PROVISIONAL SPECIFICATION

Improvements in and relating to Colour Forming Developers and Processes of Colour Development

We, KODAK LIMITED, a Company registered under the Laws of Great Britain, of Kodak House, Kingsway, London, W.C.2, do hereby declare the nature of this invention which has been communicated to us by Eastman Kodak Company, a Company organised under the Laws of the State of New Jersey, United States of America, of 349, State Street, Rochester, New York, United States of America, to be as follows:—

This invention relates to improvements in colour forming developers and in processes of colour development for use in connection with colour photography.

It is known that coloured photographic images may be formed by using a developer which forms a coloured compound on development. The coloured compound thus formed is deposited adjacent to the silver grains of the silver image during the development. It is also known that a coloured image may be formed by adding to certain developer solutions a compound which couples, during development, with the oxidation product of the developing agent and forms a colouring substance which is likewise deposited adjacent to the silver grains of the silver image during development. Such a compound, which is employed in conjunction with a developing agent for the silver and which couples with the oxidation product thereof during development, is referred to herein as a colour coupler.

The present invention concerns new or improved colour forming developers comprising an aromatic amino compound serving as the developing agent and a colour coupler as hereinafter defined and also includes a new or improved colour development process which consists in developing a reducible silver salt image in a photographic element with the aid of an aromatic amino compound in presence of a colour coupler as hereinafter defined.

The silver can be removed from the image after colour development leaving a clear transparent dye image.

The invention also includes a photographic element having at least one layer containing a clear transparent image composed essentially of the product resulting from the coupling in situ during development of a colour coupler as hereinafter defined with the oxidation product of an aromatic amino developing agent.

When a silver halide emulsion containing a latent photographic image is developed, the silver halide is reduced to metallic silver and the developing agent is oxidized. The aromatic diamino compounds which have been used as developing agents form, on oxidation, products which will couple with colour couplers during development to form dyes. If such colour couplers are added to the developer solution, or incorporated in the emulsion layer, the dye which is thus formed by coupling during development is deposited in the gelatine or other silver halide carrier adjacent to the metallic silver grain. It is desirable that the dyes thus formed should not readily wander from the place of formation. It is accordingly desirable that they should be insoluble in water. They are not physically attached to the silver grain and therefore the silver may be subsequently bleached out of the carrier layer leaving a pure dye image.

Numerous substances have hitherto been employed or proposed as colour couplers among which may be mentioned phenols, napththols, cresols, nitrophenyl-acetonitriles and acetocetic esters. It has not, however, always been possible among those hitherto available to select one which exhibits all the desired combination of properties required for any specific case. In colour-developing a gelatino-silver halide emulsion layer it is necessary to select a colour coupler which will give just the desired shade in conjunction with the colours which are produced in other layers. It is moreover important to employ a colour coupler
which gives a coloured compound which is resistant to the normal processing baths employed, although it may often be desirable to have one which gives a colour which can be destroyed and/or removed if desired during some step in the processing. Many of the color couplers employed according to the present invention are suitable in carrying out the processing described in our prior patents No. 427,472 427,516, 427,518, 427,520 440,092, 440,098 and application No. 16012/35 (Serial No. 447,092).

According to the present invention the substances employed as colour couplers are compounds having the formula:

\[ \text{CH}—\text{CH}—\text{CO}—\text{R} \]

where R is a substituted benzene nucleus or an alkyl group.

Examples of colour couplers according to the foregoing definition are:

- \( p \)-Tolylacetanilide
- \( p \)-Dimethylbenzoylacetonitrile
- \( p \)-Bromobenzoylacetonitrile
- \( m \)-Nitrobenzoylacetonitrile
- \( p \)-Nitrobenzoylacetonitrile
- \( 3 : 5 \)-Dibromo-4-methoxybenzoylacetonitrile
- 1;3-di-(o-cyanoacetyl)-benzene

Cyanoacetone

The colours given by the colour couplers named above when used in conjunction with \( p \)-aminodiethylamine, for example, as developing agent are red to magenta except the last compound which gives a yellow.

