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(11)

EP 0 895 121 B1

(12)

EUROPEAN PATENT SPECIFICATION

(45) Date of publication and mention
of the grant of the patent:
21.01.2004 Bulletin 2004/04

(51) Int Cl.7: **G03C 1/09**

(21) Application number: **98202305.3**

(22) Date of filing: **08.07.1998**

(54) **A photosensitive element containing a silver halide emulsion chemically sensitized with a chalcogenic compound**

Photoempfindliches Element, das eine mit einer Chalkogenverbindung chemisch sensibilisierte Silberhalogenidemulsion enthält

Elément photosensible contenant une émulsion à l'halogénure d'argent sensibilisée chimiquement avec un composé chalcogénure

(84) Designated Contracting States:
BE DE FR GB

(30) Priority: **01.08.1997 EP 97202395**

(43) Date of publication of application:
03.02.1999 Bulletin 1999/05

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(56) References cited:
EP-A- 0 458 278
DE-A- 19 648 008

DE-A- 19 635 098

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Description

FIELD OF THE INVENTION

[0001] The present invention relates to a photosensitive image-forming element, more particularly to a high sensitive photographic silver halide emulsion with improved photographic properties.

BACKGROUND OF THE INVENTION

[0002] In the industrial world a continuous need exists for photosensitive image-forming materials with increased sensitivity and image-quality. These characteristics are related which means that influencing one parameter has its consequences for the other parameter and vice versa. This relationship can clearly be experienced in silver halide photographic materials. An increase of sensitivity of a photographic silver halide material can for instance be realised by an increase of the mean size of the silver halide grains contained in the emulsion of the said material which in turn often results in a decrease of sharpness or contrast of the said material. This particular problem can for instance be solved by increasing the efficiency for electron trapping of the latent-image of a silver halide grain which can be realised by chemical sensitization in the presence of a compound which increases the electron-trapping efficiency mentioned hereinbefore. This compound can be for instance a sulphur compound, a gold salt or a combination of both. It has been noticed in recent years that chemical sensitization with a chalcogenic compound other than sulphur is increasingly practiced in order to enhance photographic sensitivity. These compounds in turn often give rise to an increasing tendency of fog formation and often to stability problems.

Looking at these problems which are for instance related to selenium sensitization various suggestions have already been formulated in the patent literature. Examples of selenium compounds which were suggested for use in order to get an increased sensitivity were substituted selenourea (as described e.g. in EP-A 0 280 031, EP-A 0 458 278), selenoethers (e.g. in JP-A 2-132434, JP-A 4-271341, JP-A 6-175258, US-A 5,532,120), diselenides (e.g. in EP-A 0 703 492), selenoesters (e.g. in US-A 5,306,613), selenoketones (e.g. in EP-A 0 476 345), phosphineselenides (e.g. in EP-A 0 506 009) and various others. However as mentioned hereinbefore selenium sensitization easily causes fog as a result of the depth of the electron trap of the formed sensitivity speck which is larger than the depth of the sensitivity speck created by sulphur sensitization. At the same time the stability and the gradation of a selenium sensitized emulsion decrease if no precautions are taken. An alternative way is described by G.J.Burgmaier et al in US-P 4,810,626 which provides a silver halide photographic material comprising a sensitizing amount of a tetrasubstituted urea compound with the following general structure: $N(A_1R_1)(A_2R_2)-C(=X)-N(A_3R_3)(A_4R_4)$ where X equals S, Se or Te and where S is preferred (due to the greater solubility of the thiourea compound in aqueous solutions where silver halide sensitization occurs and also to the ready availability of starting materials for thiourea synthesis). In this formula at least one of A_1R_1 to A_4R_4 contains a nucleophilic group bonded to the urea nitrogen atom through a 2 or 3 member chain and where each of R_1 , R_2 , R_3 and R_4 independently represents an alkylene, cycloalkylene, carbocyclic arylene, heterocyclic arylene, alkarylene or aralkylene group; and each of A_1 , A_2 , A_3 and A_4 independently is hydrogen or represents a carbocyclic, sulfinic, sulfonic, hydroxamic, mercapto, sulfonamido or a primary or secondary amino nucleophilic group. The chalcogenic atom in the urea compounds of the Burgmaier patent is activated by the nucleophilic group which is generally called a neighbouring group. This type of reactions is summarized by the term 'neighbouring group participation', describing all the intramolecular reactions and all the reactions which involve non-electrostatic through-space interactions between groups within the same molecule. Another example is in principle also described in EP-A 0 458 278.

In US-P 4,810,626 and EP-A 0 458 278 as well substantial increase of sensitivity can be realized with this type of chemical sensitizer but as in the other patent proposals described hereinbefore no clear demonstration is given that fog could be kept under control. Accordingly there still exists a strong demand for chemical chalcogenic sensitizers other than the corresponding S-sensitizers that can give high sensitivity in combination with an acceptable low fog level.

OBJECTS OF THE INVENTION

[0003] It is therefore a first object of the invention to provide a photosensitive element with high photosensitivity combined with low fog level.

[0004] It is also an object of this invention to provide a class of chalcogenic compounds which are functionalised with a neighbouring group facilitating the kinetics of chemically sensitizing a silver halide emulsion.

[0005] It is another object of the invention to provide a photosensitive element comprising at least one silver halide emulsion which is chemically sensitized in the presence of said new class of chalcogenic sensitizers.

