# Journal of Materials Chemistry B

Materials for biology and medicine



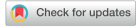
ISSN 2050-750X



### Journal of Materials Chemistry B



PAPER View Article Online
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**Cite this**: *J. Mater. Chem. B*, 2019, 7. 3958

## Glutathione from recovered glucose as ingredient in antioxidant nanocapsules for triggered flavor delivery†

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Side streams from modern lignocellulose biorefineries have found value-added applications in various industries ranging from food to medical. Here, bioproduction of glutathione from glucose recovered from man-made cellulose fiber production was investigated. Rayon fibers were enzymatically hydrolyzed and the resulting glucose and Zn in the hydrolysate were successfully used for glutathione (15.5 mg L<sup>-1</sup>) production by an engineered strain of Saccharomyces cerevisiae. Next, out of reduced glutathione (GSH) in combination with human serum albumin (HSA) and silk fibroin (SF), nanocapsules were developed. Production of HSA/SF/GSH nanocapsules was further optimized by experimental design and the resulting nanocapsules were characterized by particle size, zeta potential, chemical properties (secondary structure ratios, crosslinking, and release kinetics) and thermal stability. An average hydrodynamic radius of  $462.72\pm73.36$  nm and average zeta potential of  $-13.67\pm0.01$  mV were obtained by optimization using an experimental design approach. Increasing secondary structure ratios for HSA/SF/GSH nanocapsules indicated the successful integration of GSH into the nanocapsule shell by ultrasound induced selfassembly. Regarding possible future application as a cosmeceutical, flavor substances were encapsulated, and the release kinetics of flavor substances were studied, resulting in pH- and viscosity-dependent maximum release rates of 40.45  $\pm$  0.35% for menthol and 38.60  $\pm$  2.07% for raspberry ketone. Additionally, the radical scavenging properties of the system were evaluated, showing increased scavenging for produced HSA/SF/GSH nanocapsules compared to controls. Therefore, HSA/SF/GSH nanocapsules are seen as a promising new system in cosmeceutical approaches.

Received 8th March 2019 Accepted 1st May 2019

DOI: 10.1039/c9tb00473d

rsc.li/materials-b

#### Introduction

Side streams from modern lignocellulose biorefineries have found applications in various areas, ranging from food to medical.

For example, the hemicellulose (*i.e.* xylan) fraction resulting from rayon manufacturing is converted to xylitol (*e.g.* DuPont/Lenzing AG (Austria)). Xylitol is then used in the food and cosmetic sectors, *e.g.* as artificial bulk sweetener in chewing gums or toothpastes inhibiting salivary bacterial growth (caries-inhibitor). In a similar approach, monolithic tetracycline-containing cellulose acetate fibers based on recovered cellulose, possibly in future originating from rayon, have already been used for controlled delivery in periodontal therapy. 4,5

In the last decade, estimated total world-wide fiber consumption was 102 million metric tons. <sup>6</sup> Of these, man-made cellulosic (MMC) fibers account for more than 10 million metric tons, with an average annual growth of more than 2%. In fact, rayon is considered a future substitute for cotton materials, due to the arable land and water demand problems related to cotton production. Therefore, rayon production is expected to grow by 9.1% until 2020. <sup>6-9</sup> However, pre- and post-consumer waste MMC fibers are usually landfilled <sup>10</sup> or chemically degraded using highly concentrated chemicals such as sulfuric acid.

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<sup>†</sup> Electronic supplementary information (ESI) available: Details regarding the structure of the flame retardant pigment, detailed information on the used HPLC gradients, detailed information of the experimental designs DoE 1 and 2 including the corresponding results (ANOVA tables, response contour plots, coefficient plots), ATR-FTIR spectra and the correlated secondary structure ratios, chemical structures of the flavor substances, results of DLS based temperature stability analysis and information of artificial saliva viscosities. See DOI: 10.1039/c9tb00473d

Recently, complete recovery of glucose from rayon fibers using cellulase enzymes was demonstrated. 11 In turn, glucose could serve as a carbon source in bioproduction of value-added biomolecules with possible applications in the food or medical sector, which is the focus of this study. 12,13

Glutathione ( $\gamma$ -L-glutamyl-L-cysteinylglycine, GSH) is a natural tripeptide which is generally commercially produced by fermentation of the yeast Saccharomyces cerevisiae. GHS is increasingly used in medicine, the food and cosmetic industries due to its antioxidative properties, i.e. radical scavenging. Consequently, world-wide annual production exceeded 200 tons, 14 making both efficient production systems and cheap growth substrates important for economical processes. In fact, the commercial sector of cosmetics and cosmeceuticals is currently rapidly growing. Cosmeceuticals are defined as cosmetic products based on biologically active substances.<sup>15</sup> In general, modern systems are based on stimuli-responsive delivery of active ingredients, 16,17 where various parameters, such as temperature, ionic strength or pH, can be used as triggers. 18 Human serum albumin (HSA) and silk fibroin (SF), naturally derived from the cocoons of Bombyx mori, have been reported to show required properties for targeted delivery, such as water solubility, thermal stability (for HSA)19 and natural pH response.20 Tallian et al.21 successfully combined these properties with the development of HSA/SF nanocapsules for targeted delivery at lysosomal pH of 4.5 for the treatment of inflammations.

In the present study, GSH produced from glucose recovered from MMC rayon fibers (Rf) was combined with HSA and SF for production of nanocapsules with antioxidative properties. These properties suggest a possible new approach for gingivitis and periodontitis treatment support, as a significantly lowered pH (down to 5)<sup>22</sup> can be targeted as stimulus for release of flavour substances such as menthol or raspberry ketone. Encapsulation properties of flavour substances and their effect on the pH-responsive release behaviour in different artificial salivas were also investigated to simulate possible application for gingivitis and periodontitis treatment.

#### Results and discussion

#### Enzymatic hydrolysis of rayon fibers and glutathione production

Various components contained in side-streams of lignocellulose processing have found value-added applications. One example is xylitol produced from vegetable waste, rice straw or corncob using Candida sp. 23 and then used in chewing gums as a softener 24 and for salivary bacterial inhibition.<sup>25</sup> Similarly, residues from rayon production can provide a source of glucose after enzymatic hydrolysis and additionally contain the Zn required in glutathione production.

In this study, the enzyme cocktail Cellic CTec3 was used to hydrolyse rayon fibers. With an enzyme loading of 5 kFPU per g, 30 g of rayon fibers were completely hydrolysed, yielding a concentration of 90 g L<sup>-1</sup> of glucose. The recovered glucose solution contained some Zn (15.5 mg L<sup>-1</sup>) from rayon fiber

production. It is well known that zinc is essential in small concentrations as a redox cofactor and structural component of proteins. In fact, zinc is estimated to be necessary for the functions of nearly 3% of the S. cerevisiae proteome. Although zinc is not redox active under physiological conditions, excess zinc can be toxic to cells. 26,27

Hence, in this study, a control of commercial glucose and 15.5 mg L<sup>-1</sup> ZnSO<sub>4</sub> supplement was analysed to observe possible beneficial or toxic effects of the zinc in glutathione production. No significant difference in growth of S. cerevisiae based on OD<sub>600</sub> was found when recovered glucose was used as carbon source compared to the two controls containing pure glucose and pure glucose plus Zn. After 24 h of incubation, the OD<sub>600</sub> was slightly higher for the control without the zinc, which changed, however, after 48 h of fermentation when an OD<sub>600</sub> between 25 and 28 was reached (Fig. 1A). These data showed no toxic effect of Zn; rather, a slight biomass concentration (OD<sub>600</sub>) was found after 48 h, probably due to the role of Zn as redox cofactor.

