Modification of cotton fabrics with 2-diethylaminoethyl chloride for salt-free dyeing with anionic dyes

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Abstract A chemical modification of cotton fabrics by 2-diethylaminoethyl chloride (DEAE-Cl) was achieved, and the resulted cotton fabrics demonstrated salt-free dyeing properties with anionic dyes. Nucleophilic property of hydroxyl groups in cotton cellulose was enhanced under alkaline conditions and could react with DEAE-Cl, a chemical possessing both nucleophilic and electrophilic sites. The monolayered DEAE-grafted cotton cellulose could further react with DEAE-Cl to form multiple cationic quaternary ammonium salts (denoted as DEAE@Cotton), which are highly interactive with anionic dye molecules. The strong electrostatic interactions between the DEAE@Cotton and the dyes eliminated the use of inorganic salts in cotton dyeing process. The chemical structure and property of DEAE@Cotton were characterized and compared with untreated cotton. The DEAE@Cotton can be dyed in a salt-free system, and the dye exhaustion was faster than the conventional dyeing method due to the robust electrostatic interactions of the fabrics with anionic dyes. The dyed fabrics demonstrated outstanding color fastness under repeated washing, light exposure, and crocking. The dye adsorption process on DEAE@Cotton follows Langmuir isotherm model ($R^2 = 0.9667$). The mechanism of enhanced dyeability was experimentally proved by treating the fabric with other anionic dyes in a salt-free system, proving the process to be environmentally friendly and cost-effective.
Keywords  Cationization · Cellulose · Environmentally friendly dyeing · Direct dye · Salt-free

Introduction

Cotton is the most popular fiber used in apparels because of its hydrophilicity, comfort, and softness. Coloration of cotton textiles is an indispensable need. However, the colorations of the cotton products, especially reactive and direct dyeing processes, are major sources of water pollutions. Residual dyes, inorganic salts, and additives account for the major pollutants in the discharged wastewater. The use of salts and chemicals for dyeing cotton is necessary to achieve desired color depth and color fastness of the fabrics (Dixit et al. 2015; Kant 2012). Specifically, a large quantity of inorganic salts is commonly used in dyeing cotton fabrics with direct and reactive dyes. Even by increasing salts in the dyeing baths, residual dyes still unavoidably exist in the dyeing wastewater. The used inorganic salts in the dye effluents are not easy to be recycled, resulting in concerns to environmental health and safety (Khatri et al. 2015). Thus, many approaches have been made to reduce residual dyes or eliminate the use of salts in the dyeing processes, including: (1) synthesis of dyes with improved affinity or reactivity toward cotton cellulose (Wang et al. 2016; Xiao et al. 2017; Zhang and Zhang 2015; Zheng et al. 2012); (2) modification of cotton cellulose by introducing stronger dye interactive groups (Arivithamani and Giri Dev, 2017; Fang et al. 2017; Fang et al. 2019; Hauser and Tabba 2001; Ma et al. 2016; Ma et al. 2017; Sadeghi-Kiakhani and Safapour 2018; Samanta et al. 2015; Samanta et al. 2016; Toprak et al. 2018); and (3) varying the solvent system of dyeing (Dong et al. 2019; Mu et al. 2019; Wang et al. 2018; Xia et al. 2018; Zhao et al. 2018).

Hydrated cotton cellulose contains negative charges on surfaces in the water system, therefore, anionic dyes such as direct and reactive dyes are mostly repulsed electrostatically by the fibers, requiring the use of electrolytes (i.e., inorganic salts) in assisting dye exhaustion. Therefore, any approach that can diminish the charge repulsion between cotton and dyes could reduce the use of the salts in the dyeing bath, which has triggered the practices of synthesis of new dyes (Zheng et al. 2012) and cationic modifications of cotton fibers (Farrell et al. 2014, 2017; Fu et al. 2013; Hashem et al. 2003; Hauser and Tabba 2001).
Herein, we introduce a new strategy to modify cotton fabrics with 2-diethylaminoethyl chloride (DEAE-Cl) through a unique reaction consisted of a nucleophilic substitution followed by a self-propagation of DEAE-Cl. Relying on the coexistence of nucleophilic and electrophilic sites in DEAE-Cl, four reaction paths could occur during the modification (Scheme 1): (1) DEAE-Cl directly reacts with alkaline-activated cotton fabrics to form a monolayer of tertiary amines on cotton; (2) DEAE-Cl undergoes self-propagation and stays in the modification solution; (3) the self-propagated DEAE-Cl reacts with alkaline-activated cotton fabrics; and (4) DEAE-Cl self-propagated on alkaline-activated cotton to generate multilayered quaternary ammonium salts.

