Synergetic effect of cationic starch (ether/ester) and Pluronics for improving inkjet printing quality of office papers

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Abstract

Improving the printability of paper is still a relevant challenge, despite the fast development of digital communications. While it is well-known that cationic starches enhance ink density, their commercial paper-grade forms are limited to ethers with low degree of substitution. This work addresses the underexplored potential of highly substituted cationic starch for paper coating and its combination with tri-block polymers, namely Pluronics (P123 and F127), taking advantage of their supramolecular interactions with amylose chains. For that purpose, cationic starch ether and ester (starch betainate), both with a degree of substitution of 0.3, were synthesized by alkaline etherification and by transesterification, respectively. Paper without any surface treatment was subjected to one-side bar coating with suspensions encompassing those products and Pluronics, besides other common components. Black, cyan, yellow and magenta inks were printed on all coated papers through an inkjet printer. Key properties of printing quality such as the gamut area, gamut volume, optical density, print-through, inter-color bleed and circularity were measured in a controlled temperature-humidity environment. For instance, a formulation with cationic starch (ether/ester) and P123 improved the gamut area by 16–18% in comparison to native starch-coated paper sheets. Interestingly, the individual assessment of each component showed that cationic starch ether, starch betainate and P123 only improved the gamut area by 5.6%, 8.9% and 6.8%, respectively. Finally, but not less importantly, starch betainate was found to quench optical brightening agents to a lesser extent than cationic starch ethers.

Keywords

Cationic starch, Paper coating, Pluronics, Printing quality, Starch betainate, Whiteness
Introduction

Paper coating formulations of printing and writing papers (P&W) often comprise a number of different components such as pigments, surfactants, binders, thickeners, dispersants, crosslinkers, optical brightening agents (OBA), and/or lubricants. In each case, the composition depends on which objectives papermakers set for the end product. A careful selection of coating components can therefore be used to develop a paper surface with outstanding smoothness, enhanced barrier properties and, receiving less attention in the literature, improved printing properties (Sharma et al., 2020). Adsorption onto cellulosic fibers occurs when the paper surface is exposed to the coating suspension, but manufacturers cannot neglect the interactions between the components of such suspension, which take place beforehand, from the very moment they are mixed in an aqueous media. These interactions may include competitive adsorption, inclusion complex formation, and stabilization/destabilization (Sousa, De Sousa, Reis, & Ramos, 2014).

The inkjet printing properties of fine papers are majorly influenced by the surface properties thereof, such as charge, surface energy, roughness, permeability and surface strength (Bollström et al., 2013). A slight charge on the paper surface may lead to the effective immobilization of the ink pigments onto the coated paper surface, whereas a certain surface energy balance can favor a higher print density (Lundberg, Örtegren, Norberg, & Wågberg, 2010; Stankovská, Gigac, Letko, & Opálená, 2014). Based on that, it has been shown that highly substituted cationic starch (HCS) has a significant positive effect on the ink holdout (Lee et al., 2002), optical density, whiteness, water fastness and ink fathering properties (Gigac, Stankovská, Opálená, & Pažíný, 2016; Lamminmäki, Kettle, & Gane, 2011). These properties, along with the gamut area (GA), further increase in combination with amphiphilic polymers such as poly(vinyl alcohol) (Baptista et al., 2016), most probably due to ease of interpolymer diffusion of ink carriers during printing (Lamminmäki et al., 2011; Sousa et al., 2014).

While the biodegradability of native starch is obviously not under question, the biodegradability of its derivatives is too often taken for granted. It has been shown that HCS ethers lose biodegradability with increasing degree of substitution (DS), becoming non-biodegradable at DS ≥ 0.54 (Bendoraitiene, Lekniute-Kyzike, & Rutkaite, 2018). In this context, starch betainate (SB) rises as a convincing alternative, not only because betaine is naturally found, unlike conventional cationizing reagents, but also because the ester bonds of SB are clearly more labile than ether bonds (Auzély-Velty & Rinaudo, 2003). Likewise, starch betainate (SB), a cationic starch ester, was suggested for the improvement of paper strength since the first work reporting its synthesis (Granö, Yli-Kauhaluoma, Suortti, Käki, & Nurmi, 2000). However, as far as we know, no study has addressed the influence of SB on the printing properties of fine papers. This issue is addressed in the present work, evaluating coating formulations comprising SB and other interesting amphiphilic polymers, namely Pluronics.

Pluronics® is BASF’s trade name for the less commonly called poloxamers. This trade name comprises non-ionic, water-soluble, triblock copolymers of polyethylene oxide (PEO) and polypropylene oxide (PPO) units. Interestingly enough, they generally form inclusion complexes with starch in aqueous solution. This kind of binding has a significant effect on the dispersion performance and can be explained by hydrophobic interactions between hydrophobic parts of Pluronics® macromolecules and the cavities of the amylose helix (Petkova-Olsson, Altun, Ullsten, & Järnström, 2017). Additionally, the micellar structure of these non-ionic surfactants also influences the adsorption onto the surface of
cellulosic materials, which is enhanced in the presence of cationic polymers (Liu, Vesterinen, Genzer, Seppälä, & Rojas, 2011; Liu et al., 2010).