Glucose consumption was similar regardless whether pure or recovered glucose was used as the substrate. From a starting concentration of 20 g  $L^{-1}$  ( $t_0$ ), the glucose was almost completely consumed after 24 h in all the performed experiments. The levels of reduced (GSH) and oxidized (GSSG) glutathione were measured for the two strains of S. cerevisiae used. Regarding the sum of the reduced and oxidized forms, for both strains, a slightly higher level of glutathione was detected in the samples with the zinc supplement. In this case, the zinc level is not high enough to show a toxic effect but rather supports the S. cerevisiae metabolism for glutathione production.<sup>27</sup> A 0.34 mM concentration for strain A4 and a 0.64 mM concentration for strain A4-19 were reached after 48 h (Fig. 1B).

As previously reported, 11 S. cerevisiae can be applied for production of 2nd generation biofuels as substitutes for conventional petroleum and natural gases. The ethanol production was investigated and a level of 7 g L-1 was detected for both strains after 24 h of fermentation (Fig. 1C). The ethanol level was drastically reduced after 48 h due to evaporation from the fermentation flasks through evaporation or fermentation by S. cerevisiae.

#### Optimization of HSA/SF/GSH nanocapsules production by DoE

Next, incorporation of GSH into nanocapsules was investigated. Ultrasound assisted production of HSA/SF/GSH nanocapsules was optimized based on a production method published by Tallian et al.21 for stimuli responsive HSA/SF nanocapsules. A randomized full factorial design with 3 centre points was evaluated regarding particle size and zeta potential (ESI,† Fig. S1) to determine the influence of the tested factors on these responses. Herein, response values of zeta potential analysis ranged between -11.08 and -13.78 mV and hydrodynamic nanocapsule radii ranged from 419.20 to 664.01 nm. Both models showed a strong positive linear correlation (ESI,† Fig. S1A), with  $R^2 = 0.99$  for zeta potential (ZP) and  $R^2 = 0.93$  for hydrodynamic nanocapsule radius (radius), and a good prediction fulfilling the condition  $\Delta R^2 - Q^2 < 0.2$ –0.3 (0.07 for ZP; 0.26 for radius).

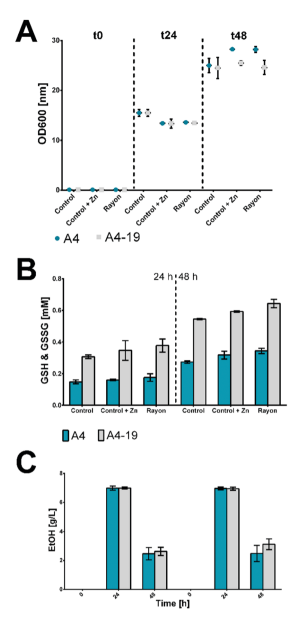


Fig. 1 (A) Growth of *S. cerevisiae* (A4 and A4–19 strains) using glucose enzymatically recovered from rayon fibers (rayon), commercial glucose (control) and commercial glucose supplemented with ZnSO<sub>4</sub> (control + Zn) as carbon sources. All experiments were performed in duplicate. (B) Glutathione production in the reduced (GSH) and oxidized (GSSG) form by *S. cerevisiae* (A4 and A4–19 strains) when grown on glucose enzymatically recovered from rayon fibers (rayon), on commercial glucose (control) and commercial glucose supplemented with ZnSO<sub>4</sub> (control + Zn) as carbon sources. All experiments were performed in duplicate. (C) Ethanol production by *S. cerevisiae* when grown on glucose enzymatically recovered from rayon fibers (rayon), on commercial glucose (control) and commercial glucose with ZnSO<sub>4</sub> supplement (control + Zn) as carbon sources. All experiments were performed in duplicate.

**∏A4-19** 

To describe the relationships of the investigated variables, obtained measurement data (Table S4, ESI†) were further fitted by multiple linear regression analysis and the fit of the model was evaluated by ANOVA tests (ESI,† Table S6). For both responses,

statistical significance of the models with a confidence level of 95% (p-value < 0.05) was achieved. Additionally, the model lack of fit evaluation, comparing the model error to replicate error with a significance level of 5% (p-value > 0.05) was evaluated, resulting in statistical significance for both models. F-tests were used in combination with p-values to evaluate the coefficient's significance (ESI,† Table S6). Scaled and centred coefficients are displayed in Fig. S1C (ESI†), showing that the predominant interaction in the developed model for zeta potential is that between PRO(Yes) × PUR (protein addition × number of purification steps). This interaction has a significant negative effect on the zeta potential as well as the hydrodynamic radius.

Based on these results, a second DoE (DoE 2) was designed, where the protein addition of HSA and SF for capsule formation was kept constant at 4.2 mg as indicated above in materials and methods. This was hypothesized to lead to the desired size of HSA/SF/GSH nanocapsules with low zeta potential. For possible future applications in cosmeceutical approaches, such as a radical scavenging toothpaste additive, sizes below 500 nm (hydrodynamic radius) were attempted. Here, in contrast to i.v. applications where particle sizes are focused between 100 to 300 nm, <sup>28,29</sup> radical properties corresponding with sufficient release behaviours are seen as the key properties of the designed system. As the number of purification steps (PUR) showed a significant positive effect on zeta potential and no significant effect on nanocapsule size, the factor range was narrowed from 1–10 to 1–5 purification steps.

In DoE 2, the relationship between the factors GLU (concentration GSH in mg mL<sup>-1</sup>) and PUR (number of purification steps) was evaluated by multiple regression analysis, from which quadratic models for both responses, zeta potential (ZP) and hydrodynamic nanocapsule radius (radius), based on the experimental data (Fig. 2 and ESI,† Table S5) were obtained (eqn (1) and (2)).

$$ZP = -2.05435 + 0.00139869x_1 - 0.0003910171x_2 + 0.00279421x_1^2 + 0.00156146x_2^2$$
 (1)

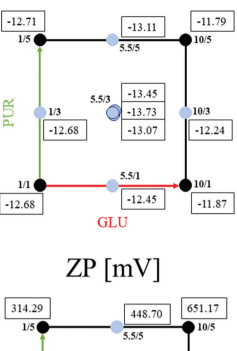
$$ZP = 540.467 + 108.024x_1 + 68.1175x_2 + 53.315x_1x_2 + 3.17916x_1^2 - 139.318_2^2$$
 (2)

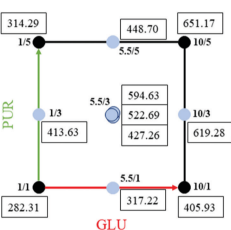
 $x_1$ : concentration GSH in mg mL<sup>-1</sup> (GLU);  $x_2$ : number of purification steps.

