

(19)



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(11)

EP 0 622 665 B1

(12)

EUROPEAN PATENT SPECIFICATION

(45) Date of publication and mention
of the grant of the patent:
02.07.1997 Bulletin 1997/27

(51) Int Cl.⁶: **G03C 1/09, G03C 1/16**

(21) Application number: **94106432.1**

(22) Date of filing: **25.04.1994**

(54) **Silver halide photographic light-sensitive material**

Lichtempfindliches, photographisches Silberhalogenid-Material

Matériau photographique à l'halogénure d'argent sensible à la lumière

(84) Designated Contracting States:
DE

(72) Inventor: **Suzuki, Tetsuya**
Hino-shi, Tokyo (JP)

(30) Priority: **26.04.1993 JP 99557/93**

(74) Representative: **Henkel, Feiler, Hänzel & Partner**
Möhlstrasse 37
81675 München (DE)

(43) Date of publication of application:
02.11.1994 Bulletin 1994/44

(73) Proprietor: **KONICA CORPORATION**
Tokyo 163 (JP)

(56) References cited:
EP-A- 0 443 453 **EP-A- 0 487 010**
EP-A- 0 512 496 **EP-A- 0 514 675**
EP-A- 0 563 708

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Description

Field of the Invention

5 This invention relates to a silver halide photographic light-sensitive material and, particularly to a silver halide photographic light-sensitive material for regular-type X-ray photographic use, that is suitable for rapid development process.

Background of the Invention

10 In recent years, a high-temperature and rapid-processing has become feasible because of the introduction of an automatic processor, and a super-rapid processing for not longer than 45 seconds in a Dry-to-Dry system has been developed. In this type of systems, on the other hand, it has been customary to make a hardening reaction in the course of the processing steps, for providing a rapid processing aptitude.

15 Conventionally, glutaraldehyde has been used as a hardener, because of a high reactivity and a rapid hardening in a developer. However, it has not been admittable that glutaraldehyde is high in safety, because it has an irritating odor, or if it adheres to a human body. Further, with raising the issue of environmental protection, a rapid processing procedure has been so developed as to eliminate a hardening development in which glutaraldehyde is used.

20 For performing a rapid photographic processing without relying on such a hardening development as mentioned above, a silver halide photographic light-sensitive material is required to be satisfactorily hardened in advance. However, in the case of a silver halide photographic light-sensitive material for medical use that is required to have a high sensitivity and a high density (Dmax) and if the light-sensitive material is satisfactorily hardened in advance, a large amount of silver coated is required to provide desired photographic characteristics. An increase in silver coverage brings about, not only increased costs is increased, but also such a serious trouble as a lowering of rapid processability, 25 an increase of fog and the deterioration of pressure resistance.

In recent years, in a silver halide photographic light-sensitive material for medical use, there have increased the transfer from a regular system to an ortho system combined with a rare-earth screen. In the latter system, a tabular silver halide emulsion has been used, because a large amount of ortho-sensitizing dyes can be adsorbed thereto; a high density can be obtained by a relatively small amount of silver coverage; a density-down is relatively small even when sufficiently hardened in advance; and desired photographic characteristics can be obtained even without further hardening. The facts mentioned above are also disclosed in, for example, Research Disclosure No. 22534, Jan., 1983, and so forth.

30 It has, however, been known that a tabular silver halide emulsion is not good in pressure resistance, such as an abrasion blackening trouble and a roller-mark produced through an automatic processor in a rapid processing, and that a pressure resistance and a processing stability are further deteriorated by increasing the hardening degrees of an emulsion layer.

35 As compared to a cubic, octahedral, spherical-formed silver halide grain, it has been known that a tabular grain having an equivalent volume is lowered in an inherent sensitivity (in a blue region) because of being thinner in thickness. As one of the improving means thereof, an increasing of an iodide content of a grain has been known to be effective. 40 However, it has been very difficult to apply a tabular grain to a regular-type (or a blue-sensitive-type) light-sensitive material, because a rapid processability is deteriorated.

On the other hand, there has been known a technique for sensitizing an inherent sensitivity by making use of a blue-sensitizing dye. However, some kind of sensitizing dyes has a property to reduce the inherent absorption of silver halide, therefore, the technique has been so disadvantageous that the technique is difficult to be properly performed.

45 Japanese Patent Publication Open to Public Inspection (hereinafter referred to JP OPI Publication) No. 59-55426/1984 discloses a technique in which the processing stability of a tabular grain having an aspect ratio of not lower than 3 is improved by making use of a blue-sensitizing dye. However, this technique is to carry out rapid processing at a high temperature in which an aldehyde type hardener (or glutaraldehyde) is contained in a developer, that is not any technique relating to a rapid processing without having a development hardening step, nor any technique for enhancing a spectral sensitization.

50 JP OPI Publication No. 4-291 338/1992 discloses a sensitizing technique in which zeromethine dye is used. However, this technique has had such a problem that not only the absolute value of a sensitivity is substantially low, but also a pressure resistance is not improved, and particularly that the characteristics are seriously deteriorated when sufficient hardening in advance for carrying out a non-hardening development. Therefore, a further improved technique has been demanded.

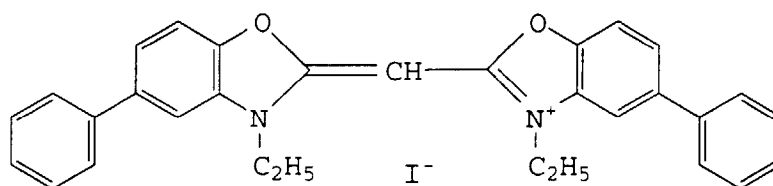
55 EP-A-0 563 708 discloses a silver halide photographic emulsion the silver halide grains of which are chemically sensitized with a selenium sensitizer after their formation. A similar disclosure can be found in EP-A-0 514 675 according to which the selenium sensitizer is added before or during the chemical sensitization. According to EP-0 512 496 a

Examples of the compounds

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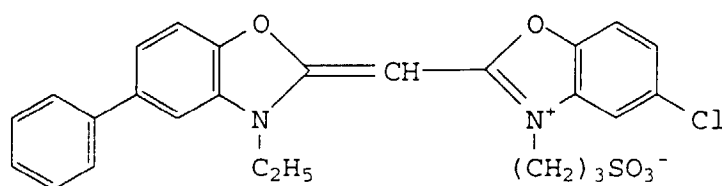
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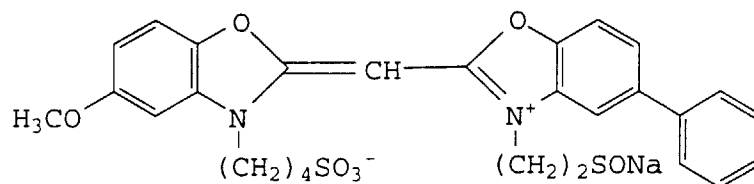


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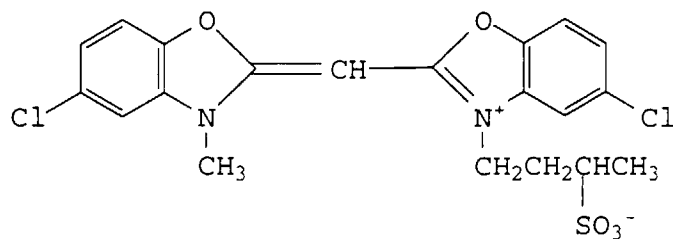
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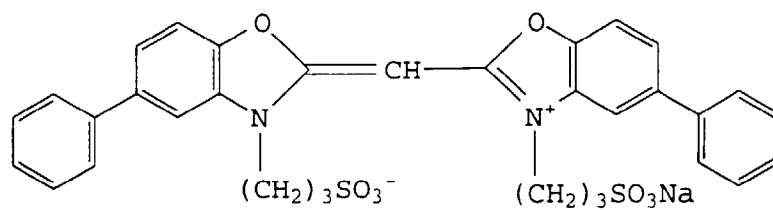


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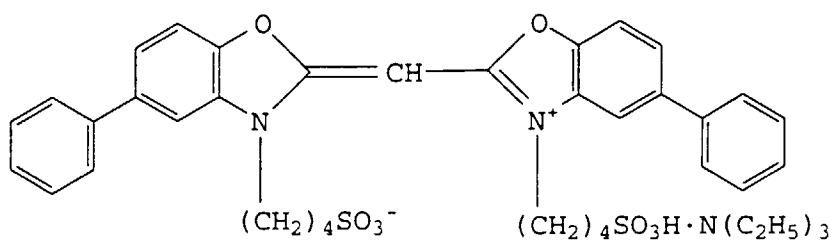


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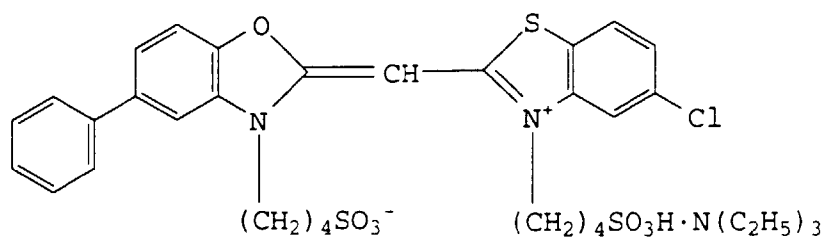
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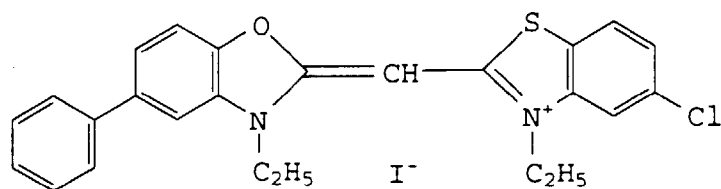


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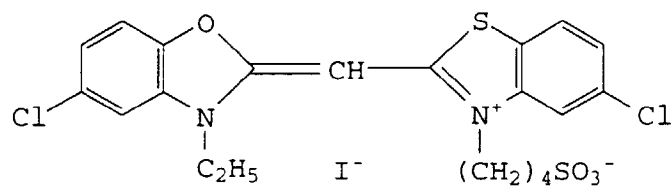


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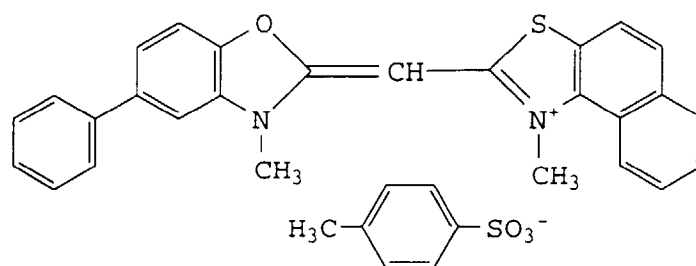


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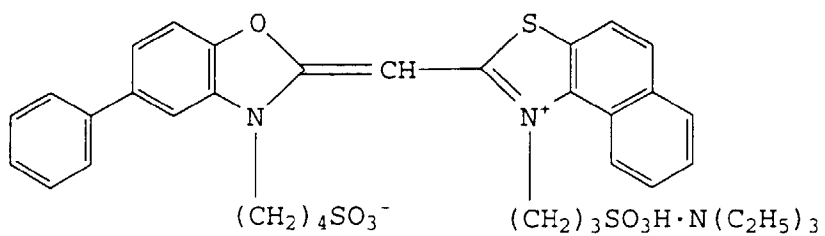


