PROCESS FOR AFTERTREATMENT OF FRESHLY COLORED POLYAMIDE FIBERS

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10 Claims

ABSTRACT OF THE DISCLOSURE

An improvement in the dyeing and printing of polyamide fibers, comprising as an after-treatment the introduction of such fibers which have been freshly dyed or printed with a fiber reactive dyestuff, prior to the conventional drying, into an aqueous bath which contains certain halogenated organic products in combination with a certain kind of tertiary saturated aliphatic di-ter or tetra-amine; the pH of the bath is adjusted to about 4 to 5.5, and the bath containing the fibers is then heated also the aforesaid after-treatment baths per se.

This application discloses and claims only subject matter disclosed in our pending application Ser. No. 637,309, filed May 9, 1967, as a continuation application of our application Ser. No. 515,774, filed Oct. 22, 1965 as a divisional application under Rule 147 of our application Ser. No. 296,392, filed July 19, 1963, both last-mentioned applications being now abandoned.

Description of the invention

This invention relates to improvements in dyeing and printing on polyamide fibers as well as, industrial products, the dyed and printed polyamide fibers treated according to the invention.

Hitherto, wool and other polyamide fibers including nylon and the like synthetic polyamide fibers which had been dyed with reactive dyestuffs containing per molecule at least one substituent which can be split off as an anion during the dyeing and/or after-treatment of the dyed fibers, had to be subjected to an after-treatment in an aqueous basic medium of a pH of preferably about 7.5 or higher which contains, for example, sodium hydroxide, sodium carbonate, or preferably ammonia or hexamethylenetetramine as the basic agent.

This known alkaline after-treatment, which serves to fix on the fibers still unreacted dyestuff present in the dyed goods, suffers from the drawback that the polyamide fibers can be injured by the alkaline medium and particularly by a subsequent drying step if the same is carried out directly on the after-treated fibers.

Therefore, in the known dyeing methods, the after-treatment with an alkaline medium is followed usually by a further treatment in an acid medium prior to final drying of the dyed or printed fibers.

Another drawback of the after-treatment with alkaline agents such as ammonia or hexamethylenetetramine resides in the tendency of the alkaline bath to remove at least part of the still unreacted dyestuff from the fibers, whereby an exact control of the shade of the fixed dyings or prints is made very difficult.

It is, therefore, an object of the present invention to provide, in a process of dyeing or printing polyamide fibers with reactive dyestuffs of the type described, an after-treatment of the dyed fibers for the purpose of fixing still unreacted residual dyestuff on the fibers, which after-treatment does not injure the fibers, permits of drying the after-treated fibers directly without further intermediate treatments, and allows for a more exact control in producing a desired shade.

These objects are attained by the treatment according to the invention, which comprises, as an after-treatment in dyeing and printing processes for producing colored polyamide fibers by heating and thereby reacting said fibers with relative dyestuff, i.e. dyestuff containing at least one mobile substituent capable of being split off as anion, in an aqueous acid medium, preferably at a pH of about 4.5 to 5, and, preferably, with subsequent rinsing of the colored fibers in water at a temperature of about 10° to preferably 60°, and not more than 70° C., and drying.

(I) Introducing the freshly colored, and preferably immediately subsequently rinsed, fibers, directly thereafter and prior to drying, into an aqueous solution of a condensation product of

(a) a naphthalene sulfonic acid, in particular a monoo trisulfonic acid, and preferably naphthalene-2-monosulfonic acid, or a 1,2,3,4-tetrahydroxynaphthalene sulfonic acid,

(b) from about 0.5 to 1.5, and preferably from 0.8 to 1.2 parts by weight, per part by weight of (a), of a compound of the formula

HO-phenylene-SO₂-phenylene-OH, preferably 4,4'-dihydroxy-diphenylsulfone,

(c) from about 0.1 to 0.2, and preferably 0.1 part by weight, per part by weight of (a), of formaldehyde, in aqueous, preferably about 30%-solution,

(II) if necessary, adjusting the pH of the aqueous solution to about 4 to 5.5, and preferably to 4.5 to 5, by the addition of acid, such as acetic or formic acid, and preferably of acetic acid, and

(III) heating the said aqueous solution and the colored fibers therein for a short time, about 10 to 60, and preferably 15 to 30 minutes, at a temperature of about 20° to 100°, and preferably 60° to 85° C.