As reported in Fig. 2, zeta potential values between -11.79 to -13.79 mV were obtained with hydrodynamic radii between 282.31 and 651.17 nm and similar strong linear correlations when compared to DoE 1, with  $R^2 = 0.85$  for ZP and  $R^2 = 0.97$  for Radius. ANOVA evaluations resulted in statistical significance for both models with p-values < 0.05 (confidence interval of 95%), as shown in Table S7 (ESI†).  $Q^2$  values and obtained model validity values for both responses indicate that the obtained models are well fitted and suitable for prediction (Fig. 3).

Scaled and centered coefficients were plotted in Fig. 3C to visualize the effects of independent factors on the ZP and Radius responses. Here, the coefficients are half of the factor effects. In the ZP model, the interaction GLU  $\times$  PUR was removed to improve  $R^2$  and  $Q^2$ .

A4





#### Radius [nm]

Fig. 2 Optimization of HSA/SF/GSH nanocapsules regarding zeta potential and hydrodynamic radius. Two-dimensional representation of the results of the central composite face-centered design (CCF; DoE 2), where the numbers in the quadratic space represent the obtained zeta potential (ZP) values in mV or the obtained hydrodynamic radii (radius) in nm for different combinations of the factors GLU (concentration of GSH) and PUR (number of purification steps).

Regarding the quadratic interactions, GLU × GLU is significant; however, the non-significant interaction of PUR imesPUR and the non-significant factor PUR were included in the model, as both coefficients were significant for the second model. In a further step, the most robust setpoint was evaluated using MODDE, where hydrodynamic radius and zeta potential were minimized. As result, 1.8 mg mL<sup>-1</sup> of GSH and 4.6 purification steps were obtained. Therefore, optimized HSA/SF/GSH nanocapsule production was set accordingly using 5 purification steps; a zeta potential of -12.95 mV and hydrodynamic radius of 407.74 nm were predicted. These predictions correlate with the visually evaluated response contour plots shown in Fig. 3B.

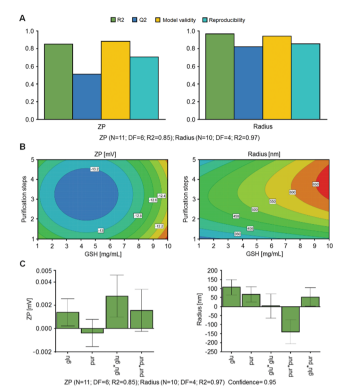


Fig. 3 Optimization (RSM) of ultrasound assisted monodisperse HSA/SF/GSH nanocapsule production via design of experiments (CCF, quadratic model). (A) Summary of fit plots for zeta potential (ZP, left) in mV and hydrodynamic particle radius (radius, right) in nm, including  $R^2$  (goodness of fit),  $Q^2$  (goodness of prediction), the model validity and the reproducibility based on three center points, where N indicates the number of experiments and DF refers to degrees of freedom. Model was fitted with MLR. (B) Response contour plots for zeta potential (ZP, left) in mV and hydrodynamic particle radius (radius, right) in nm with constant protein addition (yes). (C) Extended coefficients plots of scaled and centered coefficients for zeta potential (ZP, left) in mV and hydrodynamic particle radius (radius, right) in nm, including significant 2 factor interactions, whereby R2 refers to goodness of fit, N indicates the number of experiments and DF refers to degrees of freedom. Coefficient plots were calculated using a confidence interval of 95%.

Furthermore, the successful incorporation of GSH in the nanocapsule shells was tested during the complete optimization process using an indirect HPLC-ELSD based approach. GSH concentrations in the washing waters collected during particle production were determined. This was done to ensure that the changes in zeta potential and radius were solely based on the modifications made in the corresponding DoEs. For all samples tested, GSH concentration in the analysed washing waters was under the limit of detection, indicating that the complete amount of GSH was successfully incorporated in the HSA/SF/GSH nanocapsule shells.

#### Characterization of optimized HSA/SF/GSH nanocapsules

Based on the prediction results of DoE 2, optimized HSA/SF/GSH, as well as HSA and HSA/SF nanocapsules as controls, were produced, purified and characterized by their physical and chemical properties. Microscopic analyses were performed to investigate shape-related differences between the controls and the optimized HSA/SF/GSH nanocapsules (Fig. 4).

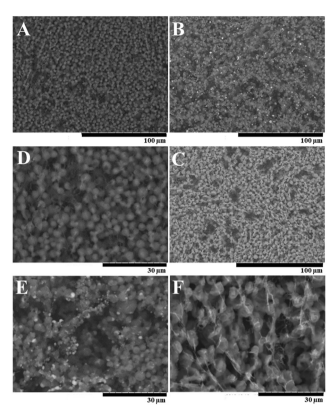
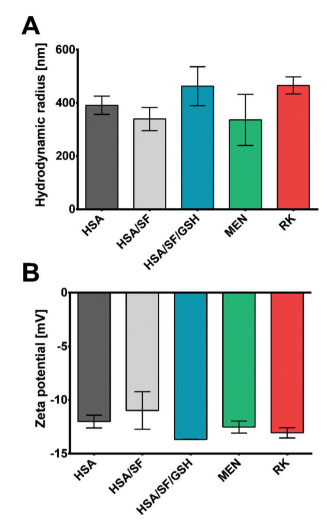


Fig. 4 Scanning electron microscopy images of (A) HSA nanocapsules, (B) HSA/SF nanocapsules and (C) HSA/SF/GSH nanocapsules at  $1000 \times \text{magnification}$  and (D) HSA nanocapsules, (E) HSA/SF nanocapsules and (F) HSA/SF/GSH nanocapsules at  $2500 \times \text{magnification}$  obtained with a SEM equipped with a cryo chamber at  $-20 \, ^{\circ}\text{C}$ .

As indicated in Fig. 4, HSA/SF/GSH nanocapsules showed a slightly bigger particle size compared to the controls; however, it was apparent that the high energy per sample surface area due to the magnification  $(2500\times)$  was affecting sample stability during the analysis. This effect resulted in visible black holes caused by nanocapsule disruption due to the electron beam. For confirmation of the differences in particle size and verification of the optimum settings for HSA/SF/GSH nanocapsules, DLS analysis was performed to investigate the average hydrodynamic radius of the monodisperse phase of each sample (Fig. 5A). HSA/SF nanocpasules showed the smallest average hydrodynamic radius (339.20  $\pm$  43.19 nm), followed by HSA (390.70  $\pm$  34.42 nm) and HSA/SF/GSH nanocapsules (462.72  $\pm$ 73.36 nm). Hence, particle sizes of all three samples were not significantly different and the predicted particle size at optimum production settings (407.74 nm) of DoE 2 was confirmed. These results correlate with previously published data by Tallian et al., 21 where hydrodynamic particle radii of HSA/SF nanocpasules containing 50% SF ranged between 339 and 507 nm. Additionally, the obtained hydrodynamic radii for HSA/SF/GSH nanocpasules fullfiled the criteria set in regard to possible future applications in cosmeceuticals, with sizes below 500 nm in radius.