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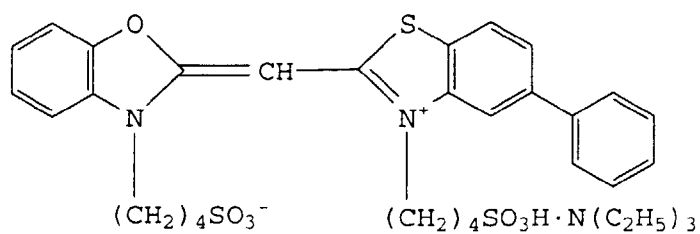


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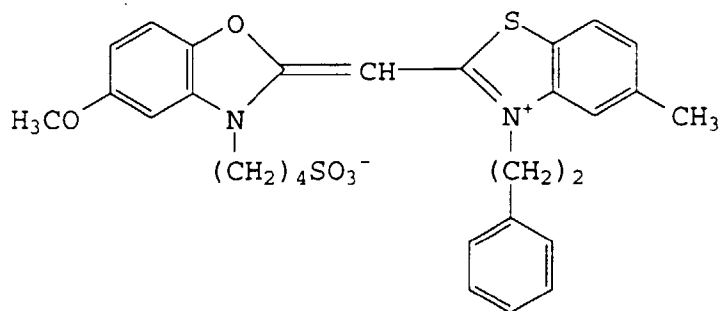
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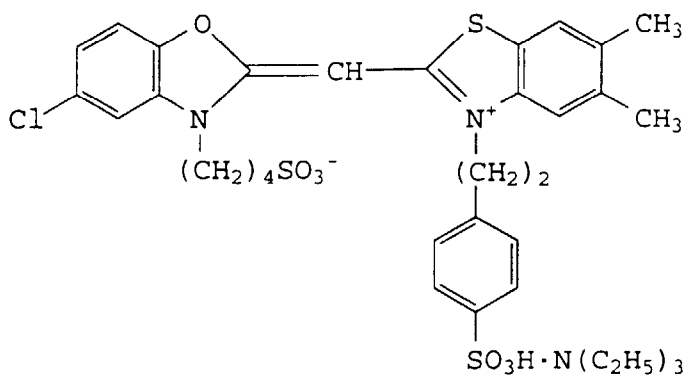


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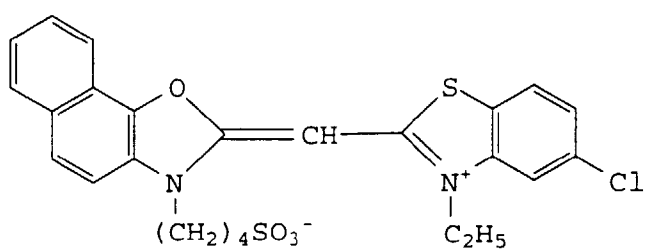
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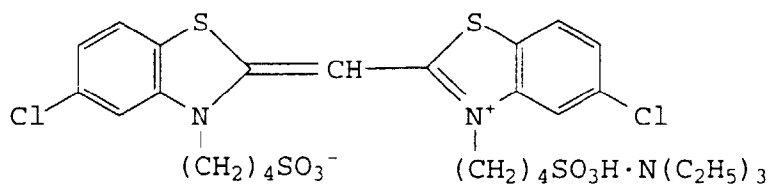


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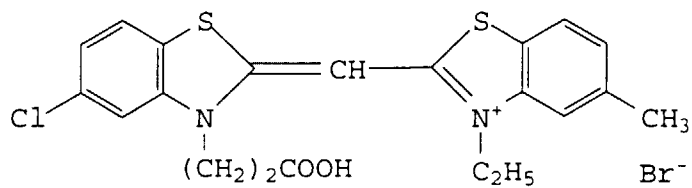
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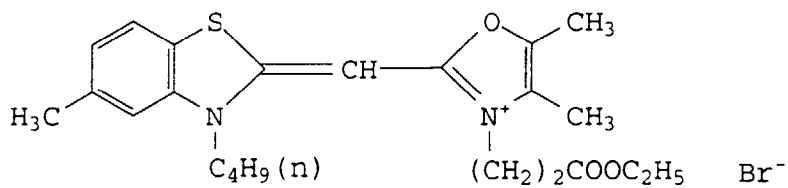


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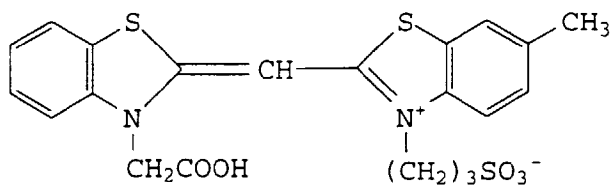
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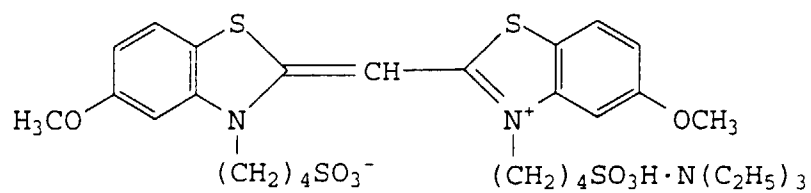


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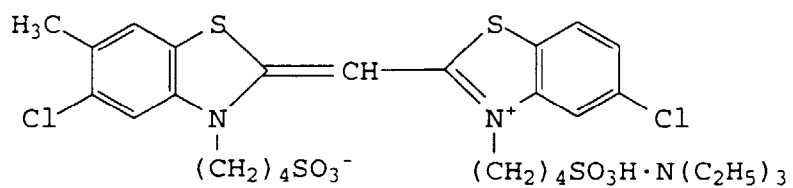
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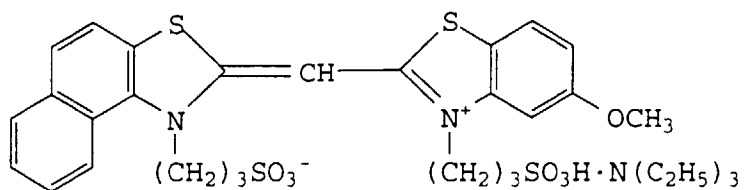
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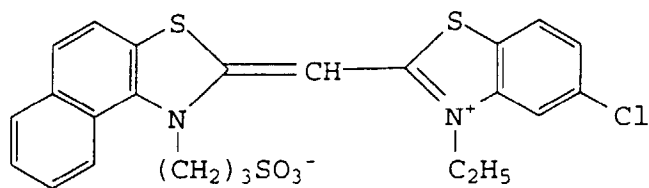
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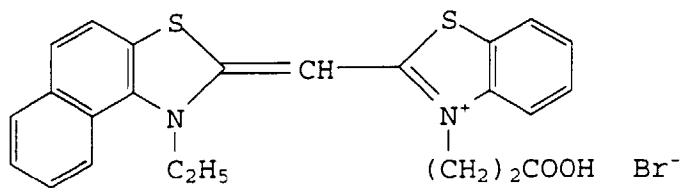
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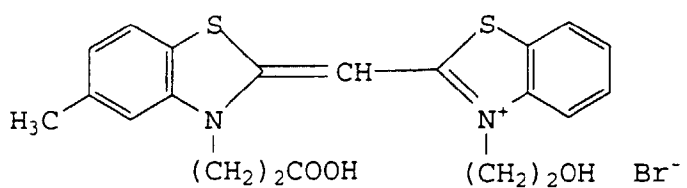
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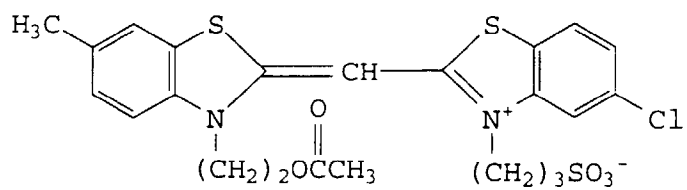


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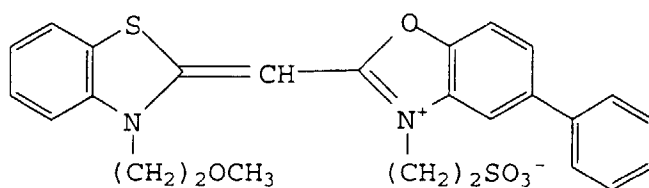
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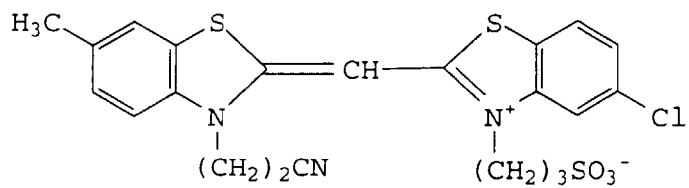
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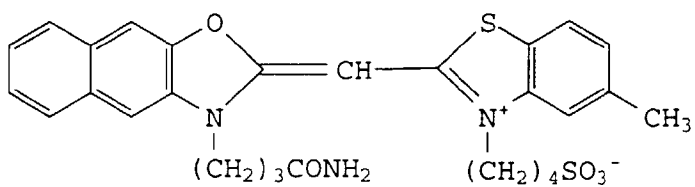
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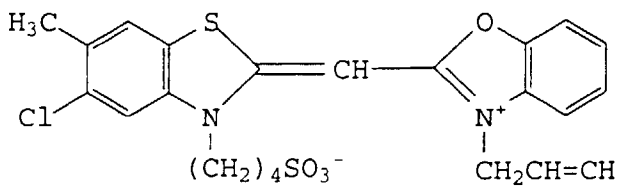
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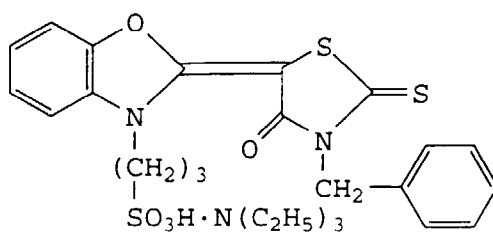
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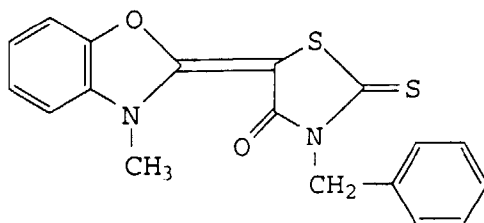


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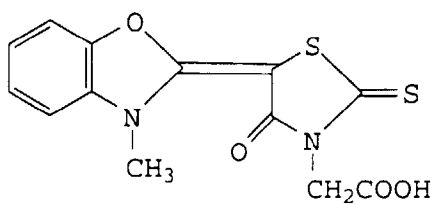


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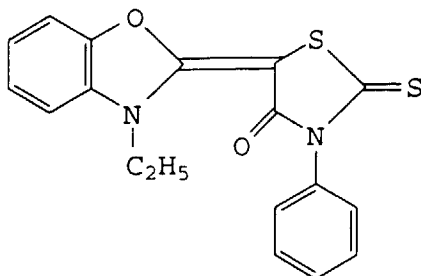


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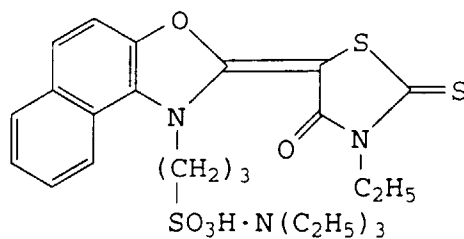


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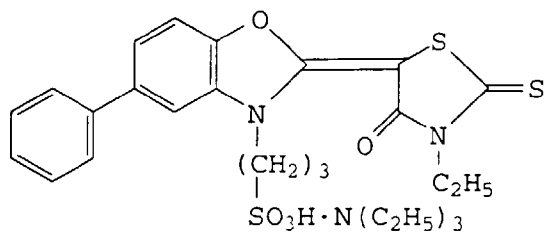