According to another mode of carrying out the treatment according to the invention in practice, the condensation product used as the fixing agent in the aqueous acid solution of steps (I) to (III) described supra, is produced by condensation of

(a) a α,ω-dichloro- or α,ω-dibromoalkane of from 2 to 7 carbon atoms, or α,ω'-dichloro-dialkylenether with a total of from 4 to 8 carbon atoms, with

(b) a tertiary saturated aliphatic amine with a total of from 6 to 24 carbon atoms and from two to four amino nitrogen atoms,

in such molar ratio of (a):(b) that approximately one halogen atom is present in the condensation reaction for every amino nitrogen atom.
According to yet another mode of carrying out the treatment according to the invention in practice, there is used in the aqueous solution of steps (i) to (III) supra, in lieu of the abovementioned condensation products, a product produced by the condensation of

(a) dicyanodiamide,
(b) at least one mole, per mole of (a), or an excess thereof, of an ammonia donor which is either urea or the urea-theta ratio of a mineral acid capable of splitting off ammonia, and
(c) at least two moles, per mole of (a), or an excess thereof, of formaldehyde in aqueous, preferably 30% by weight, solution. A concentration of about 0.5 gram of condensation product per liter of solution is the minimum.

In the second and third modes of the treatment according to the invention described above, the addition of acid to the solution in accordance with step II, supra, is mandatory.

As polyamide fibers, those of natural origin, above all wool and silk, but also synthetic polyamide fibers such as the various kinds of nylon e.g. nylon 6, nylon 66, and nylon 11 (Rilsan), are suitable for the after-treatment according to the invention.

These polyamide fibers are dyed by known methods in a dyestuff solution which, in addition to the dyestuffs mentioned, can also contain the auxiliaries usual in wool dyeing, e.g. salts such as sodium sulfate or ammonium sulfate and/or dilute acids, e.g. acetic or formic acid, and/or wetting agents, e.g. condensation products of fatty acids having at least 8 carbon atoms and lower alkanolamines such as are described in U.S. Patent 2,080,122.

The aforesaid fibers are printed by known processes in neutral or acid medium in the presence of thickeners and, optionally, urea.

The after-treatment according to the invention with the solution of the condensation products is preferably performed in a solution which contains these products in a concentration of about 0.5 to 3 g. per liter, furthermore, optionally, other additives usual in textile dyeing, and finally, acid, e.g. acetic or formic acid, as mentioned above.

The aforesaid condensation products suitable for use in the first above-described mode of carrying out the invention in practice are produced as described in German Patent 611,671, issued Apr. 5, 1935; those suitable for use in the second mode of practicing the invention described above, are produced as described in German Patent 894,237, issued Oct. 22, 1953. This class of condensation products is obtained by reacting the components mentioned above at a raised temperature, about 50 to 140° C., possibly in the presence of diolvents, e.g. butyl alcohol, ethylene glycol or diethylene glycol. Among the saturated aliphatic tertiary polynylamines defined above, there are:

Di-, tri- and tetra-amines such as N,N',N'-tetramethyl-hexamethylenediamine and homologues, N,N,N',N'-pentamethyl-di-ethylenetriamine, N,N,N',N'-diethylhexamethylenetriamine as well as industrial mixtures thereof also.