The zeta potentials of these formulations were investigated for similar reasons; in fact, HSA/SF/GSH nanocapsules showed a significantly lower zeta potential ( $-13.67\pm0.01$  mV) compared



**Fig. 5** (A) DLS based hydrodynamic radii of the monodisperse fractions of HSA, HSA/SF and HSA/SF/GSH nanocapsules and menthol (MEN) and raspberry ketone (RK) loaded HSA/SF/GSH nanocapsules. (B) Zeta potential values in mV of HSA, HSA/SF and HSA/SF/GSH nanocapsules and MENand RK-loaded HSA/SF/GSH nanocapsules measured in sodium phosphate buffer pH 7.4.

to HSA ( $-12.0 \pm 0.6$  mV) and HSA/SF ( $-10.97 \pm 1.75$  mV) nanocapsules. This enhanced negativity of the zeta potential is beneficial as it leads to low serum protein binding and therefore potentially increased stability for application as a cosmeceutical, in particular for people suffering from gingivitis and/or periodontitis.30 Release properties of nanosystems with enhanced protein binding ability can be significantly decreased, as previously published by Honary & Zahir.30 In this context, low zeta potential values were further described as beneficial for prolongation of the circulation time caused by reduced RES clearance.30 Furthermore, low zeta potential values give significant information on the colloidal stability of produced HSA/SF/GSH nanocapsules, where the previously reported low colloidal stability of SF nanoparticles by Wang et al. could be improved by the new approach of incorporating GSH into the nanocapsule shells.31 The resulting relatively stable32 HSA/SF/GSH nanocapsules showed no aggregation behaviour, as can be seen in the microscopic images in Fig. 4.

As HSA/SF/GSH nanocapsule production is based on the previously published sonochemical self-assembly approach by Tallian et al., 21 the protein secondary structure changes during nanocapsule formation were further investigated. In this context, the enhanced stability of the nanoformulations obtained by application of ultrasound has previously been linked to the formation of crystalline β-sheets. 33,34 Here, the hydrophobic sequence motifs of SF (GAGAGS, 53% of entire protein sequence) play a crucial role during particle formation.<sup>21</sup>

The formation of disulfide bonds is known to be a key driver in β-sheet formation, which can be achieved between cystein residues.35 Therefore, the addition of the tripeptide GSH is hypothesized to lead to an increase in β-sheet formation during the ultrasound-induced self-assembly process. The secondary structure ratios ( $\beta/\alpha$  ratios) of HSA, HSA/SF and HSA/SF/GSH nanocapsules were investigated by ATR-FTIR analyses of lyophilized samples (ESI,† Fig. S2). HSA/SF/GSH nanocapsules showed a significantly increased  $\beta/\alpha$  ratio compared to the controls in the amide I bands (ESI,† Fig. S3), which indicates additional formation of β-sheets due to induction of covalent disulfide bonds between, for example, the cysteine residues of GSH and the free thiol groups in SF and HSA.<sup>36</sup>

#### Studies of the particle properties, the encapsulation efficiency and the release kinetics of loaded HSA/SF/GSH nanocapsules

Conventional oral administration of bioactive molecules lacks the possibility of rate-controlled substance release, which can be surmounted by the design of targeted release systems.<sup>37</sup> Although in cosmetic approaches targeted and stimuli-responsive release of certain molecules is still not the gold standard, research findings applicable in targeted delivery of bioactive molecules can also be used to achieve these desired properties in cosmeceuticals. Therefore, analysis and maximization of encapsulation efficiencies is a necessity to reduce required substance amounts during the fabrication process and to prolong and secure the desired release rates.<sup>38</sup> In the present study, two different flavour substances (menthol (MEN) and raspberry ketone (RK); see ESI,† Fig. S4 for chemical structures) were added before HSA/SF/GSH particle production. Encapsulation efficiencies were evaluated indirectly by GC-MS (MEN) and HPLC-DAD (RK), where efficiencies for MEN of 98.46  $\pm$  0.83% and for RK of 85.16  $\pm$  0.43% were achieved. Particle properties' stability was evaluated using DLS and zeta potential analysis of MEN- and RK-loaded HSA/ SF/GSH nanocapsules and compared to unloaded formulations (Fig. 5). No significant differences in average hydrodynamic radii and zeta potential could be found, which indicated that the encapsulation of the used flavour substances does not significantly affect the optimized particle production settings and properties based on the DoE. Further temperature dependent particle stability was investigated by observation of temperatureinduced fluctuations in the hydrodynamic particle radius of loaded and unloaded HSA/SF/GSH nanocapsules (ESI,† Fig. S5). All tested formulations showed slight fluctuations at higher temperature ranges from 50 to 70 °C, but no disruption or degradation process could be observed. Hence, nanocapsule stability at release-study incubation temperatures, as well as at

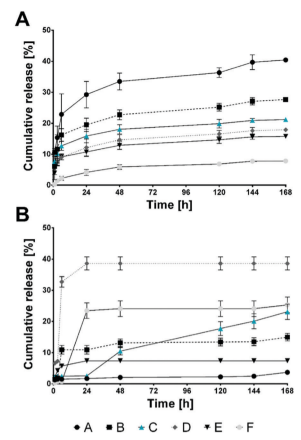


Fig. 6 Cumulative release over 168 h of (A) menthol and (B) raspberry ketone from HSA/SF/GSH nanocapsules in various artificial salivas (A-F, where A, 39 B, 44 D, 40 E41 and F are commercial certified and normed artificial salivas by Pickering laboratories and C is 100 mM sodium phosphate buffer pH 7.4).

possible storage temperatures, for future applications could be proven. Next, release studies in 4 different artificial salivas and sodium phosphate buffer with MEN- and RK-loaded HSA/SF/ GSH nanocapsules were performed and cumulative release was calculated over a time period of 168 h (Fig. 6). In the case of MEN-loaded HSA/SF/GSH nanocapsules, release was found to depend on the pH and viscosity of the artificial salivas tested. Maximum MEN release was obtained with saliva A<sup>39</sup> (pH 4.6), with 40.45  $\pm$  0.35%. As shown in Fig. 6A, the maximum release after 168 h significantly decreased with increasing pH and saliva viscosity (ESI,† Fig. S6), except in solvent C (sodium phosphate buffer pH 7.4). In the case of solvent C, the lack of media complexity combined with low viscosity visibly led to increased release compared to artificial salivas D-F. These three artificial salivas contain high concentrations of sodium carboxymethyl cellulose and, in the cases of salivas D40 and E,41 additional sorbitol, which are both substances significantly contributing to viscosity. In the release kinetics obtained for RK-loaded HSA/SF/GSH nanocapsules, the release mechanism showed dependencies on pH, viscosity, and the chemical structure of RK.

