in ethanol, while Soluplus® was dissolved in water. Both solutions
were filtered through a 0.45µm filter (FilterBio PES Syringe Filter, Labex
Ltd., Budapest, Hungary). Afterwards, the drug solution was rapidly
100 introduced into pre-cooled antisolvent under sonication using a UP 200s
Ultrasonic processor (HielscherUltrasonics GmbH, Germany) for 30 min at
4 °C and 50% amplitude. The temperature of sonication was controlled by
JulaboF32 (JULABOGmbH,Germany). LNSs were stirred at room
temperature for 24 h to remove the organic solvent. The selected LNS
105 sample was lyophilized with 5% (w/v) trehalose to produce DLNs by using a
ScanVac, CoolSafe™ freeze-dryer (LaboGene, Denmark). The selected LNS
was lyophilized at -40°C. The solvent was sublimed under a pressure of
0.01 mbar for 36 h.

2.2.3 Preparation of physical mixtures

110 Physical mixtures (PMs) corresponding to the composition of LNS were prepared by blending LOR and Soluplus[®] in a Turbula mixer (Turbula System Schatz; Willy A. Bachofen AG Maschinenfabrik, Basel, Switzerland) using 60 rpm for 10 minutes with a LOR: Soluplus ratio of 1:2.4, w/w (PM1). Moreover, PM with trehalose was prepared to figure out
115 the effect of the cryoprotectant (PM2) with a LOR: Soluplus: trehalose ratio of 1:2.4:20, w/w.

2.2.4 Particle size characterization

The MPS, PDI, and ZP of LNSs were measured by dynamic light scattering using Malvern Nano ZS zetasizer (Malvern Instrument, UK), with water
120 used as dispersant and refractive index set to 1.62. The samples were adequately diluted with distilled water and measured at 25°C and pH 5.77. 12 parallel measurements were carried out.

2.2.6 Characterization of dried nanoparticles

2.2.6.1 Scanning electron microscopy (SEM)

125 The morphology of the powder particles was investigated by scanning electron microscopy (SEM) (Hitachi S4700, Hitachi Scientific Ltd., Tokyo,

Japan) at 10 kV. The samples were coated with gold-palladium (90 seconds) with a sputter coater (Bio-Rad SC 502, VG Microtech, Uckfield, UK) using an electric potential of 2.0 kV at 10 mA for 10 min. The air pressure was
130 1.3–13.0 mPa.

2.2.6.2 X-ray powder diffraction (XRPD)

The structure of lyophilized nanoparticles and raw materials was characterized using a BRUKER D8 Advance X-ray powder diffractometer (Bruker AXS GmbH, Karlsruhe, Germany) with Cu K λ_1 radiation ($\lambda =$
135 1.5406 Å) and a VÅNTEC-1 detector. The powder samples were scanned at 40 kV and 40 mA, with an angular range of 3° to 40° 2 θ , at a step time of 0.1s and a step size of 0.01°.

2.2.6.3 Differential scanning calorimetry (DSC)

The thermal analysis was carried out using a differential scanning
140 calorimeter (Mettler Toledo DSC 821^e, Mettler Inc., Schwerzenbach, Switzerland). About 3–5 mg of powder was accurately weighed into DSC sample pans, which were hermetically sealed and lid pierced. An empty pan was used as a reference in an inert atmosphere under constant argon purge. The samples were examined in the temperature interval of 25–300 °C at a
145 heating rate of 5 °C min⁻¹.

2.2.6.4 Surface free energy and polarity investigation

The contact angle, surface free energy (SFE) and polarity of the samples were measured. 0.15 g of sample was pressed at 1-ton hydraulic press to pastille (PerkinElmer Hydraulic Press; PerkinElmer Inc., Waltham, MA, USA). Then, the surface of the pastilles was dripped with polar and non-polar solvents. The contact angle was detected for 30 seconds with DataPhysics OCA 20 device (DataPhysics Inc. GmbH, Filderstadt, Germany), and then Wu correlation was used. The solvents were distilled water ($\gamma_p=50.2$ mN/m, $\gamma_d=22.6$ mN/m) and diiodomethane ($\gamma_p=1.8$ mN/m, $\gamma_d=49$ mN/m).

2.2.6.6 Dissolution studies

The dissolution tests were performed using the modified paddle method (USP dissolution apparatus, type II Pharma Test, Hainburg, Germany). Samples were tested in 100 mL of PBS (pH 7.4). The paddles were rotated at 100 rpm at 37 °C. At a predetermined time, 5-mL aliquots were withdrawn and filtered. The concentration of LOR was measured spectrophotometrically (Unicam UV/VIS Spectrophotometer, Cambridge, UK) at λ_{\max} 248 nm.

3. Results and discussion

165 **3.1 Knowledge space development for the precipitation ultrasonication method**

The development of knowledge space could visualize the overall manufacturing process with respect to the selection of CPPs, and the definition of the required CQAs [13].