Examples of α,ω-dihalogen alkanes are 1,2-dichloro-1,2-diethoxymethane, 1,3-dichlorobutane, 1,5-dichloropentane, 1,6-dichlorohexane; α,ω-di-halogen ethers are, e.g. β,β'-di(chloroethy1 ether, β,β'-di-(2-chloroethoxy)-di-ethy1 ether, γ,γ'-dichloropropyl ether as well as glycerin dichlohydrox. The reaction products are most effective when the ratio of the reaction components is so chosen that there is about one halogen atom per N atom, that is, for example, when using a triamine, two moles of amine are quaternized with three moles of α,ω-di-halogen alkan. Particularly good results are obtained with the condensation product of 2 mols of N,N',N''-pentamethyl-diethylenetriamine and 3 mols of β,β'-dichloroethyl ether.

The condensation products usable in the third mode of carrying out the treatment according to the invention are produced from dicyanodiame and the ammonium salt of a strong mineral acid or urea, and formaldehyde, as described in German Patent 929,642, issued Oct. 22, 1953. Best results are obtained with compounds from this class which are produced in accordance with Example 2 of German Patent 929,642.

The reactive dyestuffs used according to the invention can be of the most varied classes of dyestuffs. Preferably they are nitro, azo, anthrachinone or phthalocyanine dyestuffs, which classes excel in stability. They contain water-solubilizing, acid, salt-forming groups, mainly sulfonic acid groups, possibly also carboxyl groups or sulfamyl groups, in the latter case, among others, also acylated sulfamyl groups, e.g. disulfimide and carbonyl sulfimide groups.

The advantages attained according to the invention are particularly impressive when the reactive dyestuffs conventionally used for the fast dyeing of cellulose are used which contain sulfonic acid groups, e.g. 2 to 4, per dyestuff molecule. Thus, examples of dyestuffs, dyeings of which are especially suitable for use in the treatment according to the invention are nitrodiarylamine sulfonic acids, metal-free mono- and poly-azo dyestuffs, heavy metal-containing, e.g. chromium or cobalt-containing, o-dihydroxy-, o-hydroxy-o-carboxy- and o-hydroxy-o-amino-azo-azo and dis-azo dyestuffs, copper-containing formazane dyestuffs, 1-amin-4-phenylamino anthraquinone dyestuffs, and copper phthalocyanines with substituted sulfamyl groups, all of which preferably contain at least two sulfonic acid groups. Mobile substituents which can be split off as anion are mainly: radicals of strong acids, in particular halogen atoms, of these preferably chloror, or bromine or fluo-
rine,—the mobility of which is due, for example, to the bond at β-carbon atoms in negatively substituted organic radicals, at SO3 groups in the case of fluorine, at carbon atoms adjacent to tertiary ring nitrogen in nitrogen heterocycles of aromatic character, in this case preferably 6-membered heterocycles having at least two tertiary ring nitrogen atoms—aromatically bound halogen atoms in α- and/or p-position to one (or more) electrophilic group(s), in particular fluorine or chlorine atoms; further, for example, the radical of sulfuric acid in sulfated β-hydroxy-alkyl compounds, e.g. in sulfated β-hydroxy-alkyl sulfon and sulfamyl groups; or the radical of hydroxy-aryl compounds in O-aryl urethane groups.

Dyestuffs usable according to the invention contain the substituent which can be split off as anion, for example, in the form of β-chloro- or β-bromofatty acid amide groups, in which case β-chloroacronic acid amide or β-bromopropionic acid amide groups are preferred; or the said substituent is in the form of fluoroo- or chloroo-nitrobenzoxylamino groups or fluoroo- or chloroo-nitrobenzenesulfamylamino groups in which the fluorine or chlorine atoms are in the α- or β-position of cellulose amide groups or are, preferably, in the form of chloro- or bromo-diazinylamino or triazinylamino groups, in which case it is in particular in the form of monochloro- or dichloro-a-triazinylamino groups and, preferably, di- or tri-chloro-p-aminolamino groups. Dyestuffs with at least one di- or tri-halogen pyrimidylamino and at least 2 sulfonic acid groups are preferred in the dyeing and printing processes comprising the after-treatment according to the invention.