Regarding the structural effects of RK, the keto and/or OH groups are hypothesized to lead to covalent and hydrogen bonds induced during ultrasound-assisted nanocapsule production.

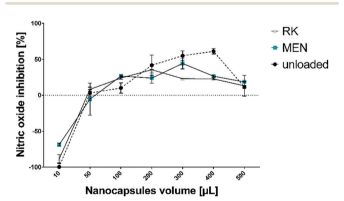
In the case of ultrasound application, the propagation of the waves occurring in the solvent leads to turbulences and microcirculations, known as acoustic streaming. 42,43

These phenomena reportedly enhance physical effects influencing further chemical processing originally limited by mass transfer. Furthermore, free radicals are generated during sonication, caused by the dissociation of vapors trapped in existing cavitating bubbles; hence, chemical reactions can be intensified, which supports the stated hypothesis. 42,43,45 The resulting bonds affect the release, resulting in a maximum release in artificial saliva  $D^{40}$  (pH 5.4) of 38.60  $\pm$  2.07%. In comparison to previously published release data by Subia and Kundu, 46 obtained maximum release rates for HSA/SF/GSH nanocapsules showed similar results. Regarding possible future application as radical scavenging and flavor delivery systems, the release rates within the observed time period of 168 h (7 days) are highly suitable. Previously published data for albumin/SF systems, however, showed release rates between 30 and 80% over a prolonged release period of 22 days. 46 Therefore, faster release was achieved due to the addition of GSH.

In general, higher media viscosities combined with lower pH seemed to be beneficial for RK release; however, for both encapsulated flavor substances, a significantly lower pH led to maximum increase. For future application of flavor-loaded HSA/SF/GSH nanocapsules as cosmeceuticals, it can therefore be expected that high release rates at lowered oral pH can be achieved. This has potential for development of new products to fight gingivitis and periodontitis, wherein product viscosity must be adjusted to support the desired flavour release based on substance structure.

#### Radical scavenging properties

During the inflammation states present in gingivitis and periodontitis, oxygen uptake in neutrophils and macrophages leads to generation of superoxide anion radicals, hydrogen peroxide and hydroxyl radicals, as well as free radicals, which are capable of damaging cell membranes or associated biomolecules. <sup>47</sup> To establish a beneficial healing environment as a future application, radical scavenging properties of unloaded and loaded HSA/SF/GSH nanocapsules, as well as of the pure substances, were tested in a nitric



**Fig. 7** Radical scavenging properties of unloaded and MEN- or RK-loaded HSA/SF/GSH nanocapsules. Nitric oxide inhibition was measured using a scavenging assay published by Alam *et al.*<sup>48</sup> and applied volumes of over 400  $\mu$ L resulted in absorbance signal inhibition.

oxide inhibition assay. Expectedly, no nitric oxide inhibition, *i.e.* radical scavenging, was observed for the pure substances (HSA, SF, MEN and RK). HSA/SF/GSH nanocapsules showed significant radical scavenging properties up to an applied volume of 400  $\mu$ L, but no significant difference between unloaded and loaded HSA/SF/GSH nanocapsules was observed (Fig. 7).

This clearly indicates that the radical scavenging effect depends on addition of GSH in the formulation, which correlated with the antioxidative properties of GSH described in literature.<sup>14</sup>

#### **Experimental**

#### **Materials and Methods**

**Materials, chemicals and enzymes.** Rayon fibers, kindly provided by Glanzstoff Industries (Lovosice, Czech Republic), had the following characteristics: twist S350 (1/m), titre 2.63 dtex, tenacity 260 cN/100 dTex, crystallinity 30% and thickness 0.018  $\pm$  0.01 mm. The rayon fibers contained about 80% w/w of cellulose-based material and 20% w/w of the phosphor-based flame retardant pigment (FRP) Exolit5060.

The enzyme preparation used, Cellic CTec3 from Novozymes (Bagsværd, Denmark), was Pointner-Rothschädl, (Salzburg, Austria). *Bombyx mori* silk cocoons were purchased from the Academy of Agricultural Sciences, Sericulture and Agriculture Experiment Station (Vratsa, Bulgaria). Human serum albumin (HSA), reduced form L-glutathione (GSH), menthol and raspberry ketone were purchased from Sigma Aldrich (Saint Louis, US) and used as received. All chemicals were of analytical grade. Artificial saliva for medical and dental research was purchased from Pickering Testing Solutions (Mountainview California, US).

Cellulase assay. A filter paper assay (FPA) was used to detect the total activity of the Cellic CTec3 enzyme preparation, as suggested by the International Union of Pure and Applied Chemistry (IUPAC) with slight modifications as previously reported. The reducing sugars were measured at 540 nm and the activity given as "filter-paper units" (FPU)  $\mu$ mol liberated per milliliter of original undiluted enzyme solution.  $^{50,51}$ 

Hydrolysis of the rayon fibers and sugars quantification. Rayon fibers (Rf) were cut and dried overnight in an oven at  $100\,^{\circ}$ C to remove humidity; after, Rf were equilibrated in a desiccator.  $30\,$ g of rayon was hydrolyzed using Cellic CTec3 (5 kFPU g $^{-1}$  of rayon) in 250 mL of 50 mM citric acid buffer at pH 4.8 and 50 °C (optimum temperature of the enzyme preparation). The incubation was carried out in a 500 mL flask in an orbital shaker at 150 rpm (IKA KS 4000 IC control, Staufen, Germany).

The hydrolyzed solution was centrifuged (Eppendorf Centrifuge 5810, Hamburg, Germany) at 25  $^{\circ}$ C for 10 min at 10 000 rpm to recover the water-insoluble FRP. The supernatant was used to quantify the released sugars. The samples for sugars determination were prepared as reported in Vecchiato *et al.*<sup>11</sup> The amount (%) of sugars was calculated using the following equation:

$$\% = \frac{(C_{\rm t} - C_0) \times V_{\rm t}}{R_{\rm i}} \times 100 \tag{3}$$

 $C_t$ : total glucose/sugars concentration (g L<sup>-1</sup>);  $C_0$ : blank glucose/sugars concentration (g  $L^{-1}$ );  $V_t$ : total volume (L);  $R_i$ : initial rayon amount (g).

Glucose recovery. After hydrolysis of the rayon fibers and glucose quantification, the supernatant was ultra-filtered using an ultrafiltration membrane (Vivaflow50, 5000 MWCO) (Sartorius, Göttingen, Germany), then passed through another filtration step with 0.1 µM vacuum filters (EMD Millipore, Austria). Glucose was quantified again via HPLC as already described to have a final concentration prior to fermentation.