In light of the aforementioned hypotheses and previous findings, Pluronics was combined with cationic starches in order to observe their effect on the inkjet printing quality parameters. Paper sheets were coated using different concentrations of SB, HCS, Pluronics (P127 and F127), precipitated calcium carbonate (PCC), alkyl ketene dimer (AKD) and optical brightening agent (OBA). This study also illustrates the use of a statistical tool to design the coating experiments and to identify the most important factors to be considered for improving the paper printability. A comparison of HCS and SB coatings was also explored, discussing their influence on the whiteness of paper, given that the interaction between cationic polymers and OBAs, generally anionic, has been pointed out as a major cause of fluorescence quenching (Shi et al., 2012).

Materials and Methods

Materials

Native corn starch (NS), α-amylase (in standard buffer solution, pH 5.8), PCC, OBA and AKD were of industrial origin. 3-Chloro-2-hydroxypropyltrimethyl ammonium chloride (CHPTAC), Pluronics® P123 (MW ~5750 g mol⁻¹, PEO~30 %wt. and CMC of 0.313 mM at 20 ºC) (Alexandridis, Holzwart hf, & Hatton, 1994) and Pluronics® F127 (MW ~12600 g mol⁻¹, PEO-70 %wt. and CMC of 0.56 mM at 25 ºC) (Thapa, Cazzador, Grønlien, & Tønnesen, 2020) were purchased from Sigma-Aldrich. Betaine hydrochloride (99%) was purchased from Alfa Aesar and used as-is for transesterification. All solvents were purified or dried prior to use the standard procedures. Other commercially available compounds were used without further purification.

Synthesis of HCS and SB

Native starch was mildly hydrolyzed with α-amylase (0.45 µL g⁻¹ of starch), under continuous stirring, at 80 ºC for 5 min. The temperature was raised up to 90–95 ºC for 15 min. Then, the starch solution was cooled down and absolute ethanol was added to precipitate the polysaccharide, hereinafter referred to as “cooked starch”. Cooked starch was then vacuum filtered, dried and stored in an oven at 50 °C. This pretreatment is common to the synthesis of both HCS and SB.

HCS was synthesized as described elsewhere (Haack, Heinze, Oelmeyer, & Kulicke, 2002). Briefly, 10 g of cooked or native starch was converted into HCS using 33.5 mL of CHPTAC (60% wt.) and 5.9 g of NaOH. The reaction was carried out for 24 h at 70 ºC in 100 ml of distilled water. The reaction mixture was then neutralized with a 0.1% HCl solution.

Starch betainate (SB) was synthesized, as described in a previous paper (Sharma, Aguado, Murtinho, Valente, & Ferreira, 2021), through the transesterification of starch with methyl betainate (MeBetCl) in polar aprotic solvents. 24 g of betaine hydrochloride was first esterified to synthesize MeBetCl using 11.3 mL of thionyl chloride and 75 mL of
methanol, under reflux, for 4 h at 70 °C. MeBetCl was recovered through evaporation of methanol followed by trituration in diethyl ether and, finally, the crude product was dried under high vacuum. Then, 10 g of starch were converted into SB using 20.8 g of MeBetCl in N,N-dimethylformamide (DMF), 100 mL. Prior to transesterification, cooked starch was pre-activated in NaOH/ethanol. The reaction was carried out for 24 h at 70 °C. HCS and SB were precipitated by adding ethanol (alcohol/water > 10, v/v), vacuum filtered and washed with absolute ethanol, followed by drying at 50 °C.

**Characterization of synthesized cationic starches**

Attenuated Total Reflectance-Fourier Transform Infrared (ATR-FTIR) spectroscopy, \(^1\)H-nuclear magnetic resonance (\(^1\)H-NMR) spectroscopy and viscometry analysis were performed to characterize the synthesized cationic starches. ATR-FTIR spectra were recorded by using an Agilent Cary 630 spectrometer, from 750 to 3000 cm\(^{-1}\), at a resolution of 4 cm\(^{-1}\) and 64 scans per sample. NMR spectra were obtained from a Bruker Biospin GmbH spectrometer, at 400 MHz, using D\(_2\)O as solvent. The degree of substitution was calculated from the area of the singlet assigned to the methyl protons of the quaternary ammonium group. The reliability of this result was confirmed by measuring the nitrogen percentage of samples on a Fisons Instruments EA 1108 CHNS-O elemental analyzer.

**Paper coating**

NS was used as a common component for preparing all formulations in this work. For that, NS was cooked as described earlier, and then cooled down to 50 °C instead of precipitated. An industrial calendered uncoated paper (base paper, BP), produced from bleached eucalyptus kraft pulp with a basis weight of ~ 78 gm\(^{-2}\), was used as substrate for performing surface coating.