An advantage of the treatment according to the invention over the known processes is the elimination of an alkali after-treatment with water or hexamethylenetetramine. This advantage becomes most apparent in the dyeing and printing of wool which is sensitive to alkali, as the danger of injury to the fibers inherent in the alkaline after-treatment is avoided.
Moreover, the control of producing the desired shade more exactly is facilitated since changes in shade due to shifting of the pH of the dyeing medium from acid to alkaline and possibly back to acid are avoided.

Dyeings or prints on polyamide fibers attained according to the invention are distinguished by pure, strong and even shades and good fastness properties. In particular, they have remarkable wet fastness properties, for example fastness to water, washing, milling and perspiration, and particularly to sea-water.

The following non-limitative examples illustrate the invention. Where not otherwise stated, parts and percentages are given by weight. The relationship of parts by weight to parts by volume is as that of grams to cubic centimeters. The temperatures are given in degrees centigrade.

**EXAMPLE 1**

100 parts of wool are dyed in 4000 parts of a dye liquor which contains the following agents:

4 parts of a dyestuff of the formula

![Chemical Structure]

0.5 part of a condensation product of 1 mol of oleic acid and 2 mols of diethanolamine (Example 5 of U.S. Patent No. 2,089,212), and 4 parts of 80% acetic acid.

The goods are introduced into the liquor at 50°, dyeing is performed for 10 minutes at this temperature, the bath is brought to the boil within 30 minutes and dyeing is continued at the boil for 45 minutes.

After dyeing, the goods are well rinsed and treated in a fresh bath for 20 minutes at 60–80° which contains 0.75 g. per liter of a condensation product of formaldehyde with dihydroxydiphenyl sulfone and naphthalene sulfonic acids, the production of which condensation product is described below, as well as 0.25 g. per liter of 80% acetic acid.

Valuable red dyeings are obtained in this way which are very wet fast.

The condensation product mentioned is obtained as follows:

100 parts of the sulfonating mixture, obtained by heating for several hours at 140–160°, 520 parts of naphthalene and 560 parts of concentrated sulfuric acid until water solubility is attained, are heated for about 1 hour at 105–110° with 100 parts of a dihydroxydiphenyl sulfone, 50 parts of water and 45 parts of formaldehyde (30%). The dihydroxydiphenyl sulfone is obtained by heating 540 parts of phenol and 180 parts of 60% oleum for 3 hours at 170–180° and distilling off excess phenol.

By using, instead of the dyestuff given in Example 1, the dyestuffs listed in column 2 of the following table, and otherwise following the procedure described in said example, wool dyeings of the shade given in column 3 and of good fastness to washing, perspiration and sea water are obtained.

<table>
<thead>
<tr>
<th>No.</th>
<th>Dyestuff</th>
<th>Shade on wool</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>![Chemical Structure]</td>
<td>Greenish-yellow.</td>
</tr>
<tr>
<td>2</td>
<td>![Chemical Structure]</td>
<td>Reddish-yellow.</td>
</tr>
<tr>
<td>3</td>
<td>![Chemical Structure]</td>
<td>Red.</td>
</tr>
<tr>
<td>4</td>
<td>![Chemical Structure]</td>
<td>Blue.</td>
</tr>
<tr>
<td>No.</td>
<td>Dyestuff</td>
<td>Shade on wool</td>
</tr>
<tr>
<td>-----</td>
<td>----------</td>
<td>--------------</td>
</tr>
<tr>
<td>5...</td>
<td><img src="image" alt="Compound" /></td>
<td>Scarlet</td>
</tr>
<tr>
<td>6...</td>
<td><img src="image" alt="Compound" /></td>
<td>Orange</td>
</tr>
<tr>
<td>7...</td>
<td><img src="image" alt="Compound" /></td>
<td>Turquoise blue</td>
</tr>
<tr>
<td>8...</td>
<td><img src="image" alt="Compound" /></td>
<td>Bordeaux</td>
</tr>
<tr>
<td>9...</td>
<td><img src="image" alt="Compound" /></td>
<td>Blue</td>
</tr>
<tr>
<td>10...</td>
<td><img src="image" alt="Compound" /></td>
<td>Black</td>
</tr>
<tr>
<td>11...</td>
<td><img src="image" alt="Compound" /></td>
<td>Orange</td>
</tr>
</tbody>
</table>