The achievement of a high density of cationic quaternary ammonium salts on DEAE@Cotton makes the dyeing performance less sensitive to pH changes, which is different from the cotton fabrics modified with amines (Zhang et al. 2007; Zhu et al. 2017). The chemical structures of DEAE@Cotton fabrics were characterized with scanning electron microscope (SEM), Fourier-transform infrared spectroscopy (FTIR), thermogravimetric analysis (TGA), and Zeta potentials. The optimal dyeing performances of the DEAE@Cotton were achieved by carefully selecting different modification parameters, and the mechanical strength, color intensity, and wash fastness of the products were compared with that of the untreated cotton. The lightfastness and crocking fastness of dyed DEAE@Cotton were also evaluated.

### Experimental section

#### Chemicals and materials

Scoured and bleached woven cotton fabrics (Style # 400 plain woven weighting 98 g/m², 60 x 60), was purchased from TestFabrics Inc. (USA). Sodium hydroxide (NaOH), 2-diethylaminoethyl chloride hydrochloric acid (DEAE-Cl), sodium chloride (NaCl), denatured ethyl alcohol (EtOH), direct red 23 (DR23), direct yellow 27 (DY27), direct blue 71 (DB71), acid red 151 (AR151), and reactive blue 4 (RB4) were purchased from Sigma-Aldrich (USA). All chemicals were directly used as received.

Modification of cotton fabrics with DEAE-Cl

Pre-washed cotton fabrics were activated in NaOH/EtOH solution (120 g/L) for 30 min at room

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**Scheme 1** Potential reactions of cotton fabrics with DEAE-Cl under alkaline conditions
temperature, then the fabrics were transferred into a DEAE-Cl/EtOH solution with a specific DEAE-Cl concentration. The pH of the final solution was adjusted to around 12.0 using 1 M NaOH if needed. The fabrics were reacted under specific temperature and time with gentle stirring. After the modification, the DEAE@Cotton fabrics were washed with ethanol (5 min) and water (5 min) to remove any unreacted chemicals. The DEAE@Cotton was dried at 80 °C for 5 min and stored for further use.

Characterization of DEAE@Cotton fabrics

Fourier-transform infrared spectroscopy (FTIR) of fabric samples was obtained from a Nicolet 6700 FTIR spectrometer (Thermo Electron Co., USA) in absorbance mode at a range of 4000–400 cm\(^{-1}\) with 64 scans and 2 cm\(^{-1}\) resolution. The sample pellets were prepared by mixing 2 mg of fabric powders with 200 mg oven-dried potassium bromide (KBr). Thermogravimetric analysis (TGA) tests were performed to evaluate thermostability of cotton fabrics before and after DEAE-Cl modification. Specifically, fabric powders (~ 8 mg) were loaded in a TGA-60 system (Shimadzu Science Instruments, Inc., USA) with an aluminum pan. The samples were first heated to 120 °C and held for 2 min under nitrogen flow (30 mL/min) to eliminate all free water in the samples. Then, a heating rate of 10 °C/min was applied to reheat the samples from 35 to 600 °C under nitrogen environment (30 mL/min). Scanning electron microscope (SEM) images were captured with Quattro ESEM (Thermo Fisher Scientific, USA). The fabrics were coated with a thin layer of gold (~ 120 Å) by a gold sputter before SEM monitoring. Zeta potentials of untreated cotton and DEAE@Cotton were measured by Zetasizer Nano ZS90 with a DTS 1070 cuvette (Malvern Panalytical, UK). Specifically, the fabric samples were cut into fine powders and disperse in distilled water with 10 min of sonication (i.e., 100 mg fabric powder in 30 mL of H\(_2\)O). Different pH of the fabric suspension was adjusted by NaOH or HCl. The freshly prepared fabric suspension was used for zeta potential tests.

