Silylated Thiol-Containing Cellulose Nanofibers as a Bio-Based Flocculation Agent for Ultrafine Mineral Particles of Chalcopyrite and Pyrite

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Abstract
A considerable amount of very fine particles can be found, e.g., stored in tailing ponds, and they can include valuable or hazardous minerals that have the potential to be recovered. Selective flocculation, i.e., the formation of larger aggregates from specific minerals, offers a promising approach to improve the recovery of ultrafine particles. This study focuses on the use of a new bio-based flocculation agent made of silylated cellulose nanofibers containing a thiol-functional moiety (SiCNF). Flocculation was performed in separated systems of ultrafine mineral dispersions of pyrite, chalcopyrite, and quartz in aqueous alkaline medium. The flocculation performance of SiCNF was addressed in terms of the turbidity reduction of mineral dispersions and the floc size, and the results were compared with the performance of a commercial anionic polyacrylamide. SiCNF exhibited a turbidity removal efficiency of approximately 90%–99% at a concentration of 4000–8000 ppm with chalcopyrite and pyrite, whereas the turbidity removal of quartz suspension was significantly lower (a maximum of approximately 30%). The sulfide particles formed flocs with a size of several hundreds of micrometers. The quartz in turn did not form any visible flocs, and the dispersion still had a milky appearance after dosing 12,000 ppm of the flocculant. These results open a promising path for the investigation of SiCNF as a selective flocculation agent for sulfide minerals.

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Graphical Abstract

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Introduction

To enable mineral processing, it is usually necessary for the ore to pass through successive fragmentation stages. Along these stages, fine and ultrafine particles, technically known as slimes, are inevitably produced. The ultrafine particles in dispersions have proven to be a challenge for several reasons. For instance, in froth flotation, slimes present a decline in recovery, which is usually attributed to low collision probability and fine particles having insufficient inertia to cross the water streamlines around the coursing bubble. Problems such as surface oxidation, adsorption of metal ions, and slime coating can also be noticed [1, 2]. As the number and grade of mineral deposits and large ore bodies have been decreasing over the years, the beneficiation of valuable minerals attributed to the fine particles has become increasingly important; moreover, there has been an increasing demand to improve conventional methods in the beneficiation of slimes [3, 4].

Selective flocculation, that is, the aggregation of the desired mineral particles (i.e., make them larger) and then separation (recovery) of the aggregates from the dispersed material, is one potential approach for enhancing the efficiency of fine particle processing [2]. Moreover, it appears to be an attractive method for recovering valuable minerals from slimes using, e.g., subsequent froth flotation [3, 5–9].

The principles of selective flocculation using different polymers as well as the surface chemistries have been reviewed earlier by several authors [1–3, 6, 10–16]. These techniques utilize the differences in the physical–chemical properties of the minerals and are based on the preferential adsorption of a flocculant in the mixture of colloidal solids, which results in selective flocculation and leaves the remaining particles suspended [1, 17].

The challenge in selective flocculation is mainly related to the tailored and feasible interactions between the flocculant and specific minerals. The great variety in the composition of slimes and interaction mechanisms makes it even more difficult to adjust the flocculation process [3, 18]. Moreover, hetero-coagulation, interference of dissolved ions, and entrapment of unwanted mineral particles in the flocs can also deteriorate the selectivity of flocculation. In general, the interactions and adsorption caused by the electrostatic forces and hydrogen bonding are unselective (the H-bonding is more specific than electrical forces), although the electrostatic forces can be manipulated by adjusting the surface potential of the minerals [3, 6, 19–21]. Therefore, specific chemical and coordination bonding seem to offer the most promising approach for the adsorption and flocculation of specific minerals. However, the widely used flocculants, such as polyacrylamides (PAMs), exhibit typically poor selectivity in flocculation despite their many benefits, such
as affordability, stability, and efficiency [22]. In addition, these petroleum-derived synthetic polymers exhibit limited biodegradability [23, 24].

Nowadays, there are strong economical, ecological, and social incentives to develop novel green chemicals from renewable resources to reduce the environmental impact of mineral processing. Natural polymers, including polysaccharides, are biodegradable and are easily available from renewable side streams or forest resources [23]. Especially chitosan, starch, cellulose, alginate, and amylopectin are abundant, affordable, and nontoxic biopolymers [25]. Cellulose, for example, has been widely studied as a green alternative to existing synthetic flocculants for the development of sustainable chemicals. Nevertheless, there are certain drawbacks in cellulosic materials, such as its low solubility in water. Cellulose nanomaterials, i.e., nanocelluloses, can in turn readily be mixed with aqueous colloidal dispersions. Moreover, they have been widely studied as sustainable dispersion and emulsification agents, surfactants, and floculants [26–30]. Cellulose nanofibers have a large surface area as well as elongated and flexible morphology; they can also be functionalized to adjust the surface characteristics [22, 31, 32]. Functionalized nanocelluloses have also been used earlier in the selective flocculation of mineral slimes, but the potential of functionalized nanocelluloses as flocculation agents still remains poorly understood [31].