The aromatic amino compounds which may be used as developing agents in the present invention include the mono-di- and tri-amino aryl compounds. Among the monoamino compounds may be mentioned the aminophenols, and aminocresols and their halogen substituted derivatives as well as the amino-naphthols. The developing agents usually used are the dianyo compounds such as para-phenylene diamine and its substitution products. These developers may be substituted in the amino groups or in the ring or in both, forming compounds such as the alkyl phenylene diamines, tolylene-diamines, alkyl-toluylene diamines and aminodiphenyamines. These compounds are usually kept in the salt form such as hydrochloride or sulphate since these are more stable than the amines themselves. As examples of developing agents of this class, there may be mentioned diethyl para-phenylene diamine, mono-methyl para-phenylene diamine, dimethyl para-phenylene diamine and ortho-amine-diethylamine.

The present invention may be utilised in the formation of coloured photographic images on plates or papers as well as on films and the dyes will be formed when gelatin or other carrier for the silver halide is used. The plates, films or papers may have differently sensitized emulsions of the mixed grain type or superimposed on one side or on both sides of the support. The dyes formed may be decolourized by an oxidizing agent such as chromic acid and colourless soluble compounds thereby formed.

The bleaching of the dye in this manner need not destroy the silver image but may convert it into a developable silver salt image which can in turn be coloured, bleached and recoloured a number of times.

Dated this 21st day of May, 1937.

W. P. THOMPSON & CO.,
12, Church Street, Liverpool, 1,
Chartered Patent Agent.

COMPLETE SPECIFICATION

Improvements in and relating to Colour Forming Developers and Processes of Colour Development

We, KODAK LIMITED, a Company registered under the Laws of Great Britain, of Kodak House, Kingsway, London, W.C.2, do hereby declare the nature of this invention which has been communicated to us by Eastman Kodak Company, a Company organised under the Laws of the State of New Jersey, United States of America, of 345, State Street, Rochester, New York, United States of America, and in what manner the same is to be performed, to be particularly described and ascertained in and by the following statement:

This invention relates to improvements in colour forming developers and in processes of colour development for use in connection with colour photography. It is known that coloured photographic images may be formed by using a developer which forms a coloured compound on development. The coloured compound thus formed is deposited adjacent to the silver grains of the silver image during the development. It is also known that a coloured image may be
formed by adding to certain developer solutions or by incorporating in the gelatino-silver halide emulsion before or after exposure a compound which couples, during development, with the oxidation product of the developing agent and forms a colouring substance which is likewise deposited adjacent to the silver grains of the silver image during development. Such a compound, which is employed in conjunction with a developing agent for the silver and which couples with the oxidation product thereof during development, is referred to herein as a colour coupler.

The present invention concerns new or improved colour forming developers comprising an aromatic amino compound serving, as the developing agent and a colour coupler as hereinafter defined and also includes a new or improved colour development process which consists in developing a reducible silver salt image in a photographic element with the aid of an aromatic amino compound in presence of a colour coupler as hereinafter defined as well as the photographic image forming therefrom. It also includes photographic sensitive elements having such a colour coupler incorporated in one or more emulsion layers. The silver can be removed from the image after colour development leaving a clear transparent dye image.

The invention also includes a photographic element having at least one layer containing a clear transparent image composed essentially of the product resulting from the coupling in situ, during development of a developable silver salt, of a colour coupler as hereinafter defined with the oxidation product of an aromatic amino developer agent.

When the silver halide emulsion containing a latent photographic image is developed, the silver halide is reduced to metallic silver and the developing agent is oxidised. The aromatic diamino compounds which have been used as developing agents form, on oxidation, products which will couple with colour couplers during development to form dyes. If such colour couplers are added to the developer solution, or incorporated in the emulsion layer, the dye which is thus formed by coupling during development is deposited in the gelatine or other silver halide carrier adjacent to the metallic silver grain. It is desirable that the dye thus formed should not readily wander from the place of formation. It is accordingly, desirable that they should be insoluble in water. They are probably not physically attached to the silver grain. The silver may be subsequently bleached out of the carrier layer leaving a pure dye image.

Numerous substances have hitherto been employed or proposed as colour couplers among which may be mentioned phenois, napthols, cresols, nitrophenylacetonitriles and acetoacetic esters. It has not, however, always been possible among those hitherto available to select one which exhibits all the desired combination of properties required for any specific case. In colour-developing a gelatino-silver halide emulsion layer it is necessary to select a colour coupler which will give just the desired shade in conjunction with the colours which are produced in other layers. It is moreover important to employ a colour coupler which gives a coloured compound which is resistant to the normal processing baths employed, although it may often be desirable to have one which gives a colour which can be destroyed and/or removed if desired during some step in the processing. Many of the colour couplers employed according to the present invention are suitable in carrying out the processing described in our prior patent Nos. 427,472, 427,516, 427,518, 427,520, 440,032, 440,089 and 447,092.