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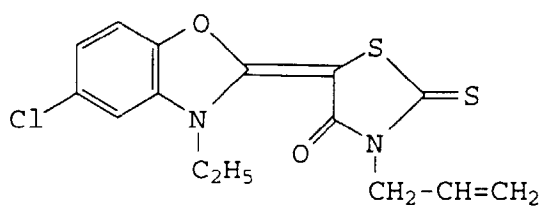


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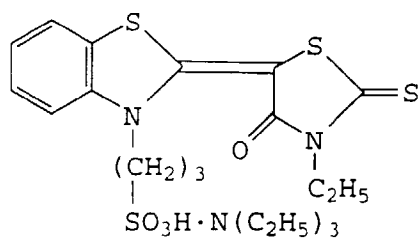


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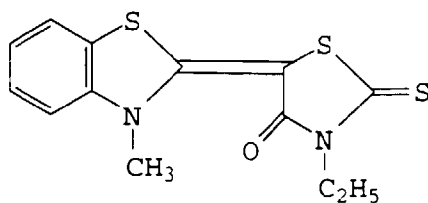


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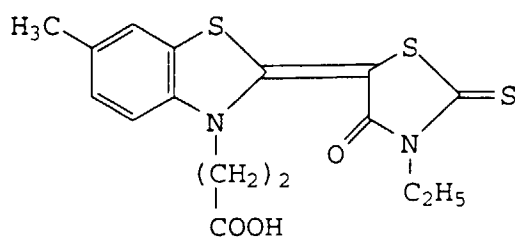
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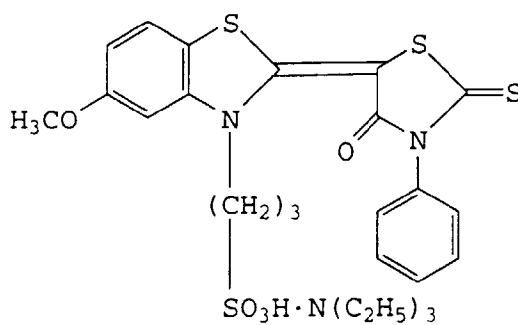


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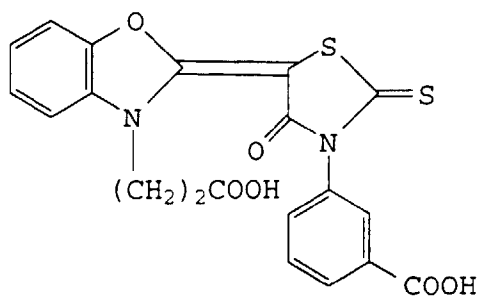


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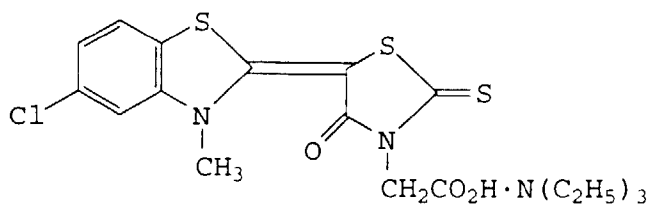


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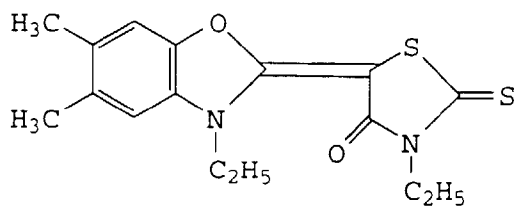
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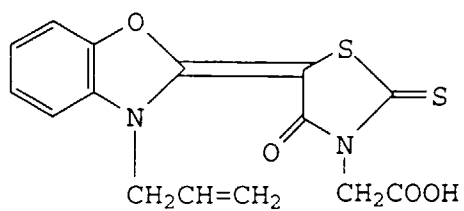


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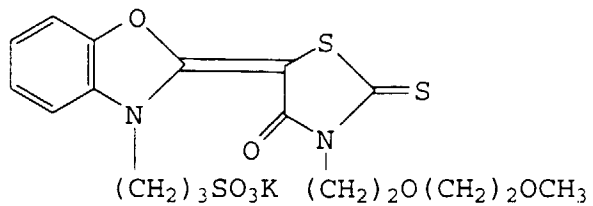
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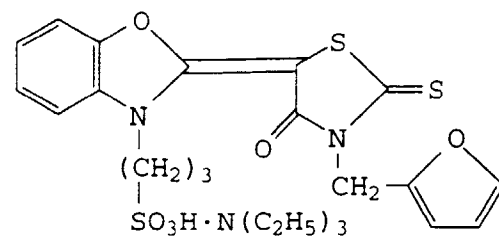


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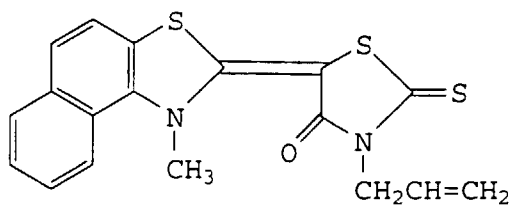
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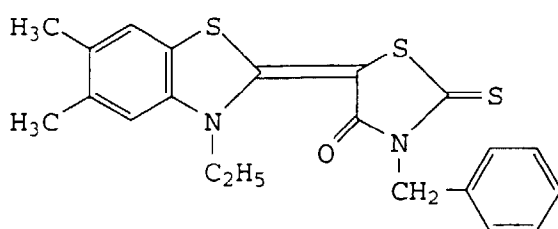


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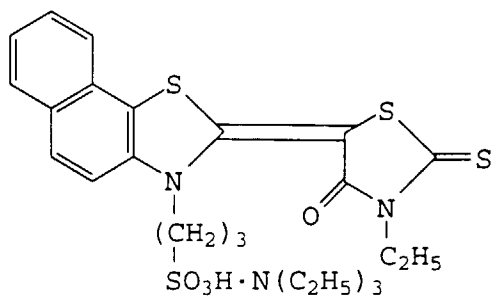


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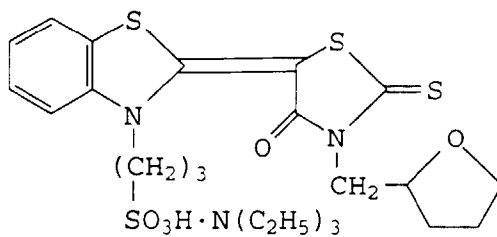


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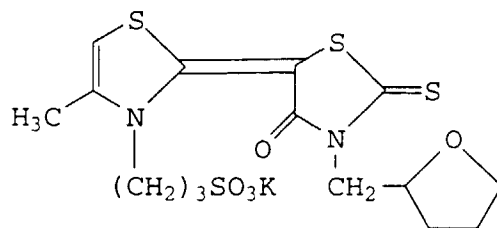
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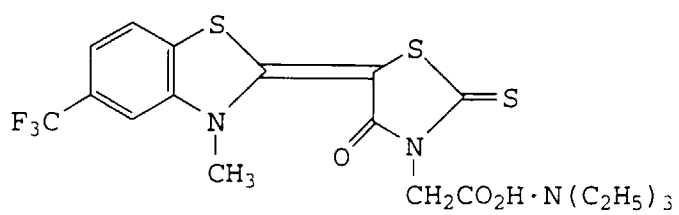


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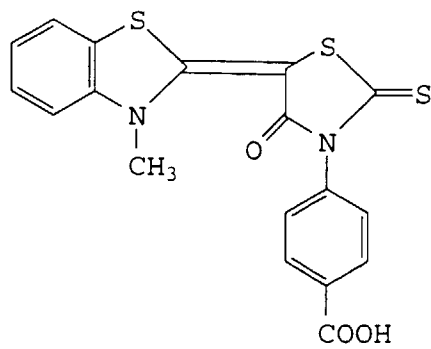


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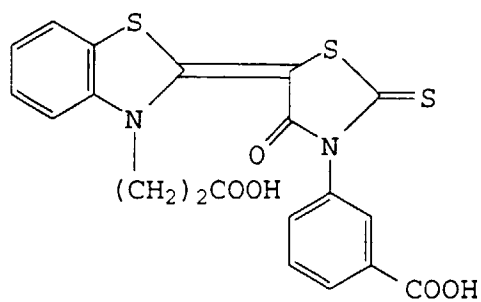
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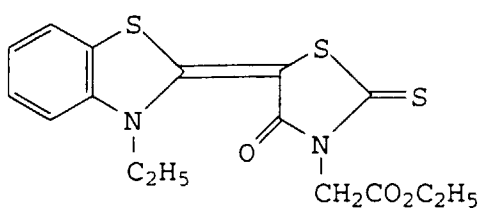


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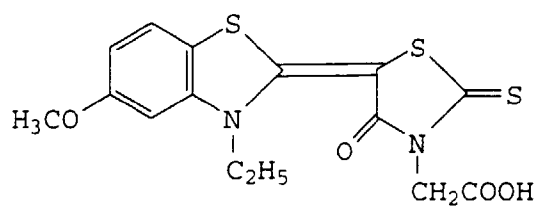
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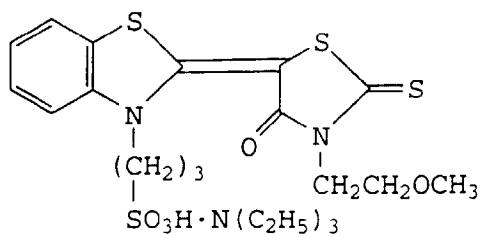


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II-28

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The sensitizing dyes represented by Formulas (I) and (II) have been well-known and they can readily be available. They can also readily be synthesized according to the descriptions of the following literature.

F.M. Hamer, "The chemistry of heterocyclic compounds; The cyanine dyes and related compounds", 1964, John Wiley & Sons, (New York & London), p. 58 & p. 536.

The sensitizing dyes relating to the invention, which are represented by Formula (I) or (II), may be used independently or in combination.

5 The dye represented by Formula (I) or (II) may be added in an amount preferably within the range of 5×10^{-6} to 5×10^{-3} mols per mol of a silver halide content of a silver halide emulsion layer comprising tabular grains.

The dye represented by Formula (I) or (II) may be added in any stage in the course of preparing a silver halide photographic light-sensitive material. It is, however, preferable to add them at any point of time between after the completion of a grain formation and before the completion of a chemical sensitization, and it is more preferable to add
10 them at any point of time between the point of time when completing the grain formation and the point of time before starting the chemical sensitization.

The sensitizing dyes relating to the invention, which are represented by Formulas (I) or (II), are preferably added in the form of a solution in which the dyes are dissolved in a suitable organic solvent such as methanol, phenoxyethanol and phenylethanol, or in a mixed solvent of these organic solvent and water or an amine.

15 The above-mentioned tabular silver halide grains are to have an average grain-diameter within the range of, preferably, 0.3 to 3.0 μm and, particularly, 0.5 to 1.5 μm .

The tabular silver halide grains relating to the invention are to have a mean ratio of grain-diameter/grain-thickness (hereinafter referred to as an average aspect ratio) of not lower than 1.2, and within the range of, preferably 1.5 to 20.0 and, particularly 2.0 to 10.0.

20 The tabular silver halide grains relating to the invention are to have an average thickness of, preferably not thicker than 1.0 μm , particularly not thicker than 0.5 μm and, further preferably not thicker than 0.3 μm .