In the flotation of sulfide minerals, selective sulfhydryl collectors, such as xanthates containing dithiocarbamates and dithiophosphates, are used [2, 33, 34]. Previously, cellulose derivatives based on these active groups, such as cellulose xanthates, have also been harnessed for the selective flocculation of lead [17], zinc [17], and copper minerals [12, 19] from a mixture containing quartz or calcite minerals [19, 35]. However, xanthates have many drawbacks in terms of occupational health and environmental safety. They pose toxicity risks to aquatic organisms even at low concentrations [2, 36–38]. Therefore, more sustainable modifications are required for the development of green alternative flocculation agents and other mineral processing chemicals from renewable sources [3, 12, 17, 19, 31, 39, 40]. In general, thiol moieties offer an appealing approach for selective interaction with the sulfide minerals. They can interact with metal ions present on the surface of the mineral to form metallic complexes [3, 19, 41].

The present study aimed to investigate silylated cellulose nanofibers (SiCNF) modified with thiol-functional moieties as an effective and sustainable flocculation agent for the dispersion of fine particles of sulfide minerals based on chalcopyrite and pyrite. The flocculation performance of SiCNF was tested in the dispersion of separate fine particles of chalcopyrite, pyrite, and quartz. Here, the focus was on the interactions, mechanisms, and flocculation efficiency of single mineral particle systems only; mixed systems will be the focus of our next studies. The effects of pH and SiCNF dosage on flocculation were investigated via batch experiments by measuring the residual turbidity of settled suspension and analyzing the characteristics of flocs under optical microscope. Furthermore, the aim was to compare the modified nanocellulose and its performance with a commercial anionic polyacrylamide (APAM).

Materials and Methods

Raw Materials and Chemicals

The mineral samples of pyrite, chalcopyrite, and quartz were obtained from Peru, Mexico, and Germany, respectively. All reagents were used as received, and deionized water was utilized throughout the experiments. The commercial polymeric flocculant, Superfloc A130 (SFA130), was supplied by Kemira (Finland). It was a high-molecular-weight PAM with medium anionic charge density (33%). A dry sheet of industrial bleached birch (Betula verrucosa) kraft pulp was purchased from UPM (Finland) and used as a cellulose raw material. 3-mercaptopropyltrimethoxysilane (MPTMS, 95%) was obtained from Sigma-Aldrich (Germany).

For pH adjustment, 0.1 M/1 M NaOH and 0.1 M HCl (Oy FF-Chemicals Ab, Finland) were used.

Preparation of Mineral Samples

A jaw crusher (SKET, LKSB 100×63, Germany) with a 1-cm discharge opening, followed by sieving, and a roll mill (own development) with 2-mm discharge opening were used to ground the chalcopyrite and the pyrite. Quartz had a smaller initial particle size and was ground only using a ball mill (Bond index, custom built) for 20 min to obtain a particle-size distribution with d80 < 100 μm. The samples were homogenized, split, and stored. The sulfides were stored protected from contact with air to avoid oxidation.

For inductively coupled plasma–optical emission spectrometry (ICP-OES) and X-ray powder diffraction (XRD) analyses, the samples underwent a further step of comminution using a vibratory disk mill (Siebtechnik, Germany) for 3 min. The resulting samples were stored in a desiccator before analysis.

For the flocculation tests, the minerals were further pulverized using a planetary ball mill PM 200 (Retsch, Germany) with two grinding stations of 125 mL at a speed of 650 min⁻¹ for 2 min. The sulfide samples were ground weekly and stored in small batches of 20 g in the fridge to avoid oxidation. Analysis of the particle-size distribution of the minerals was conducted using a Helos/BR Quixel analyzer (Sympatec, Germany).
Synthesis of Silylated Cellulose Nanofibers

A desired amount of dry sheet of bleached birch kraft pulp was dispersed in deionized water, and the suspension was homogenized using a wet disintegrator set to 30,000 rpm for 10 min. A Masuko supermasscolloider grinder (MKCA6-2J, Japan) was used to disintegrate the pulp fibers to nanofibers. The stones of the grinder were first carefully brought within close contact, and then the pretreated pulp slurry was poured into the grinder at a consistency of 1.5%. The gap of the grinding stones was modified from 0 to –90 µm. The pulp was passed three times through the grinder using a zero-grinding stone gap; then, the stones were adjusted to negative gap values to begin the actual nanofibrillation (−20, −40, −60, and finally −90 µm).