Inductively coupled plasma-mass spectroscopy (ICP-MS). Zinc (Zn) released from the rayon fibers after enzymatic treatment at was quantified different time points and after the ultrafiltration step via ICP-MS Agilent 7500 (Merck GmbH, Vienna, Austria) and ICP-OES Varian (Carl Roth, Karlsruhe, Germany). 5–1000  $\mu g L^{-1}$ Zn and Mn calibration ranges were considered, according to ÖNORM EN ISO 17294-2.53 The samples were diluted in 1% HNO<sub>3</sub> water (dilution factor 1:100).

Saccharomyces cerevisiae cultivations and glutathione production. The diploid strains of Saccharomyces cerevisiae A4 and A4-19 (collection Organobalance) were used for glutathione production. The strains were stored at -80 °C in a 10% v/v glycerol solution. 250 mL shaking flasks were used for S. cerevisiae precultures in a YPD medium (1% yeast extract, 2% peptone, 2% D-glucose) overnight at 30 °C and 180 rpm. The pre-culture pellet was collected by centrifugation, washed 3 times in MQ-water and re-suspended in 10 mL of MQ-water. The culture was re-inoculated in 20 mL of WMVIII medium (Table S1, ESI†) in 500 mL shaking flasks (baffled Erlenmeyer flasks) to give an OD (600 nm) of 0.1 in, then was incubated at 30 °C and 180 rpm. The pH of the medium was set at 5 with 2 M NaOH. A control containing 15 mg mL<sup>-1</sup> ZnSO<sub>4</sub>, as expected in glucose recovered from rayon, was analyzed. Growth was monitored by determining the optical density at 600 nm and the cell dry weight (CDW). Each cultivation was performed in duplicate.

The cell dry weight was measured by collecting 2 mL of culture, harvesting the cells, and then washing them with 0.1 M HCl. The pellet was then transferred to a glass tube and dried at 100 °C for 72 h. Each reaction was performed in duplicate.

Determination of glutathione content. Glutathione was quantified by HPLC analysis (Shimadzu Corporation, Kyoto, Japan) with a refraction index detector (RID-10A, Shimadzu). From each cultivation, 5 mL samples were collected at 0, 24 and 48 hours and centrifuged at 3200 rpm and 4 °C for 10 min; then the pellet was washed with 5 mL of 0.9% NaCl. Another centrifugation step was performed at the same conditions mentioned above and then the pellet was re-suspended in 5 mL of 0.1 M phosphoric acid. The samples were incubated for 3 min at 75 °C in a water bath and then cooled on ice for 10 min. Another centrifugation step was performed and the resulting supernatant was used for HPLC analysis. In the HPLC system, separation was achieved on an Ascentis Express RP-Amide column (150 mm  $\times$  3.0 mm; 2.7 µm; Supelco analytical, Bellefonte, PA, USA) equipped with a guard column (5 mm  $\times$  3.0 mm; 2.7  $\mu$ m; Supelco analytical, Bellefonte, PA, USA). The detection of total glutathione, which included the reduced (GSH) and oxidized (GSSG) forms,

was performed at 200 nm with a photodiode array detector (SPD-M20A, Shimadzu Corporation, Kyoto, Japan). The column was operated at 25 °C, a 0.4 mL min<sup>-1</sup> flowrate of 25 mM  $NaH_2PO_4$ , and pH = 3 ( $H_3PO_4$ ). 5 µL of each sample was injected for total glutathione analysis.

Determination of glucose and ethanol content. The concentrations of glucose and ethanol in the culture were determined by HPLC analysis (Shimadzu Corporation, Kyoto, Japan) with a Phenomenex Rezex ROA column (300 mm × 7.8 mm) and a refraction index detector (RID-10A, Shimadzu Corporation, Kyoto, Japan). The column was operated at 60 °C, a 1 mL min<sup>-1</sup> flow rate and 0.004 M H<sub>2</sub>SO<sub>4</sub> as mobile phase. HPLC samples were prepared by mixing 900 µL of sample with 100 µL 0.04 M H<sub>2</sub>SO<sub>4</sub>. The mixture was filtrated through 0.20 μm RC membrane filters and 10 µL was injected for analysis.

Silk fibroin extraction from cocoons of Bombyx mori. Silk fibroin (SF) was extracted from cocoons of Bombyx mori according to the protocol published by Rockwood et al. 54 with slight modifications. Briefly, silk cocoons were cut into small pieces and boiled in 0.02 M sodium carbonate solution for 30 min. Resulting degummed SF was washed with ultrapure water and dried overnight at room temperature. SF fibers (20% w/v) were dissolved in a 9.3 M lithium bromide solution at 60 °C for 4 hours. The obtained SF solution was dialyzed against ultrapure water for 2.5 days to achieve a SF solution with conductivity similar to ultrapure water. Purified SF solution was obtained by centrifugation in two steps and stored at 4 °C until further use.

Optimization of HSA/SF/GSH nanocapsules production with DoE. Nanocapsule production is based on the sonochemical method published by Tallian et al.,21 where the method functioned as a starting point for optimization using an experimental design approach. Proteins (ratio of 50% HSA and 50% SF) were added to 2 mL of 100 mM potassium phosphate buffer pH 8.0 in a total amount of 4.2 mg and GSH was added to the protein solution (see corresponding concentrations in the experimental design for details). Subsequently, 1.3 mL of n-dodecane was added as organic phase and the mixtures were sonicated using a Branson Sonifier 250 (Emerson, Saint Louise Missouri, US) equipped with a 13 mm diameter disruptor horn. Standard settings applied during sonication for particle production were a total sonication time of 2 min with an amplitude of 10%, where a pulse on time of 1 s was followed by a pulse off period of 2 s. After sonication, phase separation of the emulsions was obtained by storing the mixtures at 4 °C for 12 hours. Then, the organic phase was removed and the nanocapsules were washed based on the settings (number of purification steps) of the experimental design with 100 mM sodium phosphate buffer pH 7.4. The supernatant was removed after each washing step and the nanocapsules were stored in 1 mL of washing buffer at 4 °C until further use or analysis. As controls, HSA and HSA/SF nanocapsules consisting of either solely HSA or HSA and SF were also produced.

Experiments were designed and evaluated using MODDE (Umetrics, Umeå, Sweden) software with three factors and two responses in the first step. Three center points were added to the full factorial design, which resulted in a total of 11 randomized experiments. Based on the achieved results, a second design set

Table 1 Coded and uncoded levels of quantitative and qualitative independent factors of the particle production process optimization for DoE 1 (screening) and DoE 2 (RSM)

DoE 1				
Quantitative				
		Levels		
Independent factor	Abbreviation	-1	0	1
Concentration GSH Purification steps	GLU PUR	1 1	5.5 5.5	10 10
DoE 1				
Qualitative				
Independent factor	Abbreviation	Settings		
Protein addition	PRO	Yes		No
DoE 2				
		Levels		
Independent factor	Abbreviation	-1	0	1
Concentration GSH Purification steps	GLU PUR	1 1	5.5 3	10 5

up using two factors and two responses similar to the initial design (Table 1) was designed and performed.