The advantages of these tabular silver halide grains are disclosed as that a spectral sensitization efficiency can be improved and that the graininess and image-sharpness of an image can be made higher, in the patent publications including, for example, British Patent No. 2,112,157 and U.S. Patent Nos. 4,439,520, 4,433,048, 4,414,310 and
25 4,434,226. The emulsion thereof can be prepared in the processes detailed therein.

In the invention, the grain-diameter of the tabular silver halide grain is defined as the diameter of a circle having the area equivalent to the projected area of the subject grain, that is determined by observing an electron-microscopic photograph of the grain.

In the invention, the thickness of a tabular silver halide grain is defined as the distances between two parallel
30 principal plane face constituting the tabular silver halide grain.

The thickness of a tabular silver halide grain can be obtained from an electron-microscopic photograph of a subject silver halide grain with the shade thereof, or from an electron-microscopic photograph of a section of a subject sample coated with a silver halide emulsion on a support and then dried up.

For determining an average aspect ratio, a series of 100 samples at minimum are to be measured.

35 In a silver halide emulsion of the invention, the proportion of the tabular silver halide grains to the whole silver halide grain is not less than 50% by projected area, preferably not less than 60% and, particularly not less than 70%.

A tabular silver halide grain relating to the invention is preferable to be of the monodisperse type. The term, 'monodisperse type', herein means that the variation coefficient of a grain-size, i.e., (a standard deviation of a grain-size / an average grain-size x 100), is not more than 25%, preferably not more than 20% and, particularly not more than 15%.

40 A silver halide emulsion relating to the invention may have any halide compositions such as silver chloride, silver bromide, silver iodide, silver chlorobromide, silver iodobromide, and silver chloriodobromide. However, from the viewpoint of a high sensitivity, silver iodobromide is preferred. An average silver iodide content thereof is to be within the range of 0.1 to 4.0 mol% and, preferably 0.5 to 3.0 mol%. Further from the viewpoint of a rapid processability, silver chloriodobromide may also be used.

45 When making use of silver chloriodobromide, silver chloride may be contained therein at any positions within the grains. It is particularly preferable to localize silver chloride on the surface of a grain or the neighborhood thereof.

A tabular silver halide emulsion relating to the invention may have a uniform halogen composition in the grains thereof, or may also contain localized silver iodide therein. However, those containing silver iodide localized around each center of the grains thereof may preferably be used. A tabular silver halide emulsion may also be prepared with
50 reference to such a process as described in, for example, JP OPI Publication Nos. 58-113926/1983, 58-113927/1983, 58-113934/1983 and 62-1855/1987 and European Patent Nos. 219,849 and 219,850. The processes for preparing a monodisperse type tabular silver halide emulsion may be referred to that detailed in JP OPI Publication No. 61-6643/1986.

55 A tabular silver iodobromide emulsion having a high aspect ratio can be prepared in such a manner that an aqueous silver nitrate solution is added to an aqueous gelatin solution having a pBr being kept to be not higher than 3.0 so as to produce twinned seed crystals, or both of an aqueous silver nitrate solution and an aqueous halide solution are added at the same time so as to produce twinned seed crystals, and the seed crystals are then grown up in a double-jet method. The sizes of a tabular silver halide grains may be controlled by a temperature at the time of forming the

grains, an adding speed a silver salt and an aqueous halide solution, and so forth.

In a silver halide emulsion relating to the invention, an average silver iodide content or an average silver chloride content thereof may be controlled by changing the composition of an aqueous halide solution to be added, that is a ratio of a bromide, iodide or chloride content thereof. When preparing a silver halide emulsion of the invention, a silver halide solvent such as ammonia, thioether thiocyanate and thiourea may further be used, if required.

For removing a soluble salt from an emulsion (or for carrying out a desalting step), the emulsion may be subjected to such a washing process as a noodle-washing process and a flocculation-precipitation process. Among the preferable washing/desalting processes, the particularly preferable desalting processes include, for example, a process in which an aromatic hydrocarbon type aldehyde resin containing a sulfo group is used, such as described in JP Examined Publication No. 35-16086/1960 and another process in which a macromolecular type flocculant is used, such as the exemplified compounds G3 and G8 given in JP OPI Publication No. 63-158644/1988.

A silver halide emulsion relating to the invention is selenium-sensitized. The expression, "be selenium-sensitized", means that a tabular silver halide grain relating to the invention has a selenium-sensitization nucleus in any position of the grain. Thus, in the present invention, selenium-sensitization is carried out at a time during a period from before completing grain growth to before completing chemical sensitization. The preferable position thereof is the surface of the grain and/or the neighborhood of the surface.

In the invention, the expression, "the neighborhood of the surface", means an internal in a depth of a position not more than 600nm, preferably not more than 200nm and, more preferably not more than 70nm, from the surface of the grain.

As for a process for selenium-sensitizing an objective position in the neighborhood of a grain of the invention, it is preferable to carry out such a process that a grain-growing step is temporarily suspended in operation and a selenium sensitizer is added thereto so that the shell-layer surface may be selenium-sensitized, and that the grain growing step is then resumed to continue the operation.

During the period of silver halide grain growth, a selenium-sensitizer may be added either by splitting it into several parts or continuously for a certain period of time.

In a selenium-sensitizing nucleus closest to the surface of a grain to be finally formed, the depth of the nucleus may be varied by applying a selenium-sensitizing process while carrying out a grain-growing step and then by controlling an amount of silver to be added to a grain-regrowing step to be carried out thereafter.

In the present invention, selenium sensitization is performed at a time during a period from a time at which 70% or more, preferably 80% or more, more preferably 90% or more, of the whole silver salt has been added and before the addition of the silver salt is completed.

A selenium-sensitization relating to the invention can be performed in any conventionally known process. To be concrete, such a selenium-sensitization may usually be carried out by adding an labile type selenium compound and/or a non-labile type selenium compound and then by stirring an objective emulsion at a high temperature preferably at not lower than 40°C for a specific period. It is preferable to use a selenium-sensitization in which an labile type selenium sensitizer is used, such as detailed in JP Examined Publication No. 44-15748/1969. The typical unstable type selenium-sensitizers include, for example, an aliphatic isoselenocyanate such as allylisoselenocyanate; a selenourea; a selenoketone; a selenoamide; a selenocarboxylic acid and the esters thereof; and a selenophosphate. Among them, some of the preferable labile type selenium compounds will be given below.

- (1) A colloidal elemental selenium;
- (2) An organic selenium compound (that is, a compound having a selenium atom made double-bonded to a carbon atom of an organic compound, by a covalent bond);

- a) An isoselenocyanate including, for example, an aliphatic isoselenocyanate such as allylisoselenocyanate;
- b) A selenourea (including those in an enol form) including, for example, a selenourea; an aliphatic selenourea such as methyl, ethyl, propyl, isopropyl, butyl, hexyl, octyl, dioctyl, tetramethyl, N-(β -carboxyethyl)-N,N'-diethyl, N,N-dimethyl, diethyl and dimethyl; an aromatic selenourea having one or more aromatic groups, such as phenyl and tolyl; and a heterocyclic selenourea having a heterocyclic group, such as pyridyl and benzothiazolyl;

Among those selenoureas, N,N'-tetra-substituted selenourea is particularly preferable. The preferable substituents include, for example, R, RCO- and ArCO-, in which R represents an alkyl group or a perfluoroalkyl group (preferably having C of 1 to 7); and Ar represents a halogen or a phenyl group substitutable with a lower alkoxy group.

- c. A selenoketone including, for example, selenoacetone, selenoacetophenone, selenoketone having an alkyl group made bonded to =C=Se, and selenobenzophenone;
- d. A selenoamide including selenoamide; and

e. Selenocarboxylic acid and the esters thereof including, for example, 2-selenopropionic acid, 3-selenobutyric acid, and methyl-3-selenobutyrate;

(3) Others:

- a. A selenide including, for example, diethyl selenide, diethyl diselenide and triphenylphosphine selenide; and
 b. A selenophosphate including, for example, tri-p-triselenophosphate and tri-n-butylselenophosphate.

A preferable series of the labile selenium compounds are given above. However, these compounds shall not be limited thereto. Speaking of an labile selenium compound for serving as a sensitizer for a photographic emulsion, the structure thereof may not be particularly essential for the skilled ones in the art, as far as a selenium is unstable, but it has generally been recognized that the structure thereof has not any role, except that an organic portion of a selenium-sensitizer molecule carries the selenium and make it present in the unstable form in an emulsion. In the invention, an labile selenium compound having such a broad concept as mentioned above can advantageously be used.

It is also allowed to use a selenium-sensitization using a non-labile selenium sensitizer therein, such as those detailed in JP Examined Publication Nos. 46-4553/1971, 52-34492/1977 and 52-34491/1977. The non-labile selenium compounds include, for example, selenious acid, potassium selenocyanide, a selenazole, a quaternary ammonium salt of a selenazole, diaryl selenide, diaryl diselenide, 2-thioselenazolidine dione, 2-selenoxazolidine dione, and the derivatives thereof.

A non-labile selenium sensitizer and a thioselenazolidine dione compound each detailed in JP Examined Publication No. 52-38408/1977 are also effective.

Such a selenium sensitizer as mentioned above may be dissolved in water, an organic solvent such as methanol and ethanol independently or a mixed solvent thereof, and the mixture thereof is then added in the course of carrying out a chemical sensitization. Such a selenium sensitizer as mentioned above shall not be limited to a single kind thereof, but two or more kinds thereof may also be used in combination. It is also preferable to make use of an labile selenium compound and a non-labile selenium compound in combination.

An amount of a selenium sensitizer applicable to the invention to be added may be so varied as to meet the activities thereof, the kinds and sizes of silver halides and the temperatures and times for ripening the emulsion. However, it is added in an amount of, preferably, not less than 1×10^{-8} mols per mol of silver halide used and, more preferably, within the range of not less than 1×10^{-7} mols to not more than 5×10^{-5} mols.

A selenium sensitization can become more effective when it is carried out in the presence of a silver halide solvent.

The silver halide solvents applicable to the invention include, for example, (1) such an organic thioether as mentioned in U.S. Patent Nos. 3,271,157, 3,531,289 and 3,574,628 and JP OPI Publication Nos. 54-1019/1979 and 54-158917/1979; (2) such a thiourea derivative as mentioned in JP OPI Publication Nos. 53-82408/1978, 55-77737/1980 and 55-2982/1980; (3) such a silver halide solvent having a thiocarbonyl group sandwiched between an oxygen or sulfur atom and a nitrogen atom as mentioned in JP OPI Publication No. 53-144319/1978; (4) such an imidazole as mentioned in JP OPI Publication No. 54-100717/1979; (5) a sulfite; and (6) a thiocyanate.

The particularly preferable solvents include, for example, a thiocyanate and a tetramethyl thiourea. An amount of a solvent applicable thereto may be so varied as to meet the kinds thereof. However, in the case of a thiocyanate, an preferable amount thereof to be applied is to be within the range of 1×10^{-4} mols to 1×10^{-2} mols per mol of silver halide used.

A reduction-sensitization may also be applied to the interior of the grains of a silver halide emulsion of the invention. Such a reduction-sensitization may be applied to a silver halide emulsion in a process in which a reducing compound is added; another process that is so-called a silver ripening process for passing the emulsion grains through a state of excessive silver ions having a pAg of 1 to 7; a further process that is so-called a high-pH ripening process for passing the emulsion grains through the state of a high-pH having a pH of 8 to 11; or the like. The above-mentioned processes may also be applied in combination.

The a reducing compound-adding process is preferable from such a viewpoint that the reduction-sensitization degrees can delicately be controlled.