The silane solution (20 w/w%) for silylation of cellulose nanofibers was prepared by diluting MPTMS in ethanol and was then mixed with a magnetic stirrer for 10 min. The pH of the CNF suspension was adjusted to 4 with HCl (0.1 M), and the freshly prepared silane solution (0.5 times the amount of the solid content of CNF) was added dropwise using a micropipette to the pulp solution and stirred using a magnetic stirrer at room temperature for 3 h (Scheme 1). The cellulose suspension was then left in the oven at 85 °C for 4 h. Finally, the sample was washed and centrifuged with a solution of deionized water and ethanol. The degree of silylation of the fabricated silylated cellulose nanofibers (SiCNF) was determined via X-ray photoelectron spectroscopy (XPS) and sulfur analysis.

Characterization

Elemental Composition of the Minerals via ICP-OES

The elemental composition of the minerals was analyzed using ICP-OES 5100 (Agilent Technologies, USA). Melt digestion (Li2B4O7 melt leached with HCl) was employed to analyze Al, K, Mg, Si, and Ti of minerals. To analyze Ca, Cu, Fe, Mn, Pb, Si, and Zn, the samples were dissolved with HCl/HNO3, and the residue was molten with Li2B4O7 and leached with HCl. To analyze sulfur, the samples were mixed with MgO/Na2CO3, ignited 4 h at 815 °C, and leached with water.

Characterization of the Mineral Structure via XRD

XRD analysis was conducted via the powder method using an X’Pert Pro diffractometer (PANalytical, the Netherlands) with a position-sensitive detector and Co-ka radiation (45 kV/40 mA). The diffractograms were collected using the angular range (2θ) of 3°–80°, step size of 0.02°, and time per step of 25 s. The minerals were identified by comparing the diffractograms with the International Center for Diffraction Data database.

Density and Particle-Size Analysis of Minerals

The densities of the mineral powders were determined using a gas pycnometer (Micro Pycnometer MKY-1, Quantachrome Instruments, USA); helium was utilized as the displacement gas medium. The particle-size distribution of each mineral was analyzed via laser diffractometer (Beckman Coulter LS 13 320, USA) from a 0.4% (w/v) water slurry that had been ultrasonicated (37 kHz, pulse mode, 100% power) for 4 min after grinding 10 g of the mineral samples (quartz, chalcopyrite, and pyrite) in a ball mill at 300 min⁻¹ for 3 min.

Zeta Potential

Mineral Samples The powder samples were poured into the graduated cylinder containing deionized water (500 mL) and KNO3 10 mM (as a background electrolyte), after which the suspensions were homogenized by agitating the capped...
cylinder for 10 min. The suspensions settled until solid concentrations of 0.01% (w/w) composed of particles below 10 μm were reached (calculated using Stokes’ Law). Three samples (200 mL) of the supernatants were taken, and the pH was adjusted from 2 to 12 using 0.1 M HCl and 0.1 M NaOH. Then, the ζ-potential was measured as a function of pH using the Malvern ZetaSizer Pro (Malvern Panalytical, UK). The data were collected at room temperature in auto mode, and three measurements were performed for each sample.

The mixed solutions of minerals and SiCNF were prepared according to the methodology explained above. After collecting 200 mL of the supernatants, SiCNF was added (8000 ppm) and mixed for 5 min. Then, the pH was adjusted to 8, 9, and 10. Finally, the ζ-potential was measured.

**SiCNF Suspension** The zeta potential of the aqueous suspension of SiCNF was measured at a pH range of 2–12 by adjusting the pH using 0.1 M HCl and NaOH. The SiCNF suspension (0.8% w/w) was diluted to 0.1% w/w, with a background electrolyte concentration of 10 mM (KNO$_3$) before the analysis.

**Transmission Electron Microscopy of SiCNF**

To visualize the structure of the SiCNF, transmission electron microscopy (TEM) analysis was conducted using the JEM-2200FS (JEOL Ltd., Tokyo, Japan). The SiCNF suspension was placed on a sample holder using a carbon-coated copper grid. A small droplet of 0.1% poly-l-lysine suspension was added to the top of the grid, and the excess was removed by touching with the corner of a filter paper. The SiCNF suspension was first diluted with deionized water, and then a small droplet was poured onto the top of the grid. Finally, a 2% uranyl acetate solution was dropped to negatively stain the sample; the excess was removed by touching with the corner of a filter paper. The samples were dried overnight at room temperature and analyzed using an acceleration voltage of 200 kV under standard conditions. Images were captured using the EM-Menu software, whereas the ImageJ software was used to measure the length and width of the individual cellulose nanofibers.

**XPS Analysis of SiCNF**

The X-ray photoelectron spectra of SiCNF were measured using an X-ray photoelectron spectrometer Thermo Fisher Scientific ESCALAB 250Xi (Thermo Fisher Scientific, UK) with a monochromatic Al-Kα (1486.6 eV) source.