(Wherein Cu-Phtc is the copper phthalocyanine radical.)
EXAMPLE 2

Wool is printed with a printing paste of the following composition:
30 parts of the dyestuff of the formula

\[
\begin{align*}
\text{H}_2\text{N}-\text{C}=\text{C}-\text{NH}_2 \\
\text{C}=\text{C}-\text{NH}_{2} \\
\end{align*}
\]

100 parts of urea,
10 parts of the sodium salt of m-nitrobenzene sulfonic acid,
450 parts of 5% sodium alginate solution, and
410 parts of water.

The goods are then dried, steamed and well rinsed with cold water.

The rinsed wool is then treated for 20 minutes at 60°-80° in a fresh bath which contains 0.75 g. per liter of the formalddehyde/dihydroxydiphenylsulfone/naphthalene sulfonic acids condensation product used in Example 1, supra, as well as 0.5 g. per liter of aqueous 80% acetic acid.

In this way, valuable red prints are obtained which have very good wet fastness properties.

EXAMPLE 3

100 parts of nylon are dyed in 4000 parts of a dye liquor which contains 1.5 parts of a dyestuff of the formula

\[
\begin{align*}
\text{SO}_3\text{H} \\
\text{SO}_3\text{H} \\
\end{align*}
\]

and 1 part of 80% acetic acid.

The goods are entered into the liquor at 40°, dyeing is performed for 10 minutes at this temperature, the bath is brought to the boil within 30 minutes and dyeing is continued for 45 minutes at the boil.

After dyeing, the goods are well rinsed and treated for 20 minutes at 60°-80° in a fresh bath which contains 0.75 g. per liter of the formalddehyde/dihydroxydiphenylsulfone/naphthalene sulfonic acids condensation product mentioned in column 5, lines 16-32 of Example 1 and 0.25 g. per liter of 80% acetic acid.

In this way, valuable greenish yellow dyes which have very good wet fastness properties are obtained.

EXAMPLE 4

Nylon is printed with a printing paste of the following composition:

30 parts of the dyestuff of the formula

\[
\begin{align*}
\text{SO}_3\text{H} \\
\end{align*}
\]

50 parts of urea,
250 parts of boiling water,
45 parts of phenol,
30 parts of thiourea, and
450 parts of 5% sodium alginate solution, and
145 parts of water.

The goods are then dried and steamed and rinsed, first with cold and then with 60° warm water.

The rinsed goods are after-treated in a fresh bath in the manner described in the previous Example 3.

In this way, valuable greenish yellow prints are obtained, which have very good wet fastness properties.
EXAMPLE 5

100 parts of wool are dyed in 4000 parts of a dye liquor which contains the following agents:

4 parts of a dyestuff of the formula

\[
\begin{align*}
\text{Cl} & \quad \text{Cl} \\
\text{HO-S} & \quad \text{HO-S} \\
\text{NH} & \quad \text{C} \\
\text{C} & \quad \text{O} \\
\text{N} & \quad \text{N} \\
\text{O} & \quad \text{H}
\end{align*}
\]

0.5 part of a condensation product of 1 mol of oleic acid and 2 mols of diethanolamine (Example 5 of U.S. Patent No. 2,089,212) and 4 parts of 80% acetic acid.

These goods are introduced at 85°, dyeing is performed for 10 minutes at this temperature, the bath is brought to the boil within 10 minutes and dyeing is performed at the boil for 45 minutes.

After dyeing, the dyed goods are rinsed and treated for 20 minutes at 60–80° in a fresh bath which contains 0.75 g. per liter of the reaction product of 2 mols of \(N,N',N''\)-pentamethyl-diethylenetriamine with 3 mols of \(\beta,\gamma\)-dichlorodiethyl ether, and 0.25 g. per liter of 80% acetic acid. In this way, valuable vivid bluish red dyings are obtained which are very wet fast.