The tensile strengths of cotton fabrics before and after DEAE-Cl treatment were tested on an INSTRON 5566 tester (Instron Corporation; MA, USA) according to the American Society for Testing and Materials (ASTM) D5035-06 method. The tensile strength on the filling direction of the fabric was measured to represent the mechanical strength of unmodified cotton (named as Cotton in the following discussion) and DEAE@Cotton. The tensile strength recovery (TSR) was calculated from Eq. (1).

\[
\text{TSR} = \frac{T_{S_i}}{T_{S_C}} \times 100\%
\]

where \(T_{S_i}\) is the tensile strength in a unit of Newton (N) of treated fabrics and \(T_{S_C}\) is the tensile strength in N of Cotton fabrics.

A wicking test of fabrics was performed according to the American Association of Textile Chemists and Colorists (AATCC) 197 test method. Specifically, fabrics were cut in a size of 165 mm \(\times\) 25 mm and conditioned in a conditioning room (temperature = 21 °C, relative humidity = 65%) for at least 4 h before testing. Then, the fabric was vertically hung by immersing the other end of the fabric in a direct red 23 solution (50 mg/L) with a depth of 5 mm \(\pm\) 1 mm. After 30 min, the wicking distance was measured based on the length of dye vertically traveled. Each sample was tested in triplicates.

Dyeing fabrics with anionic dyes

Conventional and modified dyeing processes with direct dyes were applied on Cotton and DEAE@Cotton fabrics. Specifically, dye concentration of 2% owf was used in all dyeing processes. The dyeing baths with a liquor ratio of 50:1 were prepared with or without 5 g/L or 10 g/L NaCl. The fabrics were first rinsed with deionized water and squeezed before putting into the dyeing bath (pre-heated to 40 °C). Then, the temperature of the dyeing bath was increased to 60 °C within 5 min, and the fabrics were dyed under 60 °C with stirring for another 35 min. Finally, the dyed fabrics were washed with cold water for three times and dried at 80 °C for 5 min.

Characterization of dyed fabrics

A GretagMacbeth ColorEye 7000A tester (Akron, Ohio, USA) was used to measure the color coordinates \(L^*, a^*,\) and \(b^*\) and K/S values of the fabrics. The color difference (\(\Delta E\)) of dyed DEAE@Cotton over the conventionally dyed Cotton was calculated from Eq. (2). The color depth is evaluated by K/S values.
and can be calculated from the reflectance \( R \) of the fabrics based on Eq. (3).

\[
\Delta E = \sqrt{\left( L_C^* - L_D^* \right)^2 + \left( a_C^* - a_D^* \right)^2 + \left( b_C^* - b_D^* \right)^2}
\]

where \( L^* \), \( a^* \) and \( b^* \) refer to the color changing values from light to dark, red to green, and blue to yellow from positive to negative, respectively. The subscripts of \( C \) and \( D \) stand for the specific values of the conventionally dyed Cotton and the dyed DEAE@Cotton.

\[
K/S = \frac{(1 - R)^2}{2R}
\]

where \( K \) is the absorbance coefficient, and \( S \) is the scattering coefficient.

The wash fastness of Cotton and DEAE@Cotton fabrics after dyeing was measured with washing and drying machines according to standard method of AATCC LP1-2018. The color fastness was evaluated by the \( K/S \) values of dyed fabrics after each washing cycle.