Coating of BP was performed using a Mathis laboratory coater, with a pre-drying infrared system coupled to the applicator bar (SVA-IR-B). An applicator roll with the diameter of 0.13 mm, in conjunction with a velocity of 6 m min\(^{-1}\) and intermediate load at both sides, was used to achieve 1.5 to 3 g m\(^{-2}\) per side, on the basis of dry coating weight. Coated paper sheets were air dried at room temperature. Besides NS, BP sheets were coated with SB, HCS, P123, F127, PCC, and combinations thereof. Coatings were performed using 8%, 16% and 24% of total solids coating weight of each of these components, and several combinations of them were tested at said concentrations. The coating compositions resulting from individual components (with NS) and combinations thereof are shown in Table 1 and 2, respectively. The surface weight gain was calculated by the difference between basis weights (ISO standard 536:1995) of the air-dried coated paper sheet and the respective BP sheet. Before characterization, all coated papers were kept at controlled temperature (23 °C ± 1) and humidity (RH 50% ± 2). For each run, three numbers for paper sheets were coated and characterized for evaluating the printing quality.
Table 1. Composition of coating components expressed as %w/w, on the basis of dry coating weight.

| Ingredients | HCS/SB | P123 | F127 | PCC | Reference |
|-------------|--------|------|------|-----|-----------|
| HCS/SB      | 8      | 16   | 24   | 16  |           |
| P123        | 8      | 16   | 24   |     |           |
| F127        | 8      | 16   | 24   | 16  |           |
| PCC         | 8      | 16   | 24   |     |           |
| OBA         | 6      | 6    | 6    | 6   |           |
| AKD         | 0.4    | 0.4  | 0.4  | 0.4 |           |
| NS          | 85.6   | 77.6 | 69.6 | 69.6| 93.6      |

Table 2. Composition of coating components for the interaction study, expressed as %w/w, on the basis of dry coating weight.

| Ingredients | HCS/SB +P123 | HCS/SB +P123+PCC | Reference |
|-------------|--------------|------------------|-----------|
| HCS/SB      | 16           | 16               | 16        |
| P123        | 8            | 16               | 16        |
| PCC         | 16           | 16               | 16        |
| OBA         | 6            | 6                | 6         |
| AKD         | 0.4          | 0.4              | 0.4       |
| NS          | 69.6         | 61.5             | 93.6      |

Statistical analysis

In order to observe the interactions between coating components and their impact on the printing quality of coated papers, JMP software was used as a statistical tool for the design of experiments and data analysis. In this study, four continuous factors, namely HCS, P123, PCC and OBA, were selected, each at two levels (0 and 16%), and a full factorial design with two center points was chosen to design the coating experiments. A total number of 18 runs were performed to evaluate the effect of these factors and their ternary interactions on the printing quality, namely: GA; OD for cyan, magenta, yellow, and black; PT; inter-color bleed (ITCB), and circularity for black color in the responses.

Printing quality

The printing quality was evaluated as reported elsewhere (Lourenço, Gamelas, Sarmento, & Ferreira, 2020). Briefly, the coated papers were printed using HP Officejet Pro 6230 inkjet printer, having cyan, magenta, yellow, and black color ink cartridges. The printed sheets were air dried for 4 h under controlled conditions of temperature and humidity. The GA is the area of the hexagon resulting from the a* and b* coordinates of six printed colors (red, green, blue, cyan, magenta, and yellow), where a* axis represents the color from green to red and axis b* represents the color from blue to yellow. It was determined by measuring the values of CIE L*a*b* coordinates for six color spots, including three base colors (cyan, magenta and yellow) and other three complimentary colors (red, green and blue). For that, the “X-Rite Eye One XTreme UV Cut” spectrophotometer was placed on each printed color spot, activating the UV light (D50, 2º).
readings were taken in the sequence of red, green, blue, cyan, magenta, and yellow color spots. Additionally, CIE L*a*b* values for black and white colors were measured to estimate the gamut volume (GV) of printed paper sheets.

In order to evaluate the other printing properties, such as OD and PT, the QEA PIAS-II spectrophotometer was used with a low resolution optical module (33µm/pixel with visual area of 21.3mm × 16mm), along with the software PIAS II, based on ISO 13660 quality standards, for processing the images. The PT of a printed paper requires the measurement of L*a*b* values on the opposite side, in contrast with the non-printed area of the same paper sheet. The transmitted light intensity from a specific area of each color (black, white, cyan, magenta, and yellow) was measured using QEA PIAS-II, and thus PT and OD were calculated from the following equations:

\[ \text{Optical density} = \log_{10}(\text{Insentile light} / \text{Transmitted light}) \]  

Equation 1

\[ \text{Print through} = \sqrt{(L_p^* - L_u^*)^2 + (a_p^* - a_u^*)^2 + (b_p^* - b_u^*)^2} \]  

Equation 2

where \( L^*, a^*, b^* \) are the CIE chromatic coordinates, and the subscripts \( u \) and \( p \) refer to areas of unprinted and back of the printed black spot, respectively.