As for the foregoing reducing compounds, either of an inorganic or organic compound may be used. They include, for example, thiourea dioxide, a stannous salt, an amine or polyamine, a hydrazine derivative, a formamidine sulfonic acid, a silane compound, a borane compound, ascorbic acid and the derivatives thereof, and a sulfite. Among them, thiourea dioxide, stannous chloride and dimethylamine borane are particularly preferable. The amounts of those reducing compound to be added are varied according to the reductivity thereof, the kinds of silver halides and the emulsion preparation conditions such as dissolving conditions. However, they may be suitably added in an amount within the range of 1×10^{-8} to 1×10^{-2} mols per mol of silver halide used. It is preferable to dissolve the above-mentioned reducing compounds in water or an organic solvent such as an alcohol and then to add while growing silver halide grains.

After completing the step of a desalting process, it is preferable that a silver halide emulsion of the invention is

chemically ripened in a selenium sensitization process and in the other chemical sensitization than the selenium sensitization. Thus, a silver halide grain emulsion of the invention is a surface-sensitive emulsion, i.e., emulsion that forms latent images primarily on the surfaces of the silver halide grains. Therefore, an internal latent image-forming silver halide grain emulsion that forms latent images predominantly in the interior of the silver halide grains is outside the scope of the present invention.

The temperatures for carrying out a chemical ripening process may freely be selected. It is, however, within the range of 20°C to 80°C, preferably 30°C to 70°C and, particularly 35°C to 65°C.

The chemical sensitization processes other than the above-mentioned selenium sensitization process include, for example, a chalcogen sensitization process and such a noble metal sensitization process as a gold sensitization process. It is preferable to carry out a sulfur sensitization process and a gold sensitization process in combination.

For carrying out a sulfur sensitization process, such a sensitizer as a thiosulfate, thiourea allylthiocarbamide, cystine, p-toluene thiosulfate and rhodanine may be used. Besides the above, it is also allowed to use a sulfur sensitizer such as those described in, for example, U.S. Patent Nos. 1,574,944 and 3,656,955; German Patent No. 1,422,869; JP Examined Publication No. 56-24937/1981; and JP OPI Publication No. 55-45016/1980. Such a sulfur sensitizer as given above may be added in an amount sufficient to enhance an emulsion sensitivity effectively. The amount thereof to be added can widely be varied under various conditions such as those of pH values, temperatures and silver halide grain sizes. However, a rough standard thereof is preferably within the range of 5×10^{-8} to 5×10^{-5} mols per mol of silver contained in a silver halide emulsion relating to the invention.

For carrying out a gold sensitization process, the following gold sensitizers may be used. Namely, a chloroaurate, an gold-thiourea complex salt, potassium chloroaurate, auric trichloride, potassium auric thiocyanate, potassium iodoaurate, tetracyanoauric amide, ammonium aurothiocyanate and pyridyl trichlorogold. The amounts of those gold sensitizers to be added are widely varied under various conditions. It may be added in an amount within the range of, preferably, 5×10^{-7} to 5×10^{-3} mols and, more preferably, 2×10^{-6} to 4×10^{-4} mols, each per mol of a silver halide emulsion of the invention.

To a silver halide emulsion of the invention, it is preferable to add a finely grained silver halide emulsion, after completing the grain formation.

The above-mentioned fine grains of silver halides include, for example, silver chloride, silver bromide, silver iodide, silver chlorobromide, silver iodochloride, silver iodobromide and silver chloriodobromide. Among them, silver bromide and silver iodide are preferred. The grain size (diameter) of each of those silver halide fine grains is not larger than 0.12 μ m, preferably within the range of 0.001 to 0.10 μ m and, more preferably 0.006 to 0.06 μ m.

In the case of applying silver iodide fine grains to the invention, γ type AgI in a cubic crystal form and β type AgI in a hexagonal crystal form have generally been known as silver iodide. However, they may have either one of the crystal structures and they may also be of the mixture thereof.

In the invention, when making use of the fine grains of silver bromide, silver chloride or a silver chlorobromide, those fine grains are each preferable to be of a non-twinned crystal substantially having no twinned plane, that is so-called a normal crystal or a single twinned crystal having one twinned plane.

Silver halide fine grains applicable to the invention are preferable to be excellent in monodispersibility and to be prepared in a double-jet precipitation process while controlling the temperature, pH and pAg thereof.

When the average grain size of silver halide grains relating to the invention is regarded as d (μ m), silver halide fine grains may be added in an amount of preferably not more than $1/100d$ mols, and within the range of, more preferably $1/2000d$ to $1/300d$ mols and, most preferably $1/5000d$ to $1/500d$ mols, each per mol of a silver content of an emulsion used in the invention.

Silver halide fine grains may be added at any point of time in the course from the point of time when carrying out a chemical ripening step to the point of time immediately before starting a coating step, and they may preferably be added in the course of carrying out the chemical ripening step. The term, 'a chemical ripening step', herein means a period of time between the point of time when completing the formation of the grains of an emulsion relating to the invention and when completing a desalting treatment, and the point of time when completing chemical ripening. The methods for completing a chemical ripening step include, for example, a temperature lowering method, a pH lowering method and a method of making use of a chemical ripening stopper, each of which have been known. Taking an emulsion stability into consideration, the method of making use of a chemical ripening stopper is preferable. Those chemical ripening stoppers include, for example, a halide (such as potassium bromide and sodium chloride) and an organic compound having been known as an antifoggant or a stabilizer (such as 4-hydroxy-6-methyl-1,3,3a,7-tetraza-indene). They may be used independently or in combination.

Silver halide fine grains may be added in several parts with some time intervals, or a further chemical ripened emulsion may also be added after adding the silver halide fine grains.

When adding silver halide fine grains, the temperature of an emulsion solution relating to the invention is to be within the range of, preferably 30 to 80°C and, particularly 40 to 65°C.

The invention is preferably embodied preferably under the conditions where a part of or the whole of silver halide

fine grains are vanished in the course between the time after adding the grains and the time immediately before carrying out a coating operation and, more preferably, under the conditions where not less than 20% of the silver halide fine grains added therein are to be vanished immediately before carrying out a coating operation.

The quantitative determination of a vanished amount thereof can be accomplished in the following manner; an emulsion or a coating solution, to which silver halide fine grains were added, is centrifuged under suitable conditions and then the absorption spectra of a supernatant are measured. Thereafter, the resulting absorption spectra are compared to the absorption spectra of silver halide grains having an already known density so that the quantitative determination can be obtained.

Silver halide grains of the invention can contain a metal belonging to VIII group of the periodic table.

A metal belonging to VIII group of the periodic table may be added so as to be contained inside of silver halide grains usually in such a manner that the metal is made present as the metal compound thereof when forming the grains, and it may also be added continuously or in several parts.

It is also preferable to carry out a process in which the above-mentioned metal is added in advance to an aqueous silver salt solution and/or an aqueous halide solution and silver halide grains are precipitated by making use of the aqueous solution(s).

A compound of a metal belonging to the VIII group of the periodic table herein means a metal compound of iron, iridium, platinum, palladium, nickel, rhodium, osmium, ruthenium and cobalt. It is the matter of course that not only the above-mentioned metal compounds, but also the metal ions and metal atoms thereof may also be contained in silver halide grains relating to the invention.

A silver halide emulsion layer and non-light-sensitive hydrophilic colloidal layer each relating to the invention are hardened so that the dissolving time thereof (that is the melting time thereof) can be within the range of 5 to 150 minutes, preferably 7 to 90 minutes and, particularly 10 to 60 minutes.

The above-mentioned dissolving time can be confirmed in the following manner, for example; a sample cut into a 5mm x 20mm sized piece is dipped in an aqueous 1.5 wt% sodium hydroxide solution being kept at 50°C so as to be in a non-stirring state and then a period of time until a hydrophilic colloidal layer is eluted out is measured.

A desired dissolving time (or a desired melting time) can be obtained in such a means that the control is made with a layer hardener. For this purpose, any one of the conventionally known layer hardeners may be used independently or in combination.

For example, the following compounds may be used; namely, a chromium salt (such as chromium alum and chromium acetate), an aldehyde (such as formaldehyde, glyoxal and glutar aldehyde), an N-methylol compound (such as diethylol urea and methylol dimethyl hydantoin), a dioxane derivative (such as 2,3-dihydroxy dioxane), an active vinyl compound (such as 1,3,5-triacryloyl-hexahydro-2-triazine and 1,3-vinylsulfonyl-2-propanol), an active halogen compound (such as 2,4-dichloro-6-hydroxy-3-triazine), a mucohalogeno acid (such as mucochromic acid and mucophe-noxychromic acid), an isoxazole, dialdehyde starch, 2-chloro-6-hydroxy triazino gelatin, and a carbamoyl pyridinium compound.

It is preferable that a layer can be hardened by making use of such a hardener as mentioned above so that a swelling rate in water can be not higher than 200%, preferably not higher than 160% and, particularly not higher than 120%.

In an emulsion relating to the invention, a variety of photographic additives can be used in the steps before or after carrying out a physical or chemical ripening step. The well-known additives include, for example, the compounds given in Research Disclosure (RD) No. 71643 (Dec., 1978), *ibid.*, No. 18716 (Nov., 1979) and, *ibid.*, No. 308119 (Dec., 1989). The kinds of the compounds and the pages of the RDs where the compounds, each given in the above-mentioned three issues thereof, will be indicated below.

Additive	RD-17643		RD-18716		RD-308119	
	Page	Class	Page	Class	Page	Class
Chemical sensitizer	23	III	648 Upper R.996		996	III
Sensitizing dye	23	IV	648-649		996-8	IV
Desensitizing dye	23	IV			998	IVB
Dye	25-26	VIII	649-650		1003	VIII
Development accelerator	29	XXI	648 Upper R.			
Antifoggant-stabilizer	24	IV	649 Upper R.		1006-7	VI
Whitening agent	24	V			998	VI
Hardener	26	X	651 L.		1004-5	X
Surfactant	26-7	XI	650 R		1005-6	XI

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(continued)

Additive	RD-17643		RD-18716		RD-308119	
	Page	Class	Page	Class	Page	Class
Antistatic agent	27	XII	650 R		1006-7	XIII
Plasticizer	27	XII	650 R		1006	XII
Lubricant	27	XII				
Matting agent	28	XVI	650 R		1008-9	XVI
Binder	26	XXII			1003-4	IX
Support	28	XVII			1009	XVII

The supports applicable to a light-sensitive material relating to the invention include, for example, those described in the foregoing RD-17643, p.28 and, *ibid.*, RD-308119, p.1009.

A suitable support include those made of a plastic film. To such a support as mentioned above, an under-coat layer may also be provided and/or a corona-discharge or UV-irradiation may further be treated, so as to improve a coated layer to readily adhere to the surface of the support.

EXAMPLES

Now, the examples of the invention will be detailed below. However, the invention shall not be limited to the examples given below.

Example 1

Preparation of Tabular Silver Halide Emulsion

A seed emulsion comprising silver iodobromide was prepared by making use of the following solutions.

[A ₁]	
Hydrogen peroxide-treated ossein gelatin	40g
Potassium bromide	75.1g
Add water to make	4000ml
[B ₁]	
Silver nitrate	600g
Add water to make	803ml
[C ₁]	
Hydrogen peroxide-treated ossein gelatin	16.1g
Potassium bromide	393.7g
Potassium iodide	35.1g
Add water to make	803ml
[D ₁]	
Aqueous ammonia (in a 28% solution)	235ml

An apparatus disclosed in JP OPI Publication No. 62-160128/1987 was used, and six units each of nozzles for supplying Solutions B₁ and C₁ were provided to the lower part of each stirring propeller for mixing the solutions, respectively.