The lightfastness of dyed fabrics was evaluated based on the fabric’s color change after 3 and 8 h of daylight exposures. Specifically, the dyed fabric was irradiated in a Spectrolinker™ XL-1500 UV crosslinker equipped with six cool white bulbs (6 × 15 W). The distance between the fabric and the light source was controlled as 15 cm. After irradiation, the color of the fabric was measured by the GretagMacbeth ColorEye 7000A tester. The \( \Delta E \) values were calculated from Eq. (2). The crocking fastness of the fabric was evaluated by using an AATCC crockmeter (Atlas Electric Devices Co., IL, USA) based on AATCC Test Method 8. A white cotton fabric was used as the reference to evaluate the color transfer during crocking fastness tests. The tests were performed in duplicates.

Dyeing mechanism and kinetics

The adsorption of dyes on Cotton and DEAE@Cotton fabrics was studied to fit either Langmuir (Eq. 4) or Freundlich isotherm model (Eq. 5). The fabrics were dyed in the dyeing bath with different initial concentrations (ranging from 10 to 200 mg/L) at room temperature (25 ± 2 °C), and DR23 was selected as the anionic dye representative. When the dyeing process reached the equilibrium (24 h), the residual dyeing solutions were scanned with a UV–vis spectrophotometer (Evolution 600, Thermo Scientific, USA), and the absorption intensity at the maximum wavelength of 520 nm (\( \lambda_{\text{max}} = 520 \) nm) was recorded for the measurements of the dye concentrations in the effluent (\( C_e \)) and calculation of the amounts of dyes adsorbed on the cotton fabrics (\( Q_e \)).

\[
Q_e = \frac{Q_mC_e}{1 + k_L C_e}
\]

where \( Q_m \) (mg/g) is the theoretical adsorption capacity of the fabrics, \( k_L \) and \( k_F \) (mg/g) are the adsorption constants in Langmuir and Freundlich models, respectively. And \( 1/n \) is a constant which indicates the strength of the adsorption between adsorbate and adsorbent.

The kinetics of the dyeing on Cotton and DEAE@Cotton were monitored by measuring the residual dye concentrations at different time intervals. The exhaustions (E%) of DR23 on Cotton and DEAE@Cotton at equilibrium were calculated according to Eq. (6), which were used to demonstrate the dye adsorption rates of Cotton and DEAE@Cotton.

\[
E(\%) = \left( 1 - \frac{C}{C_0} \right) \times 100\%
\]

where \( C \) is the dye concentration in the dyeing bath at specific time interval, and \( C_0 \) is the initial concentration of dye in the dyeing bath.

Results and discussion

Modification of DEAE@Cotton

Given the unique chemical structure of DEAE-Cl possessing both nucleophilic (tertiary amine) and electrophilic (R-Cl) reactive sites, alkaline-activated cotton can be cationized through a nucleophilic substitution reaction on R-Cl accompanied with the self-propagation of DEAE-Cl. Three reaction paths shown in Scheme 1 can introduce cationic moieties onto the cellulose structure (Tang et al. 2019, 2020), with the following properties provided on the fabrics: (1) covalently bonded cationic groups possessing excellent durability for dyeing, printing, and
laundering; (2) strong affinity to anionic dyes in a salt-free dyeing system; (3) improved color fastness compared with conventional dyeing methods; and (4) retention of mechanical properties of treated cotton fabrics. To optimize the dyeability of DEAE@Cotton, critical modification parameters were carefully studied.

As shown in Fig. 1a–c, NaOH concentration in the activation step highly affected the final dyeability of DEAE@Cotton since the alkaline condition not only could swell the cellulosic fibers to open more reactive sites to react with DEAE-Cl, but also ionize hydroxyl groups in cellulose structure to be more nucleophilic. By increasing the NaOH concentration from 0 g/L (control group) to 60 g/L, the ΔE of dyed fabrics over the conventional dyed Cotton ($L^* = 48.07$, $a^* = 54.17$, $b^* = 24.88$) decreased and K/S values increased gradually. Nevertheless, the dyeability was