Nanocapsules size determination. The size of HSA, HSA/SF and HSA/SF/GSH nanocapsules was determined using a Dynapro Nanostar (Wyatt Technology Europe, Dernbach, Germany). Size measurements were performed with non-diluted samples over a measurement time of 30 min at 21  $^{\circ}$ C with 20 acquisitions of 2 s acquisition time each, based on a previously developed method by Wyatt Technology Europe published by Tallian *et al.*<sup>21</sup> Average hydrodynamic radii ( $R_{\rm H}$ ) were evaluated for the initial third of the measurement time and the measurement time between 10 and 30 min, as well as over the complete measurement period for each sample.

Laser–Doppler velocimetry. Zeta potential measurements for the evaluation of colloidal stability were performed using a Zetasizer Nano ZS (Malvern Instruments GmbH, Herrenberg, Germany) equipped with folded capillary cells. HSA/SF/GSH nanocapsules were diluted (15×) and measured in 100 mM sodium phosphate buffer pH 7.4 three times in automatic mode for each sample (fast field reversal technique). Based on the obtained electrophoretic mobilities, the zeta potentials ( $\zeta$ ) were calculated using Smoluchowski's equation and the results were averaged. A modified Thompson tau test was applied for outlier identification.

Determination of the L-glutathione incorporation efficiency. GSH incorporation efficiency was determined from the collected wash buffer of the washing steps during HSA/SF/GSH nanocapsule production using a 1260 HPLC system (Agilent, Palo Alto, CA) equipped with a 1290 ELSD detector (Agilent, Palo Alto, CA). Separation was performed using a Hypercarb PGC column (Thermo Fisher, Waltham, MA, 100 mm  $\times$  3 mm). The mobile phase was a mixture of A: 20 mM NFPA and 0.6% trifluoroacetic acid (TFA), B: 20 mM perfluoropentaoic acid

(NFPA) and 0.8% TFA, and C: acetonitrile (ACN) at a flow rate of 0.2 mL min $^{-1}$ , under gradient conditions (ESI,† Table S2). Injection volume was 5  $\mu$ L. The detector nebulizer and evaporator were set to 60  $^{\circ}$ C at 1.6 SLM nitrogen flow. Standard calibration covered the 1–1000  $\mu$ g mL $^{-1}$  range with a GLU solution in ultrapure water. Experiments were performed in triplicate and statistical outlier tests were applied.

Attenuated total reflection Fourier transformation infrared spectroscopy. Structural analyses of lyophilized HSA, HSA/SF and HSA/SF/GSH nanocapsules were performed by ATR FT-IR using a Spectrum GX spectrometer (PerkinElmer, USA) with diamond ATR accessory (Spec Ltd, UK). The spectra of HSA/SF/GSH nanocapsules were recorded in the 600–4000 cm $^{-1}$  range (spectral resolution of 1 cm $^{-1}$ ). Obtained spectra were normalized for conserved bands between 650 and 1200 cm $^{-1}$  and baseline corrected. Secondary structure ratios of  $\beta$ -sheets and  $\alpha$ -helices were calculated for the recorded amide I bands.

**DLS based thermal stability analysis.** Thermal stability tests were performed on unloaded and MEN- or RK-loaded HSA/SF/GSH nanocapsules, where non-diluted samples were analyzed over a 25–70 °C temperature range with the same acquisition settings as for size measurements.

Scanning electron microscopy. SEM images of HSA, HSA/SF and HSA/SF/GSH nanocapsules were obtained by collecting secondary electrons using a Hitachi 3030TM (Metrohm INULA GmbH, Austria) equipped with a detachable cryo stage (Deben UK Limited, Suffolk, UK) with energy dispersive X-ray spectrum (EDX) acceleration voltage and charge up reduction mode.

Radical scavenging analysis. Antioxidative properties of unloaded and MEN- or RK-loaded HSA/SF/GSH nanocapsules were analyzed using a nitric oxide scavenging activity assay according to the procedure published by Alam et al.48 Briefly, 2 mL of 10 mM nitroprusside solution (214 mg sodium nitroprusside dissolved in 72 mL 50 mM sodium phosphate buffer pH 7.4) were mixed with 0.5 mL of nanocapsule dispersion at various concentrations (10, 50, 100, 200, 300, 400, 500, 600, 800 and 1000 µL mL<sup>-1</sup> buffer). The mixtures were then incubated at 20 °C for 150 min. After incubation, 0.5 mL of the incubated mixture was mixed with freshly prepared 0.5 mL Griess reagent (10 mL sulfanilic acid reagent, containing 48.84 mg sulfanilic acid in 10 mL 20% glacial acetic acid incubated at room temperature for 5 min, was mixed with 10 mL naphthylethylenediamine dichloride 0.1% (w/v)). The mixture was then incubated at 21 °C for 30 min. Then, the absorbances of the nanoparticle dispersion mixtures with and without Griess reagent were measured at 546 nm using a Tecan Infinite M200 Pro (Tecan Trading AG, Switzerland). The amount of nitric oxide radical inhibition was calculated using the following equation.

Nitric oxide inhibition (%) = 
$$\frac{(A_{\rm br} - A_{\rm ar})}{A_{\rm br}} \times 100$$
 (4)

 $A_{\rm br}$ : absorbance before reaction (a.u.);  $A_{\rm ar}$ : absorbance after reaction (a.u.).

Determination of the encapsulation efficiency of flavour substances. Encapsulation efficiencies (EE) for two flavor substances, 5-methyl-2-(propan-2-yl)cyclohexan-1-ol (menthol, MEN)

and 4-(4-hydroxyphenyl)butan-2-one (raspberry ketone, RK), were determined from collected wash buffer of the performed washing steps during MEN- or RK-loaded HSA/SF/GSH nanocapsule production. 1.0 mg mL<sup>-1</sup> of RK and 10 mg mL<sup>-1</sup> of MEN were added to the aqueous phase before nanocapsule production. Analysis of the concentration of MEN in the washing buffer was performed using an 890A GC system equipped with a MS detector 5975C series and a DB-17MS (30 mm  $\times$  0.250 mm with 0.25 µm film) column (Agilent Technologies, US) at a flow rate of 1.2 mL min<sup>-1</sup>. The inlet temperature was set to 200 °C, the oven to 100  $^{\circ}\text{C}$  and the AUX temperature to 300  $^{\circ}\text{C}$ . For quantification, a standard dilution series of MEN in ultrapure water at concentrations of 10, 5, 1, 0.5, 0.1, 0.05, 0.01 and 0.001 mg mL<sup>-1</sup> was prepared. RK concentrations were determined using a 1260 HPLC system equipped with a G7117A DAD detector and a Poroshell 120 EC-C18 4.0  $\mu$ m (4.6 mm  $\times$  5 mm, 120 Å) guard column coupled to a Poroshell 120 EC-C18 4.0 µm  $(4.6 \text{ mm} \times 50 \text{ mm}, 120 \text{ Å})$  column (Agilent Technologies, US). The injection volume was 10 µL. The mobile phase was a mixture of A: ultrapure water, B: methanol and C: 10% formic acid in ultrapure water with a flow of 0.850 mL min<sup>-1</sup> under gradient conditions (ESI,† Table S3). DAD signals were monitored at 279.4 nm. A standard dilution series in ultrapure water with concentrations of 10, 5, 1, 0.5, 0.1, 0.05, 0.01, 0.005, 0.001, 0.0005 and 0.0001 mg mL<sup>-1</sup> was used for external calibration. The limits of detection, determination and quantification were determined based on DIN32645. The EE was calculated using eqn (5).

$$EE (\%) = \frac{m_{\text{initial}} - m_{\text{Mencapsulated}}}{m_{\text{initial}}} \times 100$$
 (5)

 $m_{\text{initial}}$ : initial amount of menthol or raspberry ketone before nanocapsule production (10 mg mL<sup>-1</sup> for MEN; 1.0 mg mL<sup>-1</sup> for RK);  $m_{\text{Mencapsulated}}$ : amount of menthol or raspberry ketone in the nanocapsules.