**Elemental Analysis of SiCNF**

The elemental composition of the fabricated cellulose nanofibers was analyzed using a LECO CS200 analyzer (LECO Corporation, USA). Since each silylated group contains only one atom of S, the sulfur content directly corresponded to the degree of substitution (DS), which was calculated to be 0.243 mmol/g.

**Fourier Transform Infrared Spectroscopy (FTIR) of SiCNF**

The FTIR spectra of the silylated cellulose nanofibers were recorded via Fourier transform infrared spectroscopy (Bruker Vertex v80, Indepro, Germany). The spectrum of the thin film was collected at room temperature in the transmittance mode at a resolution of 5 cm$^{-1}$ in the wavenumber range of 500–4000 cm$^{-1}$.

**Degree of Polymerization of SiCNF**

The average degree of polymerization (DP) of SiCNF was evaluated from the limiting viscosity, which was measured in cupriethylenediamine (CED) solution according to the ISO 5351 standard. The values of the limiting viscosity were converted to DP using Eq. 1, where $[\eta]$ denotes the limiting viscosity; C, the mass fraction of cellulose; and H, the mass fraction of hemicelluloses. This calculation makes a correction for the contribution of hemicelluloses to the limiting viscosity number and DP of cellulose [42].

$$DP = \frac{(1.65[\eta] - 116H)^{1.111}}{C}$$

According to Eq. 1, the degree of polymerization of the silylated cellulose nanoparticles was measured to be 2569.7.

**Flocculation Experiments**

Flocculation tests were conducted using 2 g of single minerals and 200 mL of deionized water (1% (w/v) suspension) in a 400-mL beaker. To disperse the solids, the suspension was ultrasonicated (37 kHz, pulse mode, 100% power, 4 min) (Elmasonic P series, Germany). The experiments were conducted at varying pH, from 8 to 10 (± 0.3), using diluted NaOH and HCl solutions. After the pH adjustment, the beaker was placed on a stirring plate (MIXdrive 6, 2mag AG, Germany) at a speed of 800 min$^{-1}$, and the flocculant was added (200–12,000 ppm SiCNF per mass of dry mineral, representing 20–120 mg/L concentrations, and 12.5–100 ppm with SFA130 reference flocculant per mass of dry mineral, representing 0.125–1 mg/L concentrations) after 30 s of stirring, followed by a decrease in the stirring
speed to 400 min\(^{-1}\) for 2 min. Reference experiments without the addition of flocculants were also conducted similarly.

Then, the suspension was allowed to stand for 1 min, after which a sample of the supernatant was removed using a 20-mL syringe from four different points, always on the same height of the beaker. The turbidity removal was calculated based on Eq. 2, by measuring the initial turbidity of the suspension and the turbidity of the supernatant after 1 min of settling time.

\[
\text{Turbidity removal}(\%) = \frac{T_0 - T_f}{T_0} \times 100, \tag{2}
\]

where \(T_0\) denotes the turbidity of a reference suspension before the addition of flocculant, and \(T_f\) denotes the turbidity of the supernatant of the treated sample after 1 min. At least, two repetitions of the turbidity measurements of each sample, using a Hach Ratio/XR Turbidimeter 43,900 (Hach, USA), were performed. The standard deviation of the turbidity measurements was always below 1%. The supernatants of chalcopyrite and quartz were diluted 20 and 5 times, respectively, before the turbidity measurement. Commercial APAM SFA130 was used as a reference flocculation agent.

In addition, a drop was taken from the bottom of the beaker to observe the appearance of the generated flocs using an MZ FLIII stereomicroscope (Leica Microsystems, Switzerland) equipped with a DFC320 digital camera (Leica Microsystems, Switzerland). The sample was prepared by dropping the suspensions onto a glass plate.

**Results and Discussion**

**Characteristics of Minerals**

**Chemical Composition**

The elemental composition of the minerals was revealed via ICP-OES analysis, and these data were used in combination with the XRD results (Fig. 1 and Table 1) to determine the mineral composition of the samples. The chalcopyrite consisted mainly of CuFeS\(_2\) (purity \(\sim\) 76 wt% CuFeS\(_2\)) with major contaminants of Fe\(_2\)S (10 wt%) and SiO\(_2\) (4 wt%), pyrite consisted mainly of Fe\(_2\)S (purity \(\sim\) 93 wt% Fe\(_2\)S), and quartz consisted mainly of SiO\(_2\) (purity \(\sim\) 98 wt% SiO\(_2\)) (Table 1).

**XRD Analysis**

According to the XRD analysis, the primary mineral phase patterns present in the quartz and pyrite samples were attributed to quartz and pyrite, respectively, whereas in the chalcopyrite sample, the main pattern of chalcopyrite appeared with small quantities of quartz and pyrite residues (Fig. 1).