**Fig. 1** a, d, and g Color differences (ΔE) and b, e, and h K/S values of DEAE@Cotton fabrics modified with various NaOH concentrations, DEAE-Cl concentrations, and DEAE-Cl modification temperatures, respectively. Standard conditions with NaOH concentration of 120 g/L, DEAE-Cl concentration of 1.0 wt%, and dyeing temperature of 25 °C were used in selection of each single variable. c, f, and i Optical images of dyeing performance of DEAE@Cotton fabrics obtained from different modification conditions.
improved significantly when cotton was pre-treated with 120 g/L NaOH by showing a ΔE and K/S value of 9.96 and 5.30 after a salt-free dyeing, which have been improved by 4.11 and 7.34 folds than that of the control group (i.e., Cotton dyed without salts), respectively (Fig. 1a, b). In fact, the alkaline activation step is similar to the mercerization process, which is a necessary preparation treatment of cotton fabrics during textile manufacturing.

The dyeability of DEAE@Cotton can also be varied according to the DEAE-Cl concentration employed in the reaction system, because the DEAE-Cl concentration will determine the occurrence of different reactions shown in Scheme 1 and the number of cations introduced by the quaternized groups. The propagation of DEAE-Cl on the DEAE-grafted cotton fabric is crucial to maximize the cationization of the fabric with limited reactive sites in cellulose structure. As shown in Fig. 1d–f, the ΔE of the dyed DEAE@Cotton compared to conventionally dyed Cotton decreased by increasing the DEAE-Cl concentration from 0 to 1.0 wt% and increased slightly when the concentration reached 5.0 wt%. In this case, the proposed reactions of (1), (3) and (4) are expected to coexist, maximizing the cationization of DEAE@Cotton. However, the further increment of DEAE-Cl concentration to 3.0 wt% or 5.0 wt% did not present better dyeing performance. This is caused by that the high DEAE-Cl concentration could accelerate the self-propagation of DEAE-Cl (reaction 2) and compete with the propagation reaction on cotton (reaction 3). Self-polymerized DEAE-Cl in reaction 2 has reduced reactivity with cellulose due to the steric hindrance to the C–Cl site and lower the number of cations on the cotton fibers. Therefore, the optimal DEAE-Cl concentration was selected as 1.0 wt% for further studies.

The reaction temperature in the DEAE-Cl modification step has a slight impact on the dyeability of DEAE@Cotton, and the color intensity on the dyed fabrics moderately increased (i.e., the ΔE changes oppositely) when raising the dyeing temperature from 25 to 80 °C (Fig. 1g–i). It further emphasized the importance of high concentration of cationic groups on the DEAE@Cotton, providing strong attractive force to anionic dyes. With an overall consideration of chemicals usages and energy costs, the optimal modification condition for preparation of DEAE@Cotton was selected as NaOH concentration, DEAE-Cl concentration, and modification temperature of 120 g/L, 1.0 wt%, and 55 °C, respectively in this study.