Release studies in artificial saliva. For the release experiments, artificial saliva (A39-B44 and D40-E41) and 100 mM sodium phosphate buffer pH 7.4 (C) were prepared and used as buffers to mimic the natural conditions of human saliva and conditions present in gingivitis and periodontitis (Table 2), e.g. pH and viscosity. Additionally, commercially available normed and certified artificial saliva for medical and dental research containing 10 g L<sup>-1</sup> sodium carboxymethyl cellulose, pH 6.8 (F; Pickering Laboratories, Mountain View, California, US) was used.

MEN and RK release experiments were performed with loaded HSA/SF/GSH nanocapsules in triplicate. Here, 100 μL of nanocapsule dispersions were incubated in 1 mL of artificial saliva/buffer at 37 °C (150 rpm) for 168 h to mimic the oral environment. Sampling was performed after 1, 3, 6, 24, 48, 120, 144 and 168 h. During the sampling procedure, the artificial saliva/buffer was separated from the nanocapsules and 1 mL of fresh artificial saliva/buffer was added to the nanocapsules, which were further incubated. The concentration of MEN was then quantified by GC-MS (see "Determination of the encapsulation efficiency of flavor substances").

For RK concentration determination, HPLC-DAD was used (see "Determination of the encapsulation efficiency of

**Table 2** Compositions and pH values of artificial saliva ( $A^{39}-B^{44} & D^{40}-E^{41}$ ) and 100 mM sodium phosphate buffer pH 7.4 (C) used for the release studies of HSA/SF/GSH nanocapsules loaded with MEN or RK and for viscosity studies. All artificial salivas and buffers were sterilized by filtration before use. Na-CMC is sodium carboxymethyl cellulose and M p-HB is methyl p-hydroxy-benzoate

	$A^{39}$	$B^{44}$	C	$D^{40}$	$E^{41}$
Ingredients [g L <sup>-1</sup> ]					
NaCl	0.4	0.4		0.674	0.87
KCl	0.4	0.4		0.960	0.62
MgCl <sub>2</sub> 6H <sub>2</sub> O				0.0408	0.1275
CaCl <sub>2</sub> 2H <sub>2</sub> O	0.795	0.906		0.1168	0.225
NaH <sub>2</sub> PO <sub>4</sub> 2H <sub>2</sub> O		0.69			
NaH <sub>2</sub> PO <sub>4</sub> H <sub>2</sub> O	0.69		138.00		
Na <sub>2</sub> HPO <sub>4</sub> 2H <sub>2</sub> O			177.99		
Na <sub>2</sub> S 9 H <sub>2</sub> O	0.005	0.005			
NaF					0.0044
KSCN	0.3				
$KH_2PO_4$				0.242	0.3
$K_2HPO_4$					0.8
Urea	1	1			
Na-CMC				8	10
M p-HB				1	1
Sorbitol				24	29.95
pH value					
•	4.8	5.3	7.4	5.4	6.86

flavor substances"). Based on the obtained data, cumulative release over 168 h was calculated for each formulation and artificial saliva/buffer; the data were tested and corrected for outliers afterward.

Viscosity analyses of artificial saliva. Artificial saliva (Table 1, A<sup>39</sup>-B<sup>44</sup> & D<sup>40</sup>-E<sup>41</sup>), including the commercial certified and normed artificial saliva by Pickering Laboratories (F) and the 100 mM sodium phosphate buffer pH 7.4 (C), were analyzed using a CVO rheometer (Bohlin Instruments Ltd, UK) equipped with a cone/plate setup (diameter 40 mm, angle 4°, gap width 0.150 mm). Pre-shear phase was performed with rotations of 10 s<sup>-1</sup> adjusted for 10 s and analyses were performed with a constant shear rate of 250 s<sup>-1</sup>. Samples were tested in duplicates of ten single measurements for each artificial saliva. In addition, samples were investigated at isothermal levels of 25 °C and 37 °C.

#### Conclusions

In this study, production of nanocapsules containing glutathione produced from glucose recovered from rayon fiber was investigated. Sustainable enzymatic hydrolysis of rayon fiber waste was used as a circular economy solution. The complete recovery of the cellulose part in the form of glucose allowed cultivation of Saccharomyces cerevisiae and therefore glutathione production. Growth, glucose consumption and glutathione formation by Saccharomyces cerevisiae were almost identical regardless of whether glucose recovered from rayon fiber or pure glucose was used as carbon source. Interestingly, zinc contained in the recovered glucose solution supported production of glutathione. Glutathione in its reduced state was successfully incorporated in nanocapsules combined with human serum albumin and silk fibroin in an ultrasound-assisted production method.

Nanocapsule production was optimized by experimental design, leading to particle sizes of  $462.72\pm73.36$  nm (hydrodynamic radius) and zeta potential values of  $13.67\pm0.01$  mV. For possible future application as a cosmeceutical in the prevention of gingivitis and periodontitis, flavour substances were encapsulated and the release kinetics were studied in artificial saliva, revealing a pH- and viscosity-dependent release which can be adjusted for the desired flavour substance. Proven radical scavenging properties of HSA/SF/GSH nanocapsules are seen as promising formulation properties for use in cosmeceutical approaches. Therefore, an economic cycle was successfully designed and the optimized HSA/SF/GSH nanocapsule formulations can be used to offer beneficial healing environments in cases of gingivitis and periodontitis, based on the obtained properties.

#### Conflicts of interest

The authors declare no competing financial interest.

#### Acknowledgements

This work has been supported by the Federal Ministry of Science, Research and Economy (BMWFW), the Federal Ministry of Traffic, Innovation and Technology (bmvit), the Styrian Business Promotion Agency SFG, the Standortagentur Tirol and ZIT – Technology Agency of the City of Vienna through the COMET-Funding Program managed by the Austrian Research Promotion Agency FFG. Additionally, the work performed has received financial funding from the European Union H2020 Programme (H2020-NMP-PILOTS-2015) under the grant agreement FOLSMART [grant number 683356] and from the NÖ Forschungs- und Bildungsges.m.b.H. (NFB) and the provincial government of Lower Austria through the Science Calls (Project ID: SC16-024).

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