The other printing properties, namely ITCB and circularity (black), were also evaluated by means of QEA PIAS-II with high resolution module (5 µm/pixel with 3.2mm × 2.4mm). This was used to measure the raggedness, which can be defined as the geometric distortion of the line and dots, given by the standard deviation of the residue from the lines and dots adjusted to their ideal limit. The higher the raggedness, the worse the ITCB and circularity.

**Paper properties**

The surface hydrophilicity for SB-coated papers was evaluated by contact angle goniometry. The static water contact angle (WCA) was measured in an OCA 20 goniometer (Dataphysics, Germany) using the sessile drop method. A droplet of deionized water (10 µL) was automatically poured onto the coated paper surface. After settling, the formed angle was measured by fitting the Young-Laplace equation to the drop profile. Bendtsen roughness (ISO 5636-3, 8791-2) and Gurley air permeability (ISO 5636/5) were also measured for coated papers using appropriate testers from Flank.

Whiteness (CIE W D65/10) of coated papers was measured using D65 illumination in the Elrepho spectrophotometer. The average value and the standard deviation of four independent measures are reported for CA, Bendtsen roughness, Gurley air permeability and whiteness. This last property was related to the performance of OBA, as fluorescence emission spectra of solutions containing OBA were recorded by means of a FluoroMax 4 spectrofluorometer from Horiba.

The kinematic viscosity was determined using a size 100 Cannon-Fenske viscometer in a thermostatic bath (TAMSON TV 2000) set at 40 ºC. Measurements followed the ISO 3105 standard. Polymer solutions were prepared with a concentration of 5 mg cm\(^{-3}\) in 1M NaOH/H\(_2\)O (for HCS) and DMSO (for SB). The thermogravimetric analysis (TGA) was carried out on a thermo-microbalance TG 209 F3 Tarsus, from Netzsch Instruments. Samples were heated from 40 ºC to 600 ºC, under a flow of nitrogen (20 mL min\(^{-1}\)), with a heating rate of 10 ºC min\(^{-1}\). TGA was performed for filter
paper coated with P123, F127 and SB. A filter paper was cut into pieces (5cm × 2 cm) and Pluronics P123/F127 were absorbed in this cellulosic substrate. This was carried out with a 10% Pluronic aqueous solution and a LayerBuilder dip coater from KSV. Cellulose substrates were dipped into the solution for 3 min, pulled out and air-dried. The same procedure was followed for 10% SB and 10% P123/F127 + 10% SB aqueous solutions.

Results and discussion

Synthesis of SB and HCS

Figure 1a presents the ATR-FTIR spectra for synthesized cationic starches from etherification and transesterification, using respectively CHPTAC and betaine hydrochloride, in comparison to the NS spectrum. The absorption peaks at 3300 cm⁻¹, 2912 cm⁻¹, 1648 cm⁻¹ can be assigned to the –OH, –CH₂ stretching vibrations, and H₂O bending vibration due to water sorption, respectively. Additionally, peaks at 994 cm⁻¹ can be attributed to the ether bonds and the absorption band at 897 cm⁻¹ can be assigned to C1–H bending in starch. Compared to cooked starch, a new prominent peak at 1473 cm⁻¹ can be observed due to the quaternary ammonium group attached to the anhydroglucose unit (AGU) (Hebeish, Higazy, El-Shafei, & Sharaf, 2010; Wang & Cheng, 2009). Furthermore, the absorption band at 1750 cm⁻¹ is assigned to the ester bond in SB.

Figure 1b shows the ¹H-NMR spectra for HCS and SB, compared to the spectrum of cooked starch. The singlet at 3.28 ppm is assigned to the nine hydrogens of methyl groups of the quaternary ammonium. The resonances from 3.5 to 4 ppm represent the hydrogens attached to carbons 2, 4, 5, 6 (H-6 and H-6'), and 3 of AGU, typically in that order. The doublet for the H-1(α) anomic proton lies downfield (5.35 ppm). There was a certain shift upfield and broadening of all signals upon cationization. No impurities were detected in SB, but the HCS spectrum displayed a singlet at 3.33 ppm and a quadruplet at 3.65 ppm, none of which belong to the canonical structure of cationic starches. The former could be due to quaternary ammonium groups arising from substitution on hydroxypropyl chains, instead of the hydroxyl groups of AGU.
Figure 1 ATR-FTIR (a) and $^1$H-NMR (b) spectra for cationic starch ether and starch betainate, compared to cooked starch.
An important hypothesis regarding the reaction is that enzymatic cooking improves its efficiency. Figure 2a shows the effect of cooking and reaction time on the DS of the synthesized HCS. It can be observed that the DS increases from 0.20 to 0.33 and from 0.34 to 0.43 with the increase in the reaction time from 3 to 24 h, when native and cooked starches were used as raw materials for the etherification reactions, respectively. This is likely due to the formation of more porous starch granules, which facilitates the access of the reagent to hydroxyl groups (Huber & BeMiller, 2001). It was also observed that the cooking of starch enables the homogeneous dispersion of starch granules in the solvent by increasing the solubility and decreasing its viscosity, as seen in Figure 2b (Gao, Luo, Fu, Luo, & Peng, 2012).