Characterization of DEAE@Cotton

Proposed chemical structure and properties of DEAE@Cotton were characterized and ascertained with SEM, FTIR, TGA, zeta potential measurements and wicking tests, as shown in Fig. 2. The incorporated cationic moieties on the DEAE@Cotton could highly improve the fabric dyeability according to an intermolecular electrostatic force shown in Fig. 2a. The positively charged quaternary ammonium groups facilitate the dye adsorption process by converting charge repulsion between Cotton and anionic dyes to an electrostatic attraction (i.e., the major interaction between DEAE@Cotton and anionic dyes). The SEM images of the cotton fabrics before and after DEAE-Cl modification present different fiber morphologies. The DEAE@Cotton revealed fibers with smoother surface but the fibers in Cotton were mostly in flat ribbon-shape (Fig. 2b). However, an external force, which applied during the cotton mercerization, was not imposed here during the alkaline treatment, so the cross-sectional view of the DEAE-Cotton fiber only shows slight swelling effect instead of presenting a round shape (Fig. 2b). The NaOH solution could effectively swell cotton fibers, ensuring the DEAE-Cl penetration into the fibers and improving the modification effectiveness. As shown in Fig. 2c, the FTIR results proved the chemical structure difference between Cotton and DEAE@Cotton. A subtracted spectrum of Cotton from DEAE@Cotton clearly shows the decrement of hydroxyl group (–OH) and the increase of methylene groups (–CH 2–) and ether groups (C–O–C) in the DEAE@Cotton, which expressed as a negative absorbance at 3480 cm$^{-1}$ and positive absorbances at 1490 cm$^{-1}$ and 1261 cm$^{-1}$, respectively (Tang et al. 2019, 2020; Yang et al. 2018). The changes of these characteristic peaks successfully demonstrated the formation of covalent linkages between cellulose and DEAE-Cl. Meanwhile, the different absorbance patterns around 2900 cm$^{-1}$ presented the difference of saturated C-H stretching between Cotton and DEAE@Cotton fabrics. The limited surface area of conventional cotton fabrics resulted in a relatively lower grafting ratio of DEAE moieties compared to that on surfaces of nanofibers (Tang et al. 2020). The DEAE moieties on
the DEAE@Cotton are lack of distinguishable FTIR characteristic peaks, making a challenge to obviously notice the structural differences of the DEAE@Cotton over the pristine cotton. However, our previous studies by adopting the same chemistry for nanofibrous membrane cationizations have already structurally demonstrated the success of proposed reaction (Tang et al. 2019, 2020).

Moreover, the TGA results (Fig. 2d) showed higher thermal degradation residues of DEAE@Cotton at 600 °C compared with Cotton, which is attributed to the addition of DEAE moieties in DEAE@Cotton. On the other hand, the cationization of cotton fabrics with DEAE-Cl is expected to increase the hydrophilicity of treated fabrics by having additional cationic moieties in DEAE@Cotton. As shown in Fig. 2e, the wicking distance of DEAE@Cotton achieved 8.2 cm while the distance of Cotton was only 6.5 cm. The tensile strength of DEAE@Cotton did not show any difference to the Cotton fabric, ensuring desired applications of DEAE@Cotton in textiles. On the other hand, the electrostatic charges (zeta potentials) of DEAE@Cotton under varied solution pH values were measured and compared with that of Cotton in aqueous system. The results are shown in Fig. 2f. With increasing the solution pH values from 3 to 11,
the zeta potentials of Cotton and DEAE@Cotton both gradually decreased, which is caused by the deprotonation of the hydroxyl groups in cellulose by hydroxyl anions. Although DEAE@Cotton only showed a positive zeta potential under an acidic condition (i.e., pH < 4), the average zeta potentials of the DEAE@Cotton were constantly around 10 mV higher than that of Cotton in the varied pH range (Fig. 2f), which is another evidence of the formation of cationic quaternary ammonium salts on the DEAE@Cotton. Since the zeta potential measurements were conducted in a fabric suspension, more negatively charged hydroxyl groups inside the fiber could expose in the suspension but were not accessible by the DEAE-Cl during the modification. The decrement of negative charges on the fabric surface weakened the charge repulsion toward anionic dyes and can eventually achieve better dyeability of DEAE@Cotton.

Dyeing performance on DEAE@Cotton

As shown in Fig. 3a–c, the dyeability of DEAE@Cotton with direct red 23 (DR23) without addition of inorganic salt is comparable to the Cotton dyed by the conventional method (10 g/L NaCl in the dyeing bath). The ΔE and K/S value of the dyed fabrics are shown as 6.471 and 8.924, respectively (Fig. 3a, b). Moreover, the addition of different amount of salt for dyeing DEAE@Cotton did not show obvious improvement in color intensity. These results further demonstrated the efficiency of the cotton cationization reaction with DEAE-Cl and the effectiveness of using DEAE@Cotton to liberate the usage of inorganic salts during cotton dyeing.