Dyings having similar good properties are obtained if, in the above example, instead of the reaction product mentioned, a nitrogen-containing condensation product of 34 parts of dicyanodiamide with 5.5 parts of ammonium chloride, 18 parts of urea and 80 parts formaldehyde is used in the after-treatment liquor and otherwise the same procedure is followed.

If, instead of the dyestuff mentioned in the example, those given in column 2 of Table II are used and otherwise the procedure described in the example is followed, then corresponding wool dyeings as given in column 3 are obtained.

<table>
<thead>
<tr>
<th>No.</th>
<th>Dyestuff</th>
<th>Shade on wool</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><img src="image1" alt="Dyestuff 1" /></td>
<td>Greenish yellow</td>
</tr>
<tr>
<td>2</td>
<td><img src="image2" alt="Dyestuff 2" /></td>
<td>Reddish yellow</td>
</tr>
<tr>
<td>3</td>
<td><img src="image3" alt="Dyestuff 3" /></td>
<td>Red</td>
</tr>
<tr>
<td>4</td>
<td><img src="image4" alt="Dyestuff 4" /></td>
<td>Blue</td>
</tr>
<tr>
<td>5</td>
<td><img src="image5" alt="Dyestuff 5" /></td>
<td>Scarlet</td>
</tr>
</tbody>
</table>
TABLE II—Continued

<table>
<thead>
<tr>
<th>No.</th>
<th>Dye stuff</th>
<th>Shade on wool</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td><img src="image1" alt="Image" /></td>
<td>Orange</td>
</tr>
<tr>
<td>7</td>
<td><img src="image2" alt="Image" /></td>
<td>Turquoise blue</td>
</tr>
<tr>
<td></td>
<td>(wherein CuPhls, is the copper phthiacusine radical)</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td><img src="image3" alt="Image" /></td>
<td>Bordeaux</td>
</tr>
<tr>
<td>9</td>
<td><img src="image4" alt="Image" /></td>
<td>Blue</td>
</tr>
<tr>
<td>10</td>
<td><img src="image5" alt="Image" /></td>
<td>Black</td>
</tr>
<tr>
<td>11</td>
<td><img src="image6" alt="Image" /></td>
<td>Orange</td>
</tr>
<tr>
<td>12</td>
<td><img src="image7" alt="Image" /></td>
<td>Greenish-yellow</td>
</tr>
<tr>
<td>13</td>
<td><img src="image8" alt="Image" /></td>
<td>Blue-black</td>
</tr>
</tbody>
</table>
EXAMPLE 6

Wool is printed with a printing paste of the following composition:

30 parts of the dyestuff of the formula

100 parts of urea,
10 parts of the sodium salt of m-nitrobenzene sulfonic acid,
450 parts of 5% sodium alginate solution, and
410 parts of water.

The goods are then dried, steamed and well rinsed with cold water.

The rinsed wool is after-treated as described in Example 1. In this way, valuable, vivid bluish red prints are obtained which have very good wet fastness properties.

Similar results are obtained if, instead of the reaction products as described in Example 1, a nitrogen-containing condensation product of 34 parts of dicyano-diamide with 12 parts of ammonium chloride, 12 parts of urea and 80 parts of formaldehyde are used.

EXAMPLE 7

100 parts of nylon are dyed in 4000 parts of a dye liquor which contains 1.5 parts of a dyestuff of the formula

and 1 part of 80% acetic acid.

The goods are introduced at 40°, dyeing is performed for 10 minutes at this temperature, the bath is brought to the boil within 30 minutes and dyeing is continued for 45 minutes at boiling temperature.

After dyeing, the goods are well rinsed and then treated in a fresh bath for 20 minutes at 60–80° which contains 0.75 g. per liter of the reaction product of 2 mols of N,N',N''-pentamethyl diethylene-triamine and 3 mols of p,p'-dichlorodiethyl ether, and also 0.5 g. per liter of 40% acetic acid. In this way, valuable greenish yellow dyings which have very good wet fastness properties are obtained.