Solutions B₁ and C₁ were each added, at a flow rate of 62.8 ml/min in a controlled double-jet method, to Solution A₁ stirred at a high speed of 430 rpm and at 40°C. From 4 min. 46 sec. after starting the addition thereof, the flow rate was gradually increased so as to be the final flow rate of 105 ml/min. The whole adding time was 10 min. 45 sec. While adding Solutions B₁ and C₁, the pBr of the solution being prepared was kept at 1.3 with a (3.5N) potassium bromide solution.

After completing the addition, the temperature of the resulting mixed solution was lowered linearly to 20°C by

taking 10 minutes, and Solution D₁ was added thereto with stirring at 460 rpm by taking 20 seconds. The resulting solution was subjected to an Ostwald ripening treatment for 5 minutes. In the course of the ripening treatment, the bromine concentration, ammonia concentration and pH of the resulting ripened solution were 0.025 mol/liter, 0.63 mols/liter and 11.7, respectively.

Immediately thereafter, acetic acid was added thereto and neutralized so that the mixture thereof could have a pH of 5.6, and the ripening treatment was then stopped in operation. For removing the excessive salts, a desalting treatment was carried out by making use of an aqueous solution of Demol N (manufactured by Kao-Atlas Corp.) and an aqueous solution of magnesium sulfate, so that seed emulsion Em-1 could be prepared.

When observing Em-1 through an electron microscope, it was proved to be comprised of globe-shaped grains having an average grain-size of 0.28 μ m and a grain-size variation coefficient of 21%.

Growth of Seed Grain Emulsion

Successively, there was prepared a silver halide grain emulsion, which is mainly comprised of tabular twinned crystals, by making use of comparative seed emulsion Em-1 and the following three kinds of solutions.

Preparation of Emulsion

The resulting seed emulsion was taken in an amount equivalent to 0.027 mols per mol of silver halide of a final emulsion. It is then so dissolved as to be dispersed in an aqueous gelatin solution having a temperature of 62°C and containing a copolymer of polypropylene oxide (or PO) and polyethylene oxide (or EO), (having an EO/PO ratio of 0.33 and a molecular weight of 1400), that was served as a defoamer. In succession, an aqueous 3.5N silver nitrate solution and an aqueous 3.4N potassium bromide solution were each minutes added by a controlled double-jet method over a period of 97 at an accelerated adding rate, so that the adding rate at the time when completing the addition could be 2.3 times as fast as at the time when starting the addition. In the whole course of adding the solutions, the temperature, pH and pAg thereof were constantly kept at 62°C, 5.8 and 8.9, respectively. Further, the resulting mixture was desalted in the same manner as in the case of the seed emulsion preparation. The resulting emulsion was proved to be a tabular silver halide grain emulsion having an average grain-size of 1.30 μ m, an average thickness of 0.32 μ m and an average aspect ratio of 4.1. The resulting emulsion is hereinafter referred to as EM-1.

Preparation of Silver Iodide Fine Grains

To 5000ml of a 5.2 wt% gelatin solution containing 0.008 mols of potassium iodide, 1500ml of an aqueous solution containing 1.06 mols each of silver nitrate and potassium iodide was added at a constant adding rate by taking 35 minutes. The temperature in the course of preparing fine grains was kept at 40°C. When confirming the resulting silver iodide grains through a 60000X electron microscope, the grains were each proved to be a mixture of β -AgI and γ -AgI having an average grain-size of 0.05 μ m.

Chemical Sensitization of Emulsion

While keeping the resulting EM-1 stirred at 52°C, the foregoing sensitizing dyes relating to the invention, I-6, I-7, I-16, II-1, II-3 and II-13, were added respectively to separate methanol solutions as shown in Table 1, so that the amount of each sensitizing dye added could be 140mg per mol of a silver content of EM-1. 10 minutes thereafter, 60mg of ammonium thiosulfate, 1.45mg of chloroauric acid and 13.8mg of sodium thiosulfate, each per mole of silver, were added thereto as the chemical sensitizers. 30 minutes thereafter, the foregoing silver iodide finely grained emulsion was further added in an equivalent amount of 1.37×10^{-3} mols and, after passing a specific period of time, 4-hydroxy-6-methyl-1,3,3a,7-tetrazaindene and 1-phenyl-5-mercaptotetrazole were each so added thereto as to be stabilized. Therefore, the resulting emulsions were chemically ripened at optimum, respectively. The resulting emulsions are hereinafter referred to as EM-2 through EM-7, respectively.

Besides the above, a selenium-sensitized emulsions EM-8 through EM-13 were each prepared in quite the same manner as in EM-2 through EM-7, except that N,N-dimethyl selenourea was further added in an amount of 8×10^{-7} together with chloroauric acid, as the chemical sensitizers.

Preparation of Internally Selenium-Sensitized Emulsion

The addition of the aqueous silver nitrate solution and halide solution of Emulsion EM-1 were temporarily stopped at the time when 80% of silver nitrate solution was added, and ammonium thiocyanate and N,N-dimethyl selenourea were then added thereto so as to be in the amounts thereof of 7.8×10^{-6} mols and 1.1×10^{-6} mols each per mol of the

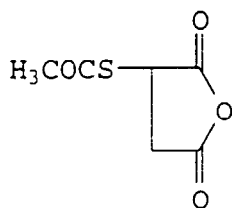
silver used in the emulsion to be finally prepared, respectively. Four (4) minutes thereafter, the remaining aqueous silver nitrate solution and halide solution were further added. When observing the resulting emulsion through an electron microscope, the configuration of the resulting emulsion was proved to be approximately the same as that of EM-1. The resulting emulsion is hereinafter referred to as EM-14.

The sensitizing dyes relating to the invention same as those used in EM-2 through EM-7 were added respectively to each of EM-14, and the same gold-sulfur sensitizers as those used in EM-2 through EM-7 were further added thereto, respectively. The resulting emulsions are hereinafter referred to as EM-15 through EM-20, respectively.

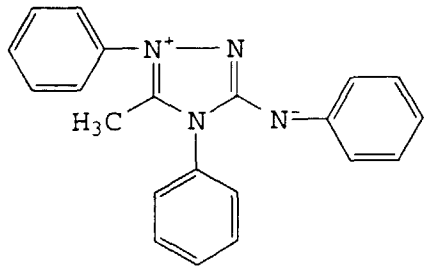
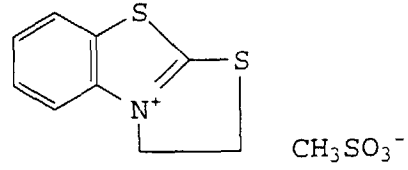
Emulsion-layer coating solutions were prepared, by adding the following additives to the resulting chemically ripened emulsions EM-1 through EM-7, EM-8 through EM-13 and EM-15 through EM-20, respectively. At the time, the following protective-layer coating solution was also prepared. The amounts of the resulting coating solutions to be coated were determined so that the silver contents and gelatin contents of the solutions could be in the amounts of 2.5 g/m² and 1.85 g/m² each per one coated surface, respectively, and each support was simultaneously coated on both-side thereof at a coating speed of 80m per minute by making use of two units of slide-hopper type coating machine. Each of the resulting emulsion-layers was dried up for 2min.20sec., so that the coated samples No. 1 through No. 18 were prepared. As for the support, there used a 175μm-thick and 0.15 density blue-colored polyethylene terephthalate film base for X-ray film use, which was prepared in the following manner; a copolymeric aqueous dispersed solution for serving as a subbing solution was prepared by diluting it so as to have a concentration of a 10 wt% copolymer comprising three monomers, glycidyl methacrylate of 50 wt%, methyl acrylate of 10 wt% and butyl methacrylate of 40 wt%, and the resulting subbing solution was then subbed on the above-mentioned film base.

The following additives were used in the emulsion. The amounts thereof used are each indicated by an amount per mol of silver halide.

1,1-dimethylol-1-bromo-1-nitromethane	70mg
t-butyl-catechol	82mg
Polyvinyl pyrrolidone (having a molecular weight of 10,000)	1.0g
Styrene-maleic anhydride copolymer	25g
Nitrophenyl-triphenyl phosphonium chloride	50mg
Ammonium 1,3-dihydroxybenzene-4-sulfonate	2.0 g
Sodium 2-mercaptobenzimidazole-5-sulfonate	1.5mg

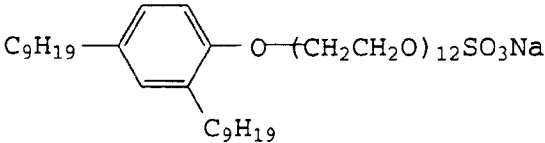
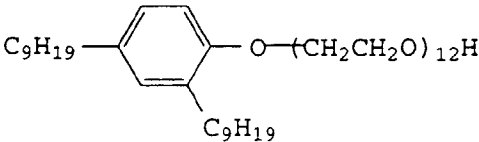
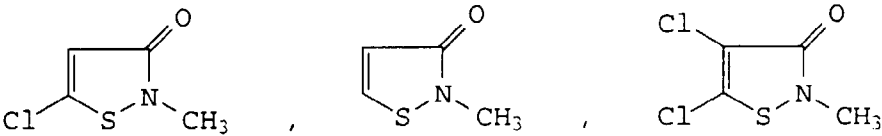


7.2 mg

5		
10		150 mg
15		
20		80 mg
25		1g
30		15mg
35		7g
40		600mg
45		2.5g

Next, the following solution was prepared for a protective layer coating solution. The amounts of the additives added thereto are each indicated by an amount per liter of the coating solution.

35		68g
40		2g
45		0.3g
50		1.1g
55		0.5g

5	Rudox AM (colloidal silica manufactured by DuPont)	230g
	(CH ₂ =CHSO ₂ CH ₂) ₂ O, (a layer hardener)	200mg
10	An aqueous 40% glyoxal solution, (a layer hardener)	2.0ml
15		1.0 g
20		0.4 g
25		
30	(50:46:4)	0.1 g

Sensitometry (Evaluation of Photographic Characteristics)

35 Sensitometry was carried out in the following manner.

A subject sample was held between two sheets of intensifying screens (manufactured by Konica Corp.) and, thereto X-rays were irradiated with a tube-voltage of 80 kvp and a tube-current of 50mA, for 0.05 seconds through an aluminum-made wedge. Thereafter, a series of developing and fixing treatments thereof were carried out by making use of a roller-transport type automatic processor and the following developer and fixer. The details thereof are as follows.

40

45

50

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Composition of Developer

5	Part A (for a 12-liter preparation)	
	Potassium hydroxide	450g
10	Potassium sulfite (in a 50% solution)	2280g
	Diethylenetetramine pentacetate	120g
	Sodium hydrogen bicarbonate	132g
15	5-methyl benzotriazole	1.2g
	1-phenyl-5-mercaptotetrazole	0.2g
20	Hydroquinone	340g
	Add water to make	5000ml
	Part-B (for a 12-liter preparation)	
25	Glacial acetic acid	170g
	Triethylene glycol	185g
30	1-phenyl-3-pyrazolidone	22g
	5-nitroindazole	0.4g
	N-acetyl-DL-penicillamine	1.2g
35	Starter	
	Glacial acetic acid	120g
40	Potassium bromide	225g
	Add water to make	1.0 liter
	Composition of Fixer	
45	Part-A (for 18-liter preparation)	
	Ammonium thiosulfate (in 70 wt/vol %)	6000g
50	Sodium sulfite	110g
	Sodium acetate·trihydrate	450g
55	Sodium citrate	50g

Gluconic acid 70g

1- (N,N-dimethylamino) -ethyl
-5-mercaptotetrazole 18g

Part-B

Aluminium sulfate 800g

In the preparation of a developer, Part-A and Part-B were added at the same time to approximately 5 liters of water and water was further added with stirring so as to make 12 liters in total. The pH thereof was controlled to be 10.40 with glacial acetic acid. The resulting solution was used as a developer.