170 To adapt to QbD-based development principles, the first step was to define the required CQAs (Table I), followed by the identification of the CPPs and affect the CQAs considering particle size the main factor based on the definition of nanosuspension and on its consequences on the other CQAs, such as solubility and dissolution (Table II). Afterwards, the RA
175 relationships between CQAs and CPPs in addition to the numeric data of the critical factors and their ranking (Pareto charts) were determined (Fig 1) to finally select the optimized **CMPs** and CPPs that support the achievement of the required CQAs. **(Table III) shows the optimized CMPs and CPPs based on our previous studies [13].**

180 **3.2 Preparation of nanosuspensions and dry nanoparticles**

MPS, PDI and ZP results are summarized in Table IV. The freshly prepared LNSs showed a significant reduction in MPS at the range of 168.3 and 245.35 nm monodispersion with low PDI index. Soluplus[®] produced LNS

with the lowest particle size compared to the commonly used stabilizers
185 [14]. Soluplus[®] is an amphiphilic compound that interacted with the
nonpolar surface area of LOR and covered the newly formed surfaces,
providing steric hindrance to prevent recrystallization from the solution and
aggregation of the primary particles. Higher concentrations of Soluplus[®]
could stabilize the NS more effectively due to weak Ostwald ripening as the
190 drug will diffuse slowly from the formed micelles [15].

The MPS of the three samples were preserved within the nanorange (Table
V). LNS3 with the smallest MPS was selected for further characterization as
dry nanoparticles (DLNS3).

DLNS3 showed a MPS in the order of 220 ± 6.23 nm, PDI range 0.21 ± 0.02
195 and ZP of -23.8 ± 4.4 mV after constitution in 5 mL of distilled water.

3.3 Morphology

SEM images (Fig. 2) showed that LOR had an irregular rod-like crystal
shape with a particle size above 5 μm and some aggregation emphasized the
broad distribution of the raw drug. DLNS3 had spherical particles at the
200 nanosized scale embedded within the carriers. The effect of stabilizer type
on morphology was expected and confirmed here as Soluplus[®] produced a

spherical shape, while F68 and F68 with PVP-K25 produced short rod morphologies [14].

3.4 Structural analysis (DSC and XRPD)

205 The thermal behaviors of the pure materials and DLNS3 are shown in Fig.3. LOR showed a single narrow peak at 134.7 °C corresponding to its melting point. The Soluplus[®] thermogram showed a wide peak, which represents water evaporation. PMs showed the crystalline state of LOR, while the absence of a LOR peak in DLNS3 indicates the presence of LOR in an
210 amorphous state. Fig 4 shows the XRPD spectra of raw materials, PMs and DLNS3. The characteristic crystalline peaks disappeared in the pattern of the dry DLNS3. This revealed the presence of LOR in its amorphous state.

3.5 Surface free energy and polarity investigation

Table VI lists the results of polarity and contact angles. Water contact angle
215 decreased for PM1, and DLNS3 showed the lowest value, indicating the highest wetting properties. When diiodomethane was used instead of water, DLNS3 showed an increase to 23.1° compared to approximately 13.5 of LOR and PM1. The increase in SFE suggests the conversion of the surface toward higher polarity. These results were confirmed by measuring the
220 polarity%, where DLNS3 showed the highest value (33.65%).

3.6 Solubility and dissolution

DLNS3 exhibited a marked increase in the solubility and dissolution of LOR (Table VII). It showed $59.39 \pm 5.18 \mu\text{g/mL}$ with 121-fold enhanced solubility compared to LOR that showed a solubility of $0.49 \pm 0.001 \mu\text{g/mL}$. Two main factors are responsible for such enhancement; the reduction in particle size and the wettability of the polymers. The dissolution of nanoparticles is enhanced based on Noyes–Whitney equation [16]. Moreover, Soluplus[®] can create a hydrophilic environment around the drug nanoparticles. PMs showed higher solubility than LOR due to the wettability enhancement of Soluplus[®]. However, trehalose slightly affects the solubility of LOR as the solubility of PM2 was comparable to that of PM1.

Fig.5 shows the dissolution profiles of the samples. LOR exhibited low drug release, less than 2% within the first 15 min, and the maximum release was approximately 5% after 2 h. PM1 and PM2 showed a release of 4.7 and 7% after 2 h, respectively. On the contrary, release from DLNS3 was high, approximately 57% in the first 15 min and 80% after 2 h.

Table VIII lists %DE values for different time periods in addition to MDT and RD60. At 30 min, the DE value of the drug is only 1.6% with a low value also for PMs, while DLNS3 showed a high release of 47.0%. Similar

240 increments were observed at 60 and 120 min with a maximum DE shown by
DLNS3 at 120 min (67.3%). Moreover, RD60 of DLNS3 showed an
observed enhancement compared to PMs. On the other hand, MDT showed a
maximum reduction with DLNS3. which emphasized the faster dissolution
of the nanoscale formulation.

245 **Conclusion**

QbD showed an efficient tool for predicting the product's quality. The use of
risk analysis for selecting high-risk factors and the further evaluation of
those factors save time and costs by providing the visual identification of
high-risk factors. The high impact relationships between **CMPs**, CPPs and
250 CQAs that were suggested by the QbD based approach were proved by
studying the effects of changing the stabilizer type. Compared to the
previously used stabilizers (e.g. HPMC, PVP-K25, F68, Tween 80 and
SLS), Soluplus[®] showed an expected difference in particle size, particle size
distribution, zeta potential, morphology, dissolution and solubility with
255 preferred effects related to lower particle size, higher zeta potential, thus
stability, higher dissolution rate and immense solubility enhancement.

Declaration of interest

The authors report no conflicts of interest related to this work.

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