Undoubtedly, due to the cleavage of 1–4 α-D-glucopyranosyl linkages of amylose and amylopectin, the inherent viscosity decreases with the enzymatic pre-treatment, from 199.4 cm³ g⁻¹ (NS) to 151.8 cm³ g⁻¹ (cooked starch). The viscosity was further reduced by 43–45% when using them in etherification. Likewise, the hydrolysis of starch molecules in highly alkaline media and at high temperature is evidenced by a loss of viscosity after functionalization. Nonetheless, after reaction times beyond 3 h, further hydrolysis is either negligible or compensated by the effects of cationization on polymer-solvent interactions.

![Figure 2](image_url)

**Figure 2** Effect of reaction time and enzymatic cooking on degree of substitution (A) and inherent viscosity (B) of HCS.

Like etherification, the increase in DS and decreasing in viscosity were also observed in the synthesis of SB. However, DS increased much more abruptly, from 0.01 to 0.33, when using NS and cooked starch in the transesterification reaction, respectively, proving the poor reaction efficiency with NS. Given that the inherent viscosity decreased by ~73%, starch faced higher depolymerization during functionalization, mostly due to the previous alkalization of starch at high temperature.

**Paper printing properties**

Both native starch and CS require cooking before adding other coating components to prepare a uniform coating solution. It should be noted that the other coating components (PCC, Pluronics, OBA and AKD) were always added after cooling down the starch solution to 50 °C. It should be stressed that the P123 and F127 have critical micelle concentrations of 0.313 mM (20°C) and 0.56 mM (25°C), respectively. These values are lower than the amount of
Pluronics used in the experiments reported here; additionally, the critical micelle temperatures of these surfactants are well below 50 °C which also support that Pluronics are in the micelle form (He & Alexandridis, 2018).

In order to prepare the coating formulations, corn native starch was used as host component and cooked using α-amylase enzyme in aqueous medium at 80 °C, for 5 min. CS was then added and cooked together with native starch at 90-95 °C for 15 min. The starch solution was cooled down for 15 min.

Gamut area

It was observed that the GA, presented in Figure 3a, increased by 8.6%, 9%, and 12.5% using 8%, 16% and 24% dry solids content of SB, respectively, compared to NS coating. Plausibly, the high DS led to higher deposition of SB on the paper surface, resulting into improved GA (Niegelhell et al., 2018).

Besides charge density, addition of SB slightly increased the hydrophobicity of the paper surface and smoothness; both also contributed to improve GA (Gigac, Stankovska, & Pazitny, 2016) (Gigac, Stankovska, et al., 2016). Table 4 can be referred to for the Bendtsen roughness, and contact angle values for SB coated papers.

Similarly to GA, GV, which considers the color luminance L, besides a and b, also increased with increasing SB concentration. The maximum increase was observed as 16.4% using 24% of SB, compared to NS coating (refer Table 4).

Figure 3 Effect of different concentrations of SB (a), Pluronics (b), PCC (c), and their combinations (d) on GA.
Table 3 Properties of coated papers using different concentrations of SB, Pluronics (P123 and F127), and their combination in coating formulations.

| Conc. (%) | Roughness (mL/min) | Gurley (mL/min) | Contact angle (°) | Whiteness | Gamut volume / 10³ | Weight gain (g/m²) |
|-----------|---------------------|-----------------|-------------------|-----------|-------------------|-------------------|
| Reference (NS) | 329 ± 13 | 318 ± 7 | 72 ± 1 | 162.8 ± 1.3 | 132 ± 1 | 1.8 ± 0.3 |
| SB | 8 | 324 ± 14 | 379 ± 14 | 75 ± 2 | 162.3 ± 0.5 | 146 ± 1 | 2.5 ± 0.2 |
| | 16 | 311 ± 7 | 397 ± 4 | 77 ± 2 | 164.5 ± 0.3 | 151 ± 1 | 1.9 ± 0.2 |
| | 24 | 376 ± 9 | 420 ± 16 | 79 ± 1 | 164.1 ± 0.7 | 153 ± 1 | 2.3 ± 0.3 |
| P123 | 8 | 368 ± 9 | 407 ± 6 | 50 ± 3 | 165.1 ± 0.2 | 153.3 ± 0.4 | 2.9 ± 0.2 |
| | 16 | 379 ± 30 | 381 ± 5 | 43 ± 2 | 165.6 ± 0.2 | 152 ± 2 | 2.9 ± 0.1 |
| | 24 | 357 ± 9 | 354 ± 9 | 47 ± 1 | 165.8 ± 0.4 | 150 ± 2 | 3.0 ± 0.2 |
| F127 | 8 | 378 ± 38 | 356 ± 5 | 63 ± 3 | 163.6 ± 0.6 | 145 ± 2 | 2.4 ± 0.2 |
| | 16 | 363 ± 28 | 325 ± 50 | 57 ± 1 | 163.4 ± 0.4 | 141 ± 3 | 2.7 ± 0.1 |
| | 24 | 329 ± 18 | 444 ± 13 | 53 ± 2 | 162.2 ± 0.7 | 150 ± 1 | 2.7 ± 0.2 |
| PCC | 8 | 414 ± 37 | 347 ± 5 | 75 ± 2 | 160.9 ± 0.3 | 149 ± 1 | 2.5 ± 0.3 |
| | 16 | 370 ± 15 | 381 ± 11 | 83 ± 2 | 161.3 ± 0.1 | 146 ± 1 | 2.5 ± 0.2 |
| | 24 | 365 ± 5 | 367 ± 2 | 80 ± 2 | 162.5 ± 1.1 | 143 ± 1 | 2.7 ± 0.1 |
| P123 (with SB) | 8 | 375 ± 22 | 376 ± 21 | 51 ± 3 | 164.9 ± 1.7 | 162 ± 3 | 2.4 ± 0.1 |
| | 16 | 383 ± 36 | 321 ± 9 | 48 ± 2 | 165.4 ± 0.8 | 161 ± 2 | 2.8 |
| | 24 | 368 ± 13 | 374 ± 13 | 40 ± 3 | 163.6 ± 0.5 | 164 ± 2 | 2.7 ± 0.1 |