The foregoing starter was added in an amount of 20 ml per liter of the developer, and the pH thereof was controlled to be 10.26, so that the resulting solution could be used.

In the preparation of a fixer, Part-A and Part-B were added at the same time to approximately 5 liters of water and water was further added with stirring so as to make 18 liters in total. The pH thereof was controlled to be 10.4.6 with sulfuric acid and NaOH. The resulting solution was used as a fixer.

The processing treatments were carried out for 90 seconds in a dry-to-dry system. The temperatures in the course of carrying out the processing were 32°C in the developing treatment, 33°C in the fixing treatment and 50°C in the drying treatment. The sensitivities each obtained therefrom were expressed in terms of the reciprocals of an exposure capable of giving a density of fog + 0.5, and indicated by a value relative to the sensitivity obtained from Sample No. 1 that was regarded as the standard value of 100.

Evaluation on Roller-mark occurrence

A pressure mark (so-called a roller-mark) produced by a roller of an automatic processor was evaluated in the following manner. A processing treatment was carried out through an automatic processor under the unexposed conditions and in the same manner as in the above-mentioned sensitometric evaluation. The roller-marks produced therein were each visually evaluated upon classifying them into the following 5 grades.

Grade 5: No roller-mark produced;

Grade 4: Few roller-marks produced;

Grade 3: A few roller-marks produced (within a practical applicability);

Grade 2: Many roller-marks produced (Out of the practical applicability); and

Grade 1: Remarkably numerous roller-marks produced.

Samples No. 1 through No. 18, each cut into a 5mm x 20mm size were each dipped in an aqueous 1.5 wt% sodium hydroxide solution being kept at 50°C. When the resulting meting time of each sample was measured, it was within the range of 15min. to 20min. Therefore, each sample after processed was satisfactory in dryness.

The results thereof will be shown in the following Table 1.

Table 1

Sample No.	Em No.	Sensitizing dye	Chemical sensitization	Position of selenium nuclei of grains	Photographic characteristics			Roller-mark
					Fog	Sensitivity	Dmax	
1 (Comparison)	EM-2	I-6	Gold-Sulfur	-	0.04	100	2.69	2
2 (Comparison)	EM-3	I-7	Gold-Sulfur	-	0.04	92	2.66	2
3 (Comparison)	EM-4	I-16	Gold-Sulfur	-	0.05	97	2.72	2
4 (Comparison)	EM-5	II-1	Gold-Sulfur	-	0.05	103	2.68	1
5 (Comparison)	EM-6	II-3	Gold-Sulfur	-	0.04	97	2.66	1
6 (Comparison)	EM-7	II-13	Gold-Sulfur	-	0.05	100	2.68	2
7 (Comparison)	EM-8	I-6	Gold-Sulfur-Selenium	Surface	0.04	120	3.10	4
8 (Comparison)	EM-9	I-7	Gold-Sulfur-Selenium	Surface	0.04	115	3.13	4
9 (Comparison)	EM-10	I-16	Gold-Sulfur-Selenium	Surface	0.04	117	3.15	4
10 (Comparison)	EM-11	II-1	Gold-Sulfur-Selenium	Surface	0.04	128	3.35	3
11 (Comparison)	EM-12	I-3	Gold-Sulfur-Selenium	Surface	0.04	125	3.28	3
12 (Comparison)	EM-13	II-13	Gold-Sulfur-Selenium	Surface	0.04	119	3.21	4
13 (Invention)	EM-15	I-6	Gold-Sulfur-Selenium	Interior	0.03	143	3.35	5
14 (Invention)	EM-16	I-7	Gold-Sulfur-Selenium	Interior	0.03	140	3.42	5
15 (Invention)	EM-17	I-16	Gold-Sulfur-Selenium	Interior	0.03	152	3.40	5
16 (Invention)	EM-18	II-1	Gold-Sulfur-Selenium	Interior	0.03	160	3.50	5
17 (Invention)	EM-19	II-3	Gold-Sulfur-Selenium	Interior	0.03	157	3.38	5
18 (Invention)	EM-20	II-13	Gold-Sulfur-Selenium	Interior	0.03	155	3.44	5

According to the invention, as is obvious from Table 1, it is shown that a light-sensitive material having the photographic characteristics high in sensitivity, low in fog and excellent in the maximum density. Resistance the roller-mark production can also be so excellent that almost no roller-mark production could be confirmed. Further, the photographic

characteristics and roller-mark production resistance were proved to be more excellent when selenium nuclei are each positioned in the interior than on the grain surfaces.

Example 2

Preparation of Hexagonal Tabular Twinned-Crystal Seed Emulsion

A hexagonal tabular seed emulsion was prepared in the following manner.

10	A ₂	
	Ossein gelatin	24.2g
15	Distilled water	9657ml
	H-[CH ₂ CH ₂ O] _m -[CH(CH ₃)-CH ₂ O] ₁₇ -[CH ₂ CH ₂ O] _n -OH (wherein m+n=5.6; the molecular weight=1700) (in a 10% methanol solution)	6.78ml
20	KBr	10.8g
	A 10% nitric acid solution	114ml
25	B ₂	
	An aqueous 2.5N AgNO ₃ solution	2825ml
30	C ₂	
	KBr	824g
35	KI	23.5g
	Add distilled water to make	2825ml
40	D ₂	
	An aqueous 1.75N KBr solution	The following silver- potential controllable amount
45		

At 35°C and by making use of a mixing-stirrer described in JP Examined Publication Nos. 58-58288/1983 and 58-58289/1983, 464.3ml each of Solutions B₂ and C₂ were added to Solution A₂ in a double-jet method by taking 2 minutes, so that a nucleus formation was carried out.

After stopping the addition of Solutions B₂ and C₂, the temperature of resulting mixture solution A₂ was raised to 60°C by taking 60 minutes and the pH thereof was set at 5.0 with a 3% KOH solution. Thereafter, Solutions B₂ and C₂ were each added again at a flow-rate of 55.4ml/min for 42 minutes by a double-jet method. The silver electrode potentials between the time when raising the temperature of Solution A₂ from 35°C to 60°C and the time when the double-jetting Solutions B₂ and C₂ again, such potentials were controlled with Solution D₃ so as to be +8mv and +16mv, respectively; (wherein a saturated silver-silver chloride electrode was used as a standard electrode, and the potential was measured with a silver-ion selection electrode.)

After completing the addition of the solutions, the pH was adjusted to be 6 with a 3% KOH solution. Immediately after that, a desalting and washing treatments were carried out. The resulting seed emulsion was proved through an

electron microscope that the seed emulsion Em-2 was comprised of hexagonal tabular silver halide grains out of which not less than 90% of the whole projected area had the maximum ratio of the adjacent sides within the range of 1.0 to 2.0, and that the average thickness of the hexagonal tabular grains was $0.06\mu\text{m}$ and the average grain-size (converted into the diameter of a circle having the same area thereof) was $0.57\mu\text{m}$.

5

Preparation of Emulsion

The resulting hexagonal tabular seed emulsion Em-2 was taken in an equivalent amount of 0.023 mols per mol of silver of the emulsion and was then so dissolved as to be dispersed in an aqueous gelatin solution being kept at 60°C and containing a copolymer of polypropylene oxide (PO) and polyethylene oxide (EO). An aqueous silver nitrate solution was prepared in advance so that an average silver iodide content of the finally formed growth grains was to be 1.05 mol%. Successively, the resulting aqueous silver nitrate solution and a halide solution comprising potassium bromide and potassium iodide were each added by a double-jet method over a period of 107 minutes, while keeping the $\text{pH}=5.8$ and $\text{pAg}=9.0$ at 60°C all along. Resultingly, a tabular silver iodobromide emulsion EM-21 having an average grain-size of $1.51\mu\text{m}$, an average thickness of $0.25\mu\text{m}$ and an average aspect ratio of 6.04 could be prepared.

10

The resulting emulsion EM-21 was chemically sensitized while keeping a temperature of 50°C . Thereto, ammonium thiocyanate of 52mg, chloroauric acid of 1.4mg and sodium thiosulfate-pentahydrate of 15.0mg were each added. After 32 minutes passed therefrom, the silver iodide fine-grain type emulsion described in Example 1 was added in an equivalent amount of 1.80×10^{-3} mols. Thereafter, 2.34g of 4-hydroxy-6-methyl-1,3,3a,7-tetrazaindene was added so that the emulsion could be stabilized and was then subjected to an optimum chemical sensitization.

15

Ten minutes before starting the chemical sensitization, sensitizing dyes relating to the invention I-6, I-7, I-16, II-3, II-13, II-23 and II-26 were each added in an amount of 160mg per mol of silver halide, so that seven kinds of emulsions were prepared, respectively.

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The resulting chemically sensitized emulsions are hereinafter referred to as EM-22 through EM-28, respectively.

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Preparation of Internally Selenium-Sensitized Emulsion

The addition of the aqueous silver nitrate solution and halide solution of Emulsion EM-21 were temporarily stopped at the time when 90% of the silver nitrate solution was added, and ammonium thiocyanate and N,N-dimethyl selenourea were then added thereto so as to be in the amounts thereof of 7×10^{-6} mols and 2×10^{-6} mols each per mol of the silver used in the growth emulsion to be finally prepared, respectively. Four (4) minutes thereafter, the remaining aqueous silver nitrate solution and halide solution were further added to form the shell portions of the grains. Immediately after forming the shells, a desalting and washing treatments were carried out, so that emulsion EM-29 could be prepared. The configuration of the resulting emulsion grains were each proved to be approximately the same as those of EM-28.

30

The sensitizing dyes relating to the invention same as those used in EM-22 through EM-28 were added respectively to each of EM-29, and the same chemical sensitizers were further added thereto, respectively. Thereafter, the surfaces of the resulting grains were gold-sulfur-sensitized. The resulting emulsions are hereinafter referred to as EM-30 through EM-36, respectively.

35

The resulting EM-22 through EM-28 and EM-30 through EM-36 were coated on the respective samples, so that Samples No. 19 through No. 32 could be prepared.

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The resulting samples were each evaluated in quite the same manner as in Example 1. The results thereof will be shown in Table 2.

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Table 2

Sample No.	Em No.	Sensitizing dye	Chemical sensitization	Position selenium-sensitized	Photographic characteristics			Roller-mark
					Fog	Sensitivity	Dmax	
19 (Comparison)	EM-22	I-6	Gold-Sulfur	-	0.05	82	2.89	2
20 (Comparison)	EM-23	I-7	Gold-Sulfur	-	0.05	85	2.92	2
21 (Comparison)	EM-24	I-16	Gold-Sulfur	-	0.05	85	2.85	2
22 (Comparison)	EM-25	II-3	Gold-Sulfur	-	0.04	81	2.90	1
23 (Comparison)	EM-26	II-13	Gold-Sulfur	-	0.05	84	2.88	1
24 (Comparison)	EM-27	II-23	Gold-Sulfur	-	0.04	82	2.90	2
25 (Comparison)	EM-28	II-26	Gold-Sulfur	-	0.04	82	2.94	2
26 (Invention)	EM-30	I-6	Gold-Sulfur-Selenium	Interior	0.03	125	3.60	4
27 (Invention)	EM-31	I-7	Gold-Sulfur-Selenium	Interior	0.03	128	3.52	4
28 (Invention)	EM-32	I-16	Gold-Sulfur-Selenium	Interior	0.03	124	3.57	4
29 (Invention)	EM-33	II-3	Gold-Sulfur-Selenium	Interior	0.03	140	3.60	5
30 (Invention)	EM-34	II-13	Gold-Sulfur-Selenium	Interior	0.03	137	3.58	4
31 (Invention)	EM-35	II-23	Gold-Sulfur-Selenium	Interior	0.03	135	3.55	5
32 (Invention)	EM-36	II-26	Gold-Sulfur-Selenium	Interior	0.03	135	3.60	4

As is obvious from Table 2, it was proved that, according to the invention, a high sensitivity, a low fog, a high

maximum density and an excellent roller mark production resistance can be achieved even in the case of a tabular silver iodobromide emulsion grown up from a hexagonal tabular twinned crystal seed emulsion.