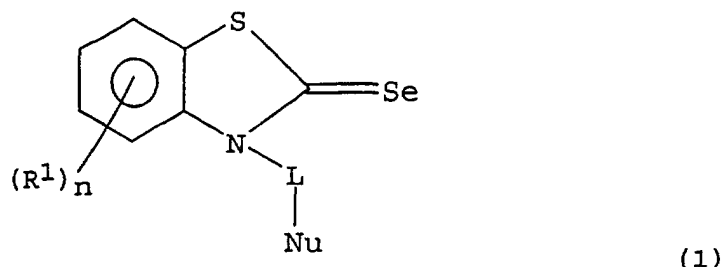
[0006] It is a further object of the invention to provide a method for making a photosensitive element comprising on a support a silver halide emulsion which is chemically sensitized in the presence of a said new class of chalcogenic

compounds.

[0007] Further objects and advantages of the invention will become apparent from the description hereinafter.

SUMMARY OF THE INVENTION

[0008] The objects just mentioned are realized by a photosensitive element which comprises on a support at least one photosensitive silver halide emulsion layer and at least one non-light sensitive layer, wherein the silver halide emulsion layer comprises a chalcogenic compound satisfying formula (1):



wherein:

- Nu represents a nucleophilic group which brings a free electron pair in heterolytic bond formation,
- L represents a divalent linking group representing each group which can position an electron donating atom in any of the 5th to the 8th position vis-a-vis the carbon atom of the carbon-chalcogen double bond,
- R¹ represents a substituted or non-substituted alkyl-, aryl-, aralkyl- or heteroaryl-group, or one or more R¹-substituents together represent the atoms necessary to form a substituted or non-substituted, saturated or unsaturated ring,
- n equals 0, 1, 2, 3 or 4.

[0009] The invention also provides a method for chemically sensitizing a silver halide emulsion by means of a chalcogenic compound according to the formula (1) at least partly in the presence of a silver halide solvent and a spectral sensitizer at a pH-value situated between 3 and 10, a pAg-value situated between 6 and 11 and a temperature in the range of from 40 °C up to 95 °C.

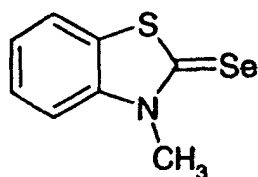
[0010] Preferred embodiments of the invention are disclosed in the dependent claims hereinafter.

DETAILED DESCRIPTION OF THE INVENTION

[0011] While the present invention will hereinafter be described in connection with preferred embodiments thereof, it will be understood that it is not intended to limit the invention to those embodiments. On the contrary, it is intended to cover all alternatives, modifications, and equivalents as may be included within the spirit and scope of the invention as defined by the appending claims.

[0012] The present invention as described in the foregoing summary will now be explained in detail starting with an illustration of the compounds satisfying formula (1) of the present invention.

[0013] Heterocyclic selenones are known for quite a long time in organic synthesis as useful synthetic intermediates for functional group transformation such as the conversion of epoxides to double bonds (Calo et al, Synthesis, (1976) 200-201). Potentially they can be used for chemical sensitization of silver halide photographic emulsions in which treatment the photosensitivity is increased. However, the reactivity of the basic products is too low to be useful under conventional circumstances, used for chemical sensitization, as illustrated by the use of 3-methyl-2-selenoxobenzothiazole (Ref-1) as chemical sensitizer.

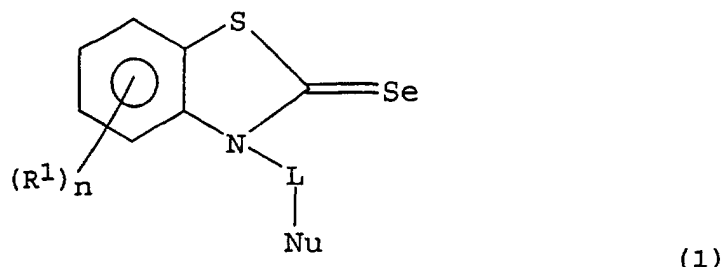


Ref-1

[0014] We found that the introduction of a nucleophilic group at a well defined position in the structure of heterocyclic selenones, rendered them usefull as chemical. sensitizer for photographic applications.

[0015] The introduction of anchimeric assistance or neighboring group participation into the structure, is capable of lowering the pH or the free silver ion concentration needed for chemical sensitization, allowing this sensitivity step under conditions where the tendency of fog formation can be controlled. A general discussion on neighboring group participation is given in Advanced Organic Chemistry, J. March, 4th edition, John Wiley and Sons, p. 308-326 and in Neighboring Group Participation Volume I, B. Capon and S. P. McManus, Plenum Press (1976).

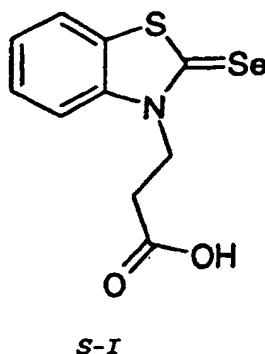
[0016] We found that compounds of the following general formula (1) are particularly useful in the present invention.

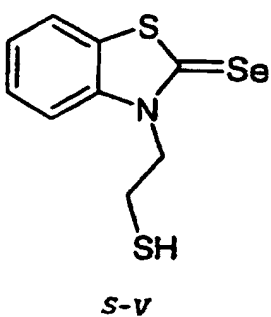