According to the present invention the substances employed as colour couplers are compounds having the formula:

\[ \text{R} - \text{CO} - \text{CH}_2 - \text{CN} \]

Where R is a substituted phenyl group (not including two closed rings) or an alkyI group.

Typical examples of suitable couplers are:

1. \[ \text{CO} - \text{CH}_3 - \text{CN} \]
   \[ \text{CH}_3 \]
   \[ p\text{-toluylacetonitrile} \]

2. \[ \text{CO} - \text{CH}_2 - \text{CN} \]
   \[ \text{H}_2\text{C} \]
   \[ 2: 4\text{-Dimethylbenzoylacetonitrile} \]

3. \[ \text{CO} - \text{CH}_2\text{CN} \]
   \[ \text{Br} \]
   \[ p\text{-Bromobenzoylacetonitrile} \]
CO—CH₃—CN
\[ m\text{-nitrobenzoylacetonitrile} \]

CO—CH₃—CN
\[ p\text{-nitrobenzoylacetonitrile} \]

CO—CH₃—CN
\[ 3:5\text{-Dibromo-4-methoxybenzoylacetonitrile} \]

Br
\[ Br \]

OCH₃

CO—CH₃—CN
\[ 1:3\text{-di-(o-cyanocetyl)-benzene} \]

CN—CH₃—CO—CH₃
\[ cyanacetone \]

These compounds, when present during the development of a silver salt with an aromatic amino developing agent, couple with the oxidation product of such developing agent forming a dye simultaneously with the formation of the silver image. The coupling occurs in each case at the methylene group contained between the carbonyl group and the cyano group of the coupler.

To prepare p-tolylacetocetitrile (M.P. 105—106° C.), p-bromobenzylacetocetitrile (M.P. 161—162° C.), m-nitrobenzylacetocetitrile (M.P. 148—150° C.), and 3:5-dibromo-4-methoxybenzoylacetonitrile (M.P. 78—81° C.), the following bromo-ketones were obtained by bromination of the methyl ketones in chloroform, carbon tetrachloride or other inert solvent.

\[ p\text{-methylphenacyl bromide (Ber. 30, 577 (1897))} \]

\[ p\text{-bromophenacyl bromide (Bl. 3, 21, 67)} \]

\[ m\text{-nitrophenacyl bromide (Ber. 22, 204)} \]

\[ 3:5\text{-dibromo-4-methoxyphenacyl bromide (M.P. 93—95° C.)} \]

The corresponding cyanomethyl ketone derivatives were prepared from the above halogenated ketones by adding a warm aqueous solution of 3 molecular amounts of sodium or potassium cyanide to a warm alcohol solution of the halogenated ketone and heating for ½ to 1½ hours on a steam bath. The product was obtained by pouring the resulting reaction mixtures into 3 volumes of dilute acid and recrystallizing the resulting crude material from a suitable solvent.

2:4-dimethyl phenacyl chloride (M.P. 61—62° C.) was prepared in a similar manner to that described by Schroeter, Müller, Ruang, Ber, 62, 655 (1929) for e-naphthacyl chloride. In the manner described above the corresponding cyanomethyl ketone was prepared, 2:4-dimethylbenzoylacetonitrile, M.P. 80—81° C.

1:3-di-(o-cyanocetyl) benzene was prepared in a manner similar to that described by Fuson and Beveridge (J.A. C.S. 53, 1896, (1931)) and melted at 156—157° C.

p-Nitrobenzoylacetonitrile was prepared by mineral acid hydrolysis of o-cyano-p-nitrobenzoylacetamide in a manner similar to that described by H. Gevekohl for the preparation of o-nitrobenzoylacetonitrile from o-nitrobenzoylacetate ester (Ann. 231, 392). The product melted at 178—180° C.