5 **Claims**

1. A silver halide photographic light-sensitive material comprising a support having on at least one side thereof a silver halide emulsion layer comprising a silver halide emulsion containing silver halide grains, which is prepared by a process comprising:

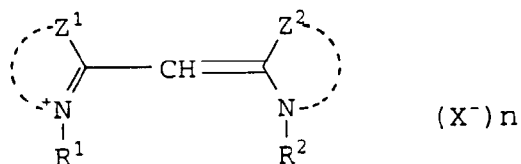
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- (i) forming the silver halide emulsion by mixing a silver salt and a halide salt in a dispersing medium and
 (ii) subjecting the emulsion formed to chemical ripening, wherein said silver halide grains are selenium-sensitized by adding a selenium compound after at least 70% of the whole silver salt used for forming the emulsion has been added and before the completion of the addition of the silver salt; said grains being spectrally sensitized with a spectral sensitizing dye represented by the following Formula (I) or Formula (II),

15

Formula (I)

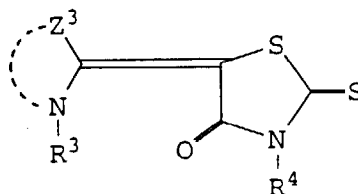
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Formula (II)

30



35

wherein Z¹, Z² and Z³ each represent a group of nonmetallic atoms necessary for forming oxazole, benzoxazole, naphthoxazole, thiazole, benzothiazole or naphthothiazole; R¹, R² or R³ each represent a substituted or unsubstituted alkyl group; R⁴ represents a substituted or unsubstituted alkyl group, or a substituted or unsubstituted aryl group; X is an acid anion; and n is an integer of 0 or 1.

40

2. The photographic material of claim 1, wherein said silver halide grains are tabular grains having an average diameter of 0.3 to 3.0 μm and having an average aspect ratio of grain diameter to grain thickness of not less than 1.2.
3. The photographic material of claim 2, wherein said tabular grains account for at least 50% of projected area of total silver halide grains contained in the silver halide emulsion.
4. The photographic material of claim 1, wherein said selenium compound is selected from an elemental selenium, an isoselenocyanate, a selenourea, a selenoketone, a selenoamide, a selenocarboxylic acid and an ester thereof, a selenide, and a selenophosphate.
5. The photographic material of claim 1, wherein selenium compound is added in the presence of a silver halide solvent selected from a thioether, a thiourea, an imidazole, a sulfite and a thiocyanate.
6. The photographic material of claim 1, wherein said sensitizing dye is added at a time during a period from the time after forming the silver halide grains to the time before completing chemical sensitization.

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7. The photographic material of claim 1, wherein silver halide fine grains having a grain size of 0.12 μm or less are added at a at a time during the course of the chemical sensitization.
8. The photographic material of claim 1, wherein said silver halide grains are silver iodobromide having an iodide content of 0.1 to 4.0 mol%.

Patentansprüche

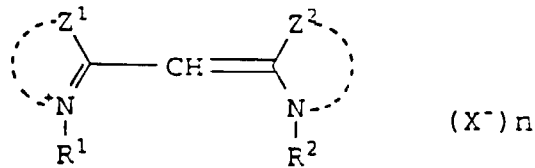
1. Lichtempfindliches photographisches Silberhalogenid-Aufzeichnungsmaterial mit einem Schichtträger und - auf mindestens einer seiner Seiten - einer Silberhalogenidemulsionsschicht, umfassend eine Silberhalogenidkörnchen enthaltende Silberhalogenidemulsion, die

(i) durch Zubereitung (der Silberhalogenidemulsion) durch Vermischen eines Silbersalzes mit einem Halogenidsalz in einem Dispergiermedium und

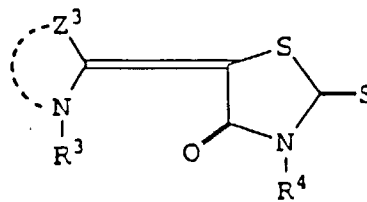
(ii) chemische Reifung (der gebildeten Emulsion)

zubereitet wurde, wobei die Silberhalogenidkörnchen durch Zusatz einer Selenverbindung nach Zugabe von mindestens 70% des gesamten zur Emulsionsbildung verwendeten Silbersalzes und vor Beendigung der Zugabe des Silbersalzes selensensibilisiert und mit einem spektralen Sensibilisierungsfarbstoff der folgenden Formel (I) oder (II)

Formel (I)



Formel (II)



worin bedeuten:

Z¹, Z² und Z³ jeweils eine Gruppe von zur Bildung von Oxazol, Benzoxazol, Naphthoxazol, Thiazol, Benzothiazol oder Naphthothiazol erforderlichen nichtmetallischen Atomen;

R¹, R² oder R³ jeweils eine substituierte oder unsubstituierte Alkylgruppe;

R⁴ eine substituierte oder unsubstituierte Alkylgruppe oder eine substituierte oder unsubstituierte Arylgruppe;

X ein Säureanion und

n eine ganze Zahl, nämlich 0 oder 1,

spektral sensibilisiert sind.

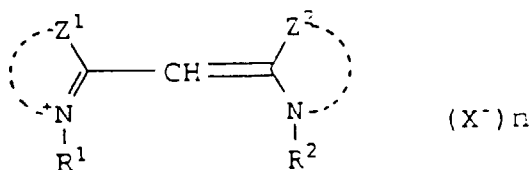
2. Photographisches Aufzeichnungsmaterial nach Anspruch 1, wobei es sich bei den Silberhalogenidkörnchen um tafelförmige Körnchen eines durchschnittlichen Durchmessers von 0,3 bis 3,0 μm und eines durchschnittlichen Seitenverhältnisses Korndurchmesser/Korndicke von nicht weniger als 1,2 handelt.

3. Photographisches Aufzeichnungsmaterial nach Anspruch 2, wobei die tafelförmigen Körnchen mindestens 50% der Projektionsfläche der gesamten in der Silberhalogenidemulsion enthaltenen Silberhalogenidkörnchen ausmachen.
- 5 4. Photographisches Aufzeichnungsmaterial nach Anspruch 1, wobei die Selenverbindung aus elementarem Selen, einem Isoselenocyanat, einem Selenharnstoff, einem Selenketon, einem Selenamid, einer Selencarbonsäure oder einem Ester derselben, einem Selenid und einem Selenophosphat ausgewählt ist.
- 10 5. Aufzeichnungsmaterial nach Anspruch 1, wobei die Selenverbindung in Gegenwart eines Silberhalogenidlösungsmittels, ausgewählt aus einem Thioether, einem Thioharnstoff, einem Imidazol, einem Sulfit und einem Thiocyanat, zugesetzt wird.
- 15 6. Photographisches Aufzeichnungsmaterial nach Anspruch 1, wobei der Sensibilisierungsfarbstoff zu einem Zeitpunkt während eines Zeitraums nach Bildung der Silberhalogenidkörnchen bis vor Beendigung der chemischen Sensibilisierung zugesetzt wird.
7. Photographisches Aufzeichnungsmaterial nach Anspruch 1, wobei zu einem Zeitpunkt während der chemischen Sensibilisierung feine Silberhalogenidkörnchen mit einer Korngröße von 0,12 µm oder weniger zugesetzt werden.
- 20 8. Photographisches Aufzeichnungsmaterial nach Anspruch 1, wobei es sich bei den Silberhalogenidkörnchen um Silberiodbromid eines Iodidgehalts von 0,1 bis 4,0 Mol-% handelt.

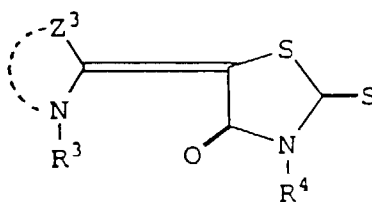
Revendications

- 25 1. Un matériau photographique à l'halogénure d'argent étant sensible à la lumière renfermant un support ayant, sur au moins une face, une couche d'halogénure d'argent comprenant une émulsion contenant des grains d'halogénure d'argent qui sont préparés par un procédé renfermant:
- 30 (i) la formation d'une émulsion d'halogénure d'argent en mélangeant un sel d'argent dans un milieu dispersif et
(ii) la soumission de l'émulsion ainsi formée au mûrissement chimique dont lesdits grains d'halogénure d'argent sont sensibilisés par du sélénium en rajoutant un composé de sélénium après qu'au moins 70% de la totalité du sel d'argent qui a été utilisé pour la formation de l'émulsion aient été rajoutés et avant la terminaison de
35 l'addition du sel d'argent; lesdits grains étant spectralement sensibilisés par un colorant spectral sensibilisant représenté par la formule suivante (I) ou la formule (II),

formule (I)



formule (II)



dans lesquelles Z^1 , Z^2 et Z^3 représentent chacun un radical d'ion non-métallique, nécessaire pour la formation d'oxazole, de benzoxazole, de naphthoxazole, de thiazole, de benzthiazole ou de naphthothiazole; R^1 , R^2 ou R^3 représentent chacun un radical d'alkyle substitué ou non-substitué; R^4 représente un radical d'alkyle substitué ou non-substitué, ou un radical d'aryle substitué ou non-substitué; X est un ion acide; et n est un nombre entier de 0 ou 1.

2. Le matériau photographique suivant la revendication 1, dans lequel lesdits grains sont des grains tabulaires ayant un diamètre moyen compris entre 0,3 à 3,0 μm et ayant un rapport d'aspect moyen entre le diamètre granulaire et l'épaisseur granulaire d'au moins de 1,2.
3. Le matériau photographique suivant la revendication 2, dans lequel lesdits grains tabulaires représentent au moins 50% du domaine projeté de la totalité de grains d'halogénure d'argent contenus dans l'émulsion d'halogénure d'argent.
4. Le matériau photographique suivant la revendication 1, dans lequel ledit composé de sélénium est sélectionné d'un sélénium élémentaire, d'un isocyanate de sélénium, d'un séléno-urée, d'un séléno-cétone, d'un séléno-amide, d'un acide séléno-carboxylique et d'un ester de ce-dernier, d'un sélényde et d'un séléno-phosphate.
5. Le matériau photographique suivant la revendication 1, dans lequel du composé de sélénium est rajouté en présence d'un solvant étant sélectionné d'un thioether, d'une thiourée, d'un imidazole, d'un sulfite et d'un thiocyanate.
6. Le matériau photographique suivant la revendication 1, dans lequel ledit colorant sensibilisant est rajouté pendant une période entre le moment suivant la formation des grains d'halogénure d'argent et le moment précédant la terminaison de la sensibilisation chimique.
7. Le matériau photographique suivant la revendication 1, dans lequel des grains fins d'halogénure d'argent ayant une taille de 0,12 μm ou moins sont rajoutés à un moment donné où la sensibilisation chimique est en cours.
8. Le matériau photographique suivant la revendication 1, dans lequel lesdits grains d'halogénure d'argent représentent du iodobromure d'argent ayant un contenu d'iodure compris entre 0,1 à 4,0 % en mole.