Similarly good dyings are obtained if, in the above example, nitrogen-containing condensation product of 34 parts of dicyanodiamide with 11 parts of ammonium chloride, 12 parts of urea and 80 parts of formaldehyde is used in the after-treatment bath instead of the reaction product mentioned and otherwise the same procedure is followed.

EXAMPLE 8

Nylon is printed with a printing paste of the following composition:

30 parts of the dyestuff of the formula

50 parts of urea,
250 parts of boiling water,
45 parts of phenol,
30 parts of thiouethylene glycol,
450 parts of 5% sodium alginate solution and
145 parts of water.

The goods are then dried, steamed and rinsed first with cold and then with 60° warm water.

The rinsed goods are after-treated as described in Example 3. In this way, valuable greenish yellow prints which have very good wet fastness properties are obtained.

Similar results are further obtained if, instead of the reaction product as described in Example 3, a reaction product obtained from corresponding amounts of N,N',N''-tetramethyl-ethylene-diamine and γ,γ'-dichloropropylyether or 1,4-dibromobutane or corresponding amounts of N,N'-tetramethyl-tetraethylene-diamine and glycoldichlorohydrin is used.

We claim:

1. In a process for producing colored polyamide fibers, the improvement comprising

(I) introducing polyamide fibers freshly colored by heating and thereby reacting said fibers in an aqueous acid medium with reactive dyestuff which contains at least one mobile substituent capable of being split off as anion, after coloration and prior to drying, into an aqueous solution the solute of which consists essentially of the condensation product of

(a) a member selected from the group consisting of dichloralkane of from 2 to 7 carbon atoms, dibromo-alkane of from 2 to 7 carbon atoms, and dichloro-dialkylether with a total of from 4 to 8 carbon atoms, with

(b) a tertiary saturated aliphatic amine with a total of from 6 to 24 carbon atoms and from 2 to 4 amino nitrogen atoms, in such molar ratio of (a): (b) that approximately one halogen atom is present in the condensation reaction for every amino nitrogen atom;

(II) adjusting the pH of the aqueous solution to about 4 to 5.5, by the addition of acid, and

(III) heating the resulting acidified aqueous solution and the colored fibers therein for about 10 to 60 minutes, at a temperature of from about 20° to 100° C.
2. The improvement described in claim 1, wherein the condensation product is that of 2 moles of N,N',N''-pentamethyl-diethylenetriamine and 3 moles of β,γ'-dichlorodiethylether.

3. The improvement described in claim 1, wherein the tertiary saturated aliphatic amine is a member selected from the group consisting of N,N'-tetra-lower alkyldimethylene diamine, N,N',N''-penta-lower alkyldiethylene triamine, and N,N',N''',N''''-hexa-lower alkyltridethylene-tetramine.

4. The improvement described in claim 1, wherein the acid added in step (II) is a member selected from the group consisting of acetic acid and formic acid.

5. The improvement described in claim 1, wherein, in step (II), the pH is adjusted to about 4.5 to 5.

6. The improvement described in claim 1, wherein the aqueous acid solution and fibers therein are heated in step (III) at a temperature of about 60° to 65° C.

7. The improvement described in claim 1, wherein said reactive dyestuff contains a halogen-substituted pyrimidylamino reactive dyestuff radical and from 2 to 3 sulfonic acid groups.

8. The improvement described in claim 1, wherein said polyamide fibers are wool.

9. The improvement described in claim 1 wherein the concentration of said solute in said aqueous solution is about 0.5 to 3 g. per liter.

10. The improvement described in claim 1 wherein said dyestuff is selected from the group consisting of nitro, azo, anthraquinone phthalocyanine dyestuffs.

References Cited

UNITED STATES PATENTS

3,104,933 9/1963 Mendelsohn et al. 8--85
3,198,595 8/1965 Mawson et al. 8--18

FOREIGN PATENTS

1,277,635 10/1961 France.
894,237 10/1953 Germany.

OTHER REFERENCES


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U.S. Cl. X.R.