The aromatic amino compounds which may be used as developing agents in the present invention include the mono-di, and tri-amino aryl compounds. Among the monamine compounds may be mentioned the aminophenols and aminocresols and their halogen substituted derivatives as well as the amino-naphthols. The developing agents usually used are the diamino compounds such as para-phenylene diamine and its substitution products. These developers may be substituted in the amino group or in the ring or in both, forming compounds such as the alkyl phenylene diamines, tolylenediamines, alkyl-tolylene-diamines and aminodiphenylamines. These compounds are usually kept in the salt form such as hydrochloride or sulphate since these are more stable than the amines themselves. As examples of developing agents of this class, there may be mentioned diethyl para-phenylene diamine, mono-methyl para-phenylene diamine, dimethyl para-phenylene diamine and ortho-amino-diethylaniline.

As would be expected from the behaviour of known colour couplers the shade of the colour obtained by coupling generally varies in accordance with the developing agent selected.

EXAMPLE:
A developing formula which may be used is the following:—
A

Diethyl para phenylenediamine hydrochloride - - 1 gram
Sodium sulphite - - - - - - - 0.5 gram
Sodium carbonate - - - - - - - 20 grams
Water - - - - - - - 1 litre

B

Colour coupler The molecular equivalent of the developer.
Water miscible solvent, such as acetone - - - 50 c.c.

For use, B is added to A.

The developing agent and the proportions of the ingredients used in the above formula may, of course, be varied. Solvents other than acetone, such as alcohols, may also be used.

The colours formed by the compounds named above on coupling with the oxidation product of the developer are red to magenta except in the case of compound No. 8 which gives a yellow.

When \( R \) is a substituted phenyl group, the colour of dyes formed from couplers having additional substituents in this group may vary according to the type of additional substituent employed.

Although we have described our invention with particular reference to the use of the colour coupler in the developing solution itself, our invention is in no way limited to this method. As an alternative method, the colour coupler may be incorporated in the photographic layer before development, and either before or after exposure. It may be absorbed upon the sensitive silver halide grains.

The present invention may be utilized in the formation of coloured photographic images on plates or papers as well as on films employing gelatine or other carrier for the silver halide. The plates, films or papers may have differently sensitized emulsions of the mixed grain type or superimposed on one side or on both sides of the support. The dyes formed may be decolourized by an oxidizing agent such as chromic acid and colourless soluble compounds thereby formed. The bleaching of the dye in this manner need not destroy the silver image but may convert it into a developable silver salt image which can in turn be coloured, bleached and recoloured a number of times.

The accompanying drawing shows the absorption characteristics of the dyes formed when \( A \), the known benzoylacetanitrite and \( B \), \( m \)-nitrobenzoylacetanitrite, respectively are employed as the couplers. The drawing represents graphs in which the density of the coloured images are represented on the ordinate and the wave lengths on the abscissa.

Having now particularly described and ascertained the nature of the said invention and in what manner the same is to be performed, as communicated to us by our foreign correspondents, we declare that what we claim is:

1. A colour forming developer comprising an aromatic amino developing agent and a colour coupler having the formula:

\[ R \text{--CO--CH}_2\text{--CN} \]

where \( R \) is a substituted phenyl group (not including two closed rings) or an alkyl group.

2. A colour forming developer as claimed in claim 1 in which \( R \) is a phenyl group containing one or more alkyl constituents.

3. A colour forming developer as claimed in claim 1 or 2 in which the developing agent is an aromatic diamino compound.

4. A colour forming developer as claimed in claim 3 in which the aromatic diamino compound is an alkyl substituted phenylene diamine.

5. A process of colour development which includes developing a reducible silver salt image with an aromatic amino developing agent in presence of a colour coupler as defined in claim 1.

6. A process of colour development which includes developing a reducible silver salt image with a colour forming developer as claimed in any of claims 1 to 4.

7. A photographic element having a layer containing a clear transparent dye image composed essentially of the product resulting from the coupling in situ,
during development of a developable
silver salt, of a colour coupler as defined
in claim 1 or 2 with the oxidation pro-
duct of an aromatic amino developing
agent and subsequent removal of
metallic silver.
8. A sensitive photographic element
having a colour coupler as defined in
claim 1 or 2 incorporated in one or more
emulsion layers.
9. The colour forming developers and
methods of colour development employ-
ing the colour couplers hereinbefore par-
ticularly described, in conjunction with
aromatic amino developing agents, sub-
stantially as herein described.
10. In the production of colour photo-
graphic records especially multi-layer
records, the employment in conjunction
with aromatic developing agents, of 20
colour couplers of the nature herein
defined.

Dated this 21st day of May, 1937.
W. P. THOMPSON & CO.,
12, Church Street, Liverpool,

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