In Figure 3b, the GA for different concentrations of Pluronics P123 and F127 is presented. GA is improved by 14.6% using 8% of P123 in the coating solution; however, the GA is further reduced by increasing the P123 concentration from 8% up to 24%. For F127, the 8% addition improves the GA by 10.5%, but further increase in the concentration of F127, from 8 up to 16%, reduces the GA as well. However, the use of 24% of F127 showed almost equal GA increase as 24% of P123, 11.8% and 12.8%, respectively. The increase of GA can be explained by the amphiphilic nature of Pluronics, which facilitates the strong adsorption of these components on cellulosic surfaces (Liu et al., 2010). Additionally, Pluronics form inclusive complexes with starches, leading to the formation of self-supporting flocs in the coating formulation, and enhancing the dispersion of other coating components (Petkova-Olsson et al., 2017; Petkova-Olsson, Ullsten, & Järnström, 2016). Interestingly enough, the lowest amount of P123 and F127 (8%) was found to be more favorable to improve GA. This area was also improved by ~7.9% in the presence of PCC at a concentration of 8% or 16% (Figure 3c), which is related to the gain in hydrophobicity of the paper surface. However, roughness increased with the presence of PCC and GA was further decreased by 3.7% with a large content of PCC (24%) in the coating formulation.

The effect of P123 coatings in combination with SB (16%) and SB (16%)/PCC (16%) is displayed in Figure 3d. It is observed that GA increases by 8.5-9% using P123 or a mixture of P123 and PCC. It was further improved significantly by 16-20% and 19-22% with the presence of SB/P123 and SB/P123/PCC, respectively. This increase in GA can be explained by the sorption of Pluronics on the cellulosic surface, which increases in combination with a highly cationic polymer (Liu et al., 2011, 2010). Moreover, formation of amyllose-Pluronics inclusion complexes may also facilitates the immobilization of the ink pigments on the coated paper surface, improving GA.

TGA contributes to understand the adsorption of Pluronics in the presence of a cationic polymer. Figure 4 (a) and 4 (b) represent the TGA and DTG curves of dip-coated paper samples, respectively. It can be seen that the major
decomposition areas can be divided into three zones, 275 to 350 °C, 300 to 350 °C and 325 to 400 °C for SB coatings, filter papers and Pluronics coatings, respectively. However, an increase in the major decomposition area of both Pluronics was observed when filter papers were coated with the presence of SB (+Pluronics), which can be attributed to an increased adsorption of Pluronics in the presence of SB.

Figure 4 TGA (a) and DTG (b) curves for filter paper, paper + SB, paper + P123, paper + F127, paper + P123 + SB and paper + F127 + SB.

Optical density

OD is an important parameter to evaluate the depth of the color tone in the printed papers, which clearly affects the perceived saturation of a color (Hu, Fu, Chu, & Lin, 2017). Figure 5 represents the OD of the black color with increasing concentration of SB (A), Pluronics (B), PCC (C) and SB/P123/PCC (D). Figure 5c shows the effect of PCC concentration on OD. OD for PCC coating correlates with the Gurley permeability, attaining deeper tones as the sheet became more resistant to air flow (Kasmani, Mahdavi, Alizadeh, Nemati, & Samariha, 2013). The highest improvement was observed at 8% of PCC. Likewise, OD followed the same trends as GA, and thus it increased with increasing concentration of SB and P123/F127. Above all, Figure 5d shows that the highest increase in OD was achieved with the combination of SB-P123-PCC in the coating formulation.
Figure 5 Effect of different concentrations of SB (a), Pluronics (b), PCC (c) and their combinations (d) on OD (0%: reference coating using NS).