**Particle-Size Distribution**

Table 2 and Fig. 2 present the particle-size distributions of the minerals. The volumetric median diameters of chalcopyrite, pyrite, and quartz were 9.3 \(\mu\)m (\(\pm\) 0.3 \(\mu\)m), 4.3 \(\mu\)m (\(\pm\) 0.4 \(\mu\)m), and 8.1 \(\mu\)m (\(\pm\) 0.4 \(\mu\)m), respectively. All minerals

![Fig. 1 XRD patterns of the quartz, pyrite, and chalcopyrite samples](image-url)

![Table 1 Chemical composition of the minerals](table-url)
presented similar particle-size distribution, but pyrite contained a slightly larger proportion of small particles, whereas chalcopyrite had still a larger particle fraction left.

**Zeta Potential**

The zeta potential values of chalcopyrite, pyrite, and quartz were measured in aqueous solutions as a function of pH, and the results are presented in Fig. 3. All the minerals exhibited a negative charge density above the pH of approximately 4 (except pyrite, which demonstrated only negative values), which decreased almost linearly with all minerals as a function of pH, from −30 to −40 mV at a pH of 12. The isoelectric point (IEP) of chalcopyrite and quartz occurred around pH 3.0–3.5 and around pH 3.5–4.0 respectively. However, the IEP of pyrite was not observed within the studied pH range (2–12), as the zeta potential demonstrated only negative values.

Variable values for the IEP of chalcopyrite, pyrite, and quartz have been previously reported. Dhar et al., Rath et al., and Reyes-Bozo et al. [43–45] determined the IEP of chalcopyrite to be around pH 3.0–4.2, which agrees with the results of the present work. However, Mitchell et al. [46] found the IEP at pH 5.5 for a sample containing 65% chalcopyrite. According to Dhar et al. and Somasundaram [45, 47], the IEP of quartz was observed at pH 2.1, which is a lower value than that observed in the current study. However, Liang et al. [48] have reported that the zeta potential of quartz remained negative and decreased from −23 to −49 mV as pH increased from 4 to 10, demonstrating a similar trend to that of the present work. Moreover, Yukselen-Aksoy and Kaya [49] found that the zeta potential changed from −30.2 mV at pH 3 to −65.4 mV at pH 11, whereas Wu et al. and Xu et al.

### Table 2 Cumulative particle size of chalcopyrite, pyrite, and quartz obtained using a laser diffractometer

| Mineral   | $d_{10}$ (μm) | $d_{20}$ (μm) | $d_{50}$ (μm) | $d_{75}$ (μm) | $d_{90}$ (μm) | $d_{100}$ (μm) |
|-----------|---------------|---------------|---------------|---------------|---------------|----------------|
| Chalcopyrite | 0.5           | 0.9           | 3.4           | 9.8           | 23.8          | 103.0          |
| Pyrite    | 0.2           | 0.4           | 2.0           | 6.1           | 11.6          | 37.7           |
| Quartz    | 0.9           | 1.4           | 4.6           | 11.8          | 20.8          | 47.0           |

![Fig. 2 Particle-size distributions of chalcopyrite, pyrite, and quartz](image)

![Fig. 3 The zeta potential (mV) of chalcopyrite, pyrite, and quartz as a function of pH at a constant ionic strength of 10 mM KNO₃. Lines do not necessarily represent linear trends](image)
[50, 51] corroborated that the zeta potential remained negative from pH 3 to 10 with the IEP at a pH < 2.5. Mitchell et al., Patra and Natarajan, and Reyes-Bozo et al. [46, 52, 53] reported that the IEP of pyrite occurs between pH 2 and 2.5. In addition, Liu and Huang [54] have reported that pyrite has an IEP at pH lower than 3.0. However, Reyes-Bozo et al. [44] observed that the IEP of pyrite occurs at pH 8.1. Here, the results suggest that the IEP of pyrite occurred at a very low pH (< 2).

Overall, different factors affect the electrokinetic behavior of sulfide minerals, such as the purity of the mineral, surface oxidation, sample pretreatment, source of the mineral, and background electrolytes [44, 54]. Therefore, the zeta potential values of the same minerals can significantly vary.

Properties of Silylated Cellulose Nanofibers

Morphology

Functionalized cellulose nanofibers with thiol moieties were synthesized using an aqueous silylation reaction based on MPTMS. The silylated nanofibers, which were individualized using ultrafine grinding, exhibited a typical elongated and flexible nanofibrillar morphology, as revealed by the TEM images (Fig. 4) [32]. The width of the smallest, individual SiCNFs varied from 2 to 5 nm (as analyzed of > 300 individual nanofibers using the ImageJ software), whereas the length ranged from 100 nm to several micrometers according to visual observations. However, the length of the individual particles was difficult to determine, as the nanofibers formed an entangled network structure. SiCNF consisted of separate cellulose nanofibers but also contained a notable portion of larger bundles of nanofibers with a diameter of several tens of nanometers.