The direct dyed cotton fabrics normally have low wash fastness. By introducing a strong electrostatic interaction between cotton fabric and direct dye, the wash fastness of DR23 dyed DEAE@Cotton was highly improved (Fig. 3d). The K/S values of DR23 dyed Cotton dropped dramatically from 10.13 to 7.94 after one wash, whereas the K/S value of the DEAE@Cotton with one unit of salt, respectively. c Optical images of dyed fabrics. d Wash fastness of dyed fabrics within 20 cycles. e Scheme of the mechanisms of wash fastness on Cotton and 1.0 wt% DEAE@Cotton.
DEAE@Cotton dyed without salts only decreased by 0.48 units (changed from 10.57 to 10.09) (Fig. 3d). Furthermore, the K/S value of the dyed-DEAE@Cotton decreased gradually within the first four cycles of washing, and the color intensity remained constant at K/S = 8.62 after 20 washings, which was ∼2 units higher than that of the dyed-Cotton (Fig. 3d). The addition of salts in dyeing DEAE@Cotton presented slightly better wash fastness within the first five washing cycles. After 20 cycles of washing, however, it reached the similar color intensity (K/S = 8.52) as the one without addition of the salt (Fig. 3d). As shown in Fig. 3e, the presence of electrostatic interaction between DEAE@Cotton and DR23 not only accelerated the dye adsorption during the dyeing process, which resulted in higher color intensity on the fabric, but also reduced the dye desorption during the initial laundering, a sign of improved interactive force between dyes and the fabric. Overall, the incorporation of cationic DEAE moieties on cotton fabrics provided dual functions of eliminating the usage of salts in the dyeing process and enhancing the color fastness during daily laundering.

On the other hand, the lightfastness and crocking fastness of the dyed cotton products are critical factors to further evaluate the colorfastness of the DEAE@Cotton. After daylight irradiation for 3 and 8 h, the DR23 dyed-DEAE@Cotton only showed a ΔE value of 1.52 ± 1.05 and 1.92 ± 0.08, respectively. This color change is barely noticeable by naked eyes. Although using the cationized cotton for salt-free dyeing resulted in more dye molecules stayed on the fiber surfaces, the lack of photoactivity of DR23 and the increased distance between dyes and hydrogen donors (i.e., the cellulose back bone) by DEAE chains contributed to the good lightfastness of the dyed DEAE@Cotton fabrics. The crocking fastness of the dyed DEAE@Cotton fabrics was evaluated under dry and wet conditions according to AATCC Test Method 8-1996. The crocking fastness ratings were 5 (no color transfer to the reference cotton) and 3–3.5 (a slight color staining to the reference cotton) under dry and wet crocking, respectively.

Thermodynamics and kinetics of dyeing on DEAE@Cotton

Equilibrium adsorption isotherms of the dyeing processes were investigated to understand the dye adsorption mechanism and explicate the different dyeing properties of Cotton and DEAE@Cotton. Langmuir and Freundlich isotherm models were applied to describe the adsorptions of DR23 on the fabrics under equilibrium (24 h), and the results are shown in Fig. 4a, b and Table 1. Freundlich isotherm better described the dyeing process of Cotton with a higher correlation coefficient (R² = 0.9976), whereas the dyeing process of DEAE@Cotton is more consistent with the Langmuir isotherm model (R² = 0.9667), consistent with the electrostatic monolayer adsorption of anionic dyes on the cationized cotton fibers. The large number of cationic quaternary ammonium salts on the DEAE@Cotton provided strong attractive force to anionic dye molecules, thereby enhancing the subsequent dyeing process.

In addition, DEAE@Cotton can completely exhaust the dye molecules in solutions and leave no residual dyes in the dyeing bath due to the strong electrostatic interaction when the initial dye concentration was lower than 50 mg/L. And the exhaustions of 100 mg/L and 200 mg/L dyeing solutions reached 95.70% and 41.27%, respectively, which are 4.86 and 3.09 times higher than that of the Cotton. The dyeing kinetics within the dyeing duration (40 min) are shown in Fig. 4d. Nearly 50% of the dyes in the dyeing bath were exhausted by the DEAE@Cotton after 40 min dyeing, whereas less than 5.0% of the dyes were fixed onto the Cotton, which exacerbated burden to subsequent wastewater treatment. In this way, the improved dye exhaustion and adsorption kinetics of DEAE@Cotton further emphasized the advantage of introducing DEAE moieties onto cotton fabrics as a more environmentally friendly process.