Inter-color bleed (ITCB), print-through (PT) and circularity for black color

Figure 6 presents the ITCB, PT and circularity (black dots) of SB, PCC, P123, SB/P123 and SB/P123/PCC coated papers. Similarly to GA, ITCB was also improved (i.e., reduced) upon the addition of these components. The highest decrease in ITCB, 15.9%, was observed with SB/P123/PCC coatings. Unlike GA and ITCB, PT of SB/P123 or SB/P123/PCC coated paper showed a higher PT at the concentrations used in this work, due to decrease in viscosity of the coating formulation, letting the formulation go deeper into the cellulose matrix, which increased the see-through of ink from the other (non-coated) side of the paper. The presence of PCC on the cellulosic surface provided a better improvement in the PT compared to SB or P123 coated papers. Circularity of black dots generally correlates with the ITCB, improving with the formulation containing SB and P123, due to better fixation of ink particles onto the surface.
Whiteness and fluorescence quenching

Whiteness, positively correlated with ISO brightness, represents a paper’s ability to equally reflect a balance of all wavelength of light across the visible spectrum (Hu et al., 2017). The addition of OBA on the paper surface is a cost-effective solution in papermaking to increase the whiteness of printing and writing papers (Shi et al., 2012). Therefore, the interaction between OBA and the other coating components is important. From Table 4, it can be noted that the presence of OBA improved the whiteness of the coated paper but the presence of HCS quenched this agent, resulting in lower whiteness (Figure S1, Supplementary Information). It is also worth mentioning that the presence of P123 and PCC did not show any further improvement in the whiteness.

Table 4 Study of interactions among factors. Percentages are related to the total solids content (on the basis of dry weight) of coating formulations.

| HCS (%) | P123 (%) | PCC (%) | OBA (%) | GA | OD (cyan) | OD (Magenta) | OD (yellow) | OD (black) | PT | ITCB | Circularity (Black) | Whiteness |
|---------|-----------|---------|---------|----|-----------|--------------|-------------|------------|----|------|-------------------|-----------|
| 0       | 0         | 0       | 0       | 6512 | 0.76      | 0.87         | 1.31        | 1.23       | 1.6 | 15.5 | 1.95              | 146       |
| 0       | 0         | 0       | 6       | 6301 | 0.74      | 0.84         | 1.31        | 1.17       | 1.97 | 15.8 | 1.81              | 162       |
| 0       | 0         | 16      | 0       | 6596 | 0.76      | 0.88         | 1.33        | 1.20       | 1.56 | 15.4 | 1.98              | 146       |
In comparison to NS coatings, whiteness increased by 11% with the addition of OBA. As aforementioned, HCS (with OBA) reduced the whiteness of coated papers by ~10.85% due to the OBA quenching, irrespective of the presence of any other components. Interestingly, such loss of whiteness was not observed when SB was used instead of the cationic starch ether (Figure S1).

To understand the OBA quenching effect in the presence of HCS and SB, fluorescence emission spectra were recorded for solutions containing OBA (1.84 ppm) and either cationic starch (6.1 ppm), so as to keep the same ratio as in coating formulations (6% OBA / 16% CS). Fluorescence quenching was clear in the presence of all cationic starches but, in the case of HCS, the intensity of the emission of blue light (~440 nm) decreased almost by a factor of 4 (Figure 7). Quenching was possibly due to the formation of a non-fluorescent complex, where the sulfonate groups of OBA donate electrons to the quaternary ammonium groups of HCS. Still, the most plausible explanation is the aggregation-caused quenching, where aggregation is promoted by electrostatic interactions. The reason for this is that solutions at higher concentration, such as 9.2 ppm OBA / 24.4 ppm HCS, showed Rayleigh scattering to such extent that no reliable spectrum could be obtained, even though a concentration of 24.4 ppm lies much below the solubility limit of HCS. In other words, there was a phase transition from solution to dispersion when both solutions, each of them displaying negligible light scattering, were mixed. However, regardless of the quenching mechanism, neither this aggregation nor that extent of quenching was observed when using SB/OBA at the same concentrations, supporting the previously described retention of paper whiteness. Given that SB and HCS had the same DS, it may be concluded that the cationic starch ester possesses a key advantage over its ether counterpart. This advantage should, undoubtedly, be further explored.