Elemental Analysis

The amount of silylated groups in SiCNF was determined using an elemental analyzer. Since each silylated group contains only one atom of S (Scheme 1), the sulfur content corresponded directly to the amount of silylated groups, which was calculated to be 0.243 mmol/g. This result indicated that the CNF was successfully functionalized with MPTMS. The sulfur and carbon contents were also investigated via XPS analyses, indicating that SiCNF has a C content of 55.65 wt. % (46.33 mmol/g) and S content of 0.53 wt. % (0.165 mmol/g). The S content observed via the XPS analysis was lower compared with the value obtained via the elemental analysis, likely because the XPS analyzer gives chemical information about the 10-nm surface layer of the sample, whereas the whole sample is analyzed via the elemental analysis.

Chemical Structure

FTIR was further employed to analyze the chemical structure of SiCNF and to prove the successful reaction between the CNF and MPTMS (Fig. 5). Silylation was confirmed by the existence of several characteristic peaks associated with the silylated product. Vibration peaks at 837–900 cm⁻¹ are assigned to glycosidic bonds (related to C₁–H deformation vibrations), peaks at 1042–1135 cm⁻¹ are related to Si–O–Si and Si–O–cellulose bonds, whereas peak at 1181 cm⁻¹ corresponds to Si–CH bond. The band at 1462 cm⁻¹ is related to the bending vibrations of H–C–H and O–C–H, and the
peak at 1642 cm\(^{-1}\) is the bending vibration of the OH groups of the adsorbed water. The typical peak of the thiol group (S–H) was observed at 2550 cm\(^{-1}\), whereas the band at 2914 cm\(^{-1}\) is assigned to C–H (bond with SH). The peaks at 3129–3512 cm\(^{-1}\) represented O–H stretching of the cellulose [42, 55–61].

**Zeta Potential**

The zeta potential of SiCNF was measured as a function of pH (2–12) in the aqueous solution with a background electrolyte concentration of 10 mM (KNO\(_3\)); the results are presented in Fig. 6. The SiCNF exhibited a negative zeta potential, which decreased almost linearly as a function of pH, being between −3 and −27 mV. The IEP of the modified cellulose was not observed within the studied pH range.

**Flocculation of Mineral Suspensions Using SiCNF**

**Turbidity Removal**

The flocculation performance of SiCNF in the dispersion of the fine particles of chalcopyrite, pyrite, and quartz (1% w/v) was analyzed in terms of turbidity removal via batch experiments (Figs. 7, 8, 9). A commercial APAM, Superfloc A130 (SFA130), was used as a reference flocculation agent. The efficiency of the turbidity removal was observed to be always better for chalcopyrite and pyrite suspensions compared with quartz suspensions both with SiCNF and the reference flocculation agent, indicating a more efficient and effective flocculation of sulfide minerals.

High efficiency of turbidity removal (maximum of approximately 90%–99%) was noted with chalcopyrite at all studied pH values (8, 9, and 10) both with SiCNF and the commercial flocculation agent (Fig. 9a and b). A similar trend was also observed with pyrite (the maximum turbidity removal for SiCNF and SFA130 was approximately 90% and 99%, respectively), but the flocculation performance of SiCNF was more affected by pH (Fig. 10a and b). At pH 9 and 10, the maximum flocculation efficiency dropped to approximately 75% and 65%, respectively, with SiCNF.

The turbidity removal of quartz suspension was notably lower than that of sulfide minerals with both flocculants (Fig. 11a and b). The flocculation performance was also strongly attributed to the pH of the suspension, and the maximum turbidity removal of approximately 30% was reached at pH 8. The clear difference between the flocculation behavior of sulfide minerals and quartz was also observed in the visual appearance of mineral suspensions (Figs. 9c, 10c, 11c). The chalcopyrite and pyrite dispersions exhibited a gradual increase in transparency as a function of SiCNF concentration, whereas the quartz dispersion maintained its milky and turbid appearance despite the increase in SiCNF dosage.

The most significant difference between SiCNF and the commercial flocculation agent was shown in the required optimal flocculant dosage. The maximum turbidity removal of SFA130 was obtained at a concentration of 50 ppm with all minerals. The required concentrations for SiCNF were in turn much higher, i.e., 4000–8000 ppm. The reason behind the higher concentrations of SiCNF is partly related to the fact that SiCNF was based on fibrous nanoparticles with larger spatial dimensions and lower specific surface area than that of soluble polymeric flocculant. The hydrodynamic diameter of polymer molecules with a molecular weight of 2 ∙10\(^6\) was reported to be approximately 85 nm [22], whereas SiCNF had a length ranging from 100 nm to several micrometers. Previously, efficient agglomeration of hematite was reported with dosages of 200–500 ppm using anionic dicarboxylic acid (DCC) and sulfonated (ADAC) cellulose nanofibers [31].

The low DS of SiCNF may have also influenced the required flocculant dosages [62]. Cationic cellulose nanocrystals (CNC-EPTMAC) synthesized through a reaction of the 3-chloro-2-hydroxypropyltrimethylammonium chloride (CHPTAC) in basic medium was found to have a better performance in the flocculation of the SiO\(_2\) suspension when the DS, i.e., the cationic charge density of CNCs, increased. Grenda et al. [63] in turn observed that the influence of flocculant dosage was more pronounced for cellulose-based flocculants with a lower DS. In addition, the high charge density and DS of cationic cellulose-based polyelectrolytes led to a superior flocculation performance in terms of floc growth, especially at lower flocculant dosages.

Different from the present study, previously anionic cellulosic flocculants, with diverse lignin contents,
resulted in a lower turbidity reduction in effluent treatment when compared with the performance of the synthetic APAM with the same pH levels [63]. This weaker flocculation performance of cellulose-based flocculant was attributed to a slightly lower charge density and molecular weight than that of APAM.

Properties of Mineral Flocs Formed by SiCNF

Figure 10 presents the optical microscopy images of flocs of pyrite and chalcopyrite formed using 2000 ppm of SiCNF at pH 8. In both cases, mineral particles formed typical irregular fluffy flocs with the largest mean
particle size of several hundreds of micrometers; however, some smaller, more round-shaped aggregates were also observed. The flocs resembled the flocs previously reported for ultrafine hematite particles and anionic cellulose nanofibers [31] and those observed in the flocculation of pyrite fine particles using corn starch [24]. The flocs of pyrite were found to be denser and were likely able to resist better shear forces without breakage compared with those of chalcopyrite. The quartz in turn did not form any visible flocs (Fig. 11c), and the dispersion still exhibited a turbid appearance after dosing the flocculant, which supported the finding that SiCNF had a better efficiency toward sulfide minerals.

Considering the evolution of the floc size with time, it was observed that in the first seconds of the flocculation process predominated and resulted in large aggregates. After achieving the maximum floc size, the large flocs started to break; subsequently, the equilibrium between breakage and re-aggregation was achieved, and the floc size stabilized. A similar behavior was observed in the previous works in the flocculation of colored effluent using PAM and anionic and cationic cellulose-based polyelectrolytes [63] as well as
in the flocculation of pyrite fine particles using corn starch [24].

**Zeta Potential of Mixed Dispersions of Single Minerals and SiCNF**

The zeta potential of mixed dispersions of single mineral particles and SiCNF was measured as a function of pH (2–12) using a flocculant dosage of 8000 ppm (Fig. 11). The measurements were performed under a constant ionic strength (10 mM KNO₃). Both the minerals and SiCNF initially exhibited a negative charge density at pH 8, 9, and 10; the same pH range used in the flocculation tests.

All the mixed dispersions demonstrated a negative zeta potential value, which decreased as a function of pH, similar to the original zeta potential values of both separate mineral and SiCNF samples (Figs. 3 and 6). The zeta potential values of a mixture of quartz and SiCNF were close to that of the original quartz suspension. This behavior indicated that the suspension charge properties were dominated by the mineral particles (concentration of quartz > > concentration of SiCNF) instead of SiCNF, which likely did not adsorb on mineral surfaces and stayed in the suspensions as separate
entities. Moreover, the flocculation results supported this finding. For the mixtures of chalcopyrite and SiCNF as well as pyrite and SiCNF, the level of zeta potential was in turn different from that of pure minerals at pH 8, 9, 10 (except at pH 9 with chalcopyrite and SiCNF). This alteration in the zeta potential was presumably caused by the interaction between SiCNF and the sulfide minerals, which resulted in the adsorption of SiCNF on the surface of chalcopyrite and pyrite particles. Previous work by Sresty and Somasundaran

Fig. 10 Microscopic image of a chalcopyrite, b pyrite, and c quartz after flocculation using SiCNF (2000 ppm of SiCNF at pH 8)

Fig. 11 The zeta potential (mV) of a chalcopyrite and 8000 ppm of SiCNF, b pyrite and 8000 ppm of SiCNF, and c quartz and 8000 ppm of SiCNF as a function of pH at a constant ionic strength of 10 mM KNO₃. Lines do not necessarily represent linear trends
[19] investigated the zeta potential of fine hematite particles as a function of concentration of polystyrene sulfonate flocculant at pH 7.8, in which both the mineral and polymer were negatively charged. It was pointed out that the adsorption of the polymer molecules on the mineral surface led to a significant shift in the shear plane of mineral particles and that the change in the zeta potential was considered to be a good measure of the adsorbed polymer layer rather than that of the original particles themselves [19].

**Interaction and Flocculation Mechanism of SiCNF**

In the studied pH 8, 9, 10, both the flocculants and minerals demonstrated a negative surface charge (Figs. 3 and 6). Similar surface charges can result in electrostatic charge repulsion and prevent efficient interaction between the flocculant and minerals as well as further formation of flocs. However, this is often not the case in practice [22]; the adsorption of anionic polymers on negative surfaces is a common phenomenon and is promoted by specific chemical interactions [3, 23], contrary to physical adsorption, which is driven by the opposite surface charges of a reagent and surface [64].

Previously, mine tailings consisting of negatively charged clay particles were flocculated with high-molecular-weight APAMs in alkaline medium [22]. The electrostatic repulsion between polymer chains and the negatively charged surfaces of the clays caused the polymer to adsorb primarily on the neutral edges of the clay particles, thus, increasing the possibility that the same PAM chain can extend into the solution and adsorb on the surface of other particles and form large and porous aggregates. An anionic cellulose-based flocculant was in turn found to interact with negatively charged bentonite–dye complex through the sulfonate groups (electrostatic) or hydroxyl groups (hydrogen bonding) of the cellulose structure [63].

Here, the interactions between SiCNF and sulfide minerals were likely attributed to the thiol (–SH), and the hydroxyl functional moieties of the cellulose polymer surface, bonding with the hydroxyl species and metal compounds that formed on the mineral surface. A similar mechanism of interaction was observed in the flocculation of sulfide minerals with insoluble cross-linked starch xanthate [40], in which a covalent bond formation between the xanthate group and metal components on the mineral surface of chalcopyrite was observed.

The specific interaction between SiCNF and pyrite or chalcopyrite caused flocculation of sulfide minerals, which was presumably mediated by a bridging mechanism. This mechanism was supported by the fast flocculation rate and relatively open aggregates (Fig. 10). However, it is likely that the electrostatic repulsion between the mineral particles and polymers could allow only a limited polymer adsorption. However, the polymer molecule expansion arising from the charge repulsion induces the production of loops and tails, which lead to the formation of large open-structure flocs. This effect has been found to be effective in the flocculation of negatively charged clay dispersions as cited in the study by Nasser and James [65]. The bridging mechanism is also favored by long-chain length and elongated structure of the flocculant [66, 67]. Here, both SiCNF and SFA130 consisted of long and flexible ribbons.

Both SiCNF and SFA130 demonstrated poor flocculation performance with quartz. Apparently, SiCNF did not show a strong specific affinity for quartz surfaces in an aqueous solution; moreover, there was no (or very low) adsorption of SiCNF on the quartz particles as supported by the zeta potential measurement (Fig. 11). Similar results were obtained in previous works on the flocculation of fine hematite particle and quartz suspensions with anionic cellulose nanofibers [31] and in a selective flocculation of complex sulfides using cellulose xanthate [17]. This effect was explained by the high repulsive forces between the anionic flocculants with quartz. Moreover, the strong electrostatic repulsion between the negatively charged quartz particles and the APAM appeared to prevent the adsorption of APAM on the quartz surface as cited in the study by Somasundaran [9]. Therefore, the effect of the surface charge of the minerals is of high importance in flocculation, as it can potentially prevent efficient flocculation by providing a strong electrostatic repulsion [3, 6, 19]. Calcite–rutile, calcite–quartz, and alumina–quartz mixtures were readily separable on this basis via selective flocculation [6].

In the current study based on single mineral systems, SiCNF strongly reduced the turbidity of the pyrite and chalcopyrite suspensions by promoting the aggregation of fine minerals to larger flocs while exerting a minor effect on the flocculation of quartz suspensions under the studied conditions. Consequently, these results indicate that SiCNF could promote efficient flocculation and separation of chalcopyrite and pyrite from quartz minerals; however, more research on the mixed systems is required.

**Conclusions**

The performance of SiCNF was evaluated on flocculation of ultrafine mineral particles of chalcopyrite, pyrite, and quartz, separately in single mineral systems. The flocculation promoted by SiCNF was then evaluated via the turbidity removal in the dispersion of separate fine particles.

The efficiency of turbidity removal was observed to be always better for the chalcopyrite and pyrite suspensions compared with the quartz suspension both with SiCNF and the reference flocculation agent, indicating a more efficient flocculation of sulfide minerals. Higher concentrations of SiCNF were required to achieve results similar to those
obtained using a commercial polyacrylamide in terms of reduced turbidity and floc size. However, SiCNF represents a valuable environmentally friendly alternative, and the results indicate the use of SiCNF as a flocculant for sulfide minerals separation is promising.

The specific interaction between SiCNF and pyrite or chalcopyrite led to the flocculation of sulfide minerals, which was presumably mediated by the bridging mechanism. This mechanism was supported by the fast flocculation rate and the relatively open aggregates.

The selectivity of SiCNF on flocculation of ultrafine mineral particles of chalcopyrite, pyrite, and quartz, in mixed mineral systems must be further analyzed and corroborated.

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Declarations

Conflicts of interest On behalf of all authors, the corresponding author states that there is no conflict of interest.

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