Affinity of DEAE@Cotton with other anionic dyes

Possessing the cationic DEAE moieties, the DEAE@Cotton is expected to present an improved affinity to other anionic dyes under a salt-free condition as well. Therefore, four water-soluble anionic dyes that are classified in three different dye groups (direct dye, acid dye, and reactive dye) were used to dye Cotton and DEAE@Cotton (dye structures are shown in Fig. 5). The ΔE and K/S values of dyed fabrics were measured and displayed in Fig. 6. Not surprisingly, more intense colors and higher K/S values were achieved on the DEAE@Cotton without exceptions. Specifically, the ΔE between Cotton and
DEAE@Cotton dyed by DB71, DY27, AR151, and RB4 without adding salts are 24.53, 14.89, 17.82, and 21.10, respectively (Fig. 6a). Correspondingly, the K/S values of the dyed-DEAE@Cotton fabrics showed 7.37, 4.00, 3.81, and 7.85 times higher than the cases that were dyed on Cotton without addition of salts (Fig. 6b). The optical images of dyed fabrics are shown in Fig. 6c.

Table 1 Parameters of Langmuir and Freundlich isotherm models

| Fabrics       | Isotherm model | Langmuir | Freundlich |
|---------------|----------------|----------|------------|
|               |                | $k_L$ (mg/g) | $Q_m$ (mg/g) | $R^2$  | $k_F$ (mg/g) | $n$ | $R^2$ |
| Cotton        | Langmuir       | 0.059    | 2.239      | 0.8806 | 0.424       | 3.05 | 0.9976 |
| DEAE@Cotton   | Freundlich     | 3.825    | 6.376      | 0.9667 | 3.388       | 6.64 | 0.7317 |
Conclusion

DEAE@Cotton fabrics were fabricated through a nucleophilic substitution reaction between hydroxyl groups in cotton and 2-diethylaminoethyl chloride (DEAE-Cl), followed by a DEAE-Cl propagation on the fabric. The coexistence of nucleophilic and electrophilic sites in DEAE-Cl is attributed to the graft of tertiary amines onto cotton fabrics and a continuous growth of cationic quaternary ammonium salts, and eventually formed multiple cationic structures on one hydroxyl group. The proposed chemical structure of DEAE@Cotton was well demonstrated with FTIR, SEM, and TGA. DEAE@Cotton showed improved dyeability toward anionic dyes by introducing electrostatic interaction as the main force between anionic dyes and cotton fabrics.

Fig. 5 Anionic dyes used in this study

Fig. 6 a Color differences (ΔE) between dyed Cotton and dyed DEAE@Cotton and b K/S values of dyed fabrics with different anionic dyes. The labels in x-axial refer to different dyes and cotton samples (C refers to cotton and M refers to DEAE@Cotton). c Optical images of dyes fabrics with direct blue 71 (DB71), direct yellow 27 (DY27), acid red 151 (AR151), and reactive blue 4 (RB4) (from left to right). Cotton are on the left-hand side and DEAE@Cotton are on the right-hand side.
direct dyes and the fabric. The dyeing performance was competitive to the color intensity on the Cotton that was dyed under conventional condition yet with much better color fastness against daily laundering. The cationization of cotton fabrics with DEAE-CI successfully eliminated the application of inorganic salts in traditional cotton dyeing process, making the anionic dyeing of cotton fabrics more environmentally friendly.

Author contributions P. Tang: Conceptualization, Methodology, Validation, Writing-Original Draft; L. E. Lockett: Investigation, formal analysis; M. Zhang: Investigation; G. Sun: Conceptualization, Supervision, Writing-Review & Editing.

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Declarations

Conflict of interest Authors claim no conflict of interest.

Ethical approval No animal studies or human participants were involved in this study.

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