| 0   | 0   | 16  | 6   | 7074 | 0.80 | 0.91 | 1.36 | 1.21 | 1.76 | 14.4 | 1.81 | 162  |
| 0   | 16  | 0   | 0   | 6758 | 0.79 | 0.89 | 1.33 | 1.22 | 1.76 | 14.5 | 1.80 | 146  |
| 0   | 16  | 0   | 6   | 6922 | 0.80 | 0.90 | 1.34 | 1.20 | 1.86 | 13.7 | 1.76 | 162  |
| 0   | 16  | 16  | 0   | 7004 | 0.81 | 0.90 | 1.34 | 1.20 | 1.80 | 14.2 | 1.74 | 146  |
| 0   | 16  | 16  | 6   | 6802 | 0.81 | 0.90 | 1.33 | 1.21 | 1.69 | 14.0 | 1.71 | 163  |
| 16  | 8   | 8   | 3   | 7122 | 0.82 | 0.90 | 1.34 | 1.19 | 1.65 | 13.8 | 1.80 | 148  |
| 16  | 8   | 8   | 3   | 7212 | 0.82 | 0.91 | 1.33 | 1.20 | 1.69 | 14.2 | 1.81 | 147  |
| 0   | 16  | 0   | 0   | 6610 | 0.80 | 0.78 | 1.33 | 1.25 | 1.99 | 16.0 | 1.79 | 144  |
| 0   | 16  | 0   | 6   | 6656 | 0.84 | 0.77 | 1.37 | 1.24 | 1.78 | 16.2 | 1.79 | 145  |
| 16  | 0   | 16  | 0   | 6971 | 0.77 | 0.88 | 1.31 | 1.20 | 1.82 | 15.6 | 1.95 | 144  |
| 16  | 0   | 16  | 6   | 6739 | 0.79 | 0.78 | 1.31 | 1.23 | 1.82 | 15.6 | 1.84 | 146  |
| 16  | 16  | 0   | 0   | 7479 | 0.83 | 0.92 | 1.36 | 1.22 | 1.80 | 12.7 | 1.83 | 146  |
| 16  | 16  | 0   | 6   | 7443 | 0.86 | 0.85 | 1.50 | 1.42 | 1.66 | 16.3 | 1.89 | 146  |
| 16  | 16  | 16  | 0   | 7427 | 0.83 | 0.91 | 1.35 | 1.20 | 1.72 | 13.9 | 1.76 | 145  |
| 16  | 16  | 16  | 6   | 7670 | 0.85 | 0.93 | 1.37 | 1.22 | 1.84 | 13.2 | 1.71 | 148  |
Figure 7 Fluorescence emission spectrum of OBA in presence of HCS and SB. An excitation wavelength of 350 nm was used to record all spectrum.

**Statistical analysis**

Figures 8a and 8b show the half-normal plots, as obtained from JMP software, for all the studied factors. As indicated in Figure 8a, three major factors are clearly falling off from the red straight line. In other words, P123, HCS and their interaction (HCS*P123) can be considered as the most significant factors to affect the GA, whereas PCC and OBA, like their interaction, were found to be insignificant. The model was further optimized through prediction plots and ANOVA study, and insignificant factors were removed. Figure 8b indicates the half-normal plot of the model, considering only the significant factors.
A prediction profile for GA and a complete report of the model is provided in the Supplementary Information. From this statistical study, it can be inferred that the combination of HCS and Pluronics in the coating formulation, together with the presence of PCC and OBA, led to improve the printability of coated papers. The statistical study of these components has shown that, for GA, the incorporation of P123, the presence of HCS and their interaction, have significant effect in the selected range. Regarding the most important variables for the other printing properties, P123 has significant effect on OD for cyan/magenta/yellow and on ITCB, whereas HCS impacts OD of black color. The ternary interaction HCS-P123-OBA showed a good impact on PT (not seen with SB-P123-OBA) and the binary interaction P123-PCC affects the circularity of black color. The corresponding half-normal plot, the analysis of variance and the prediction profiles have also been included in the Supplementary Information. All considered, the combination of both HCS and P123, accounting for a total solids content of 16%, has the greatest impact on the overall printing quality.

Conclusions

The effect of an unconventional combination of coating components, highly substituted cationic starch and Pluronics, on the printing quality of office papers was investigated. As a key novelty, cationic starch refers not only to its typical ether form, but also to starch betainate, an ester that has been suggested for bulk addition in sheet forming but not (as far as the authors are concerned) for paper coating. A 24% coating weight of starch betainate increased the gamut area by 12.5%, whilst Pluronics P123 and F127 (8% coating weight) attain improvements of 14.6% and 11.8%, respectively. Both cationic starches, ether and ester, showed the same outcome for improving the paper printing properties in presence and absence of Pluronics. Nonetheless, while the ether caused a certain loss of whiteness, as it quenches the fluorescence emission of the optical agent, such loss was not found when starch betainate was used. The ability of starch betainate of keeping the whiteness gain of an anionic brightening agent is a key finding of this work.
Remarkably, the statistical analysis indicated that besides the aforementioned individual effects of cationic starch and Pluronics, the binary interaction thereof had a significantly positive influence on the gamut area. Furthermore, the optical density (cyan, magenta, yellow and black), print-through, inter-color bleed and circularity were successfully correlated with the independent variables. It was shown, for instance, that the print-through was significantly affected by the presence of conventional cationic starch, OBA and Pluronic P123.

Declarations

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Conflicts of interest/Competing interests

The authors declare that there is no conflict of interest and that they do not have competing interests.

Availability of data and material

All data are displayed in the article and its electronic supplementary information.

Code availability

Not applicable.

Authors' contributions

All authors made substantial contributions to the conception of the work, the acquisition and interpretation of data, and writing. All authors approve the manuscript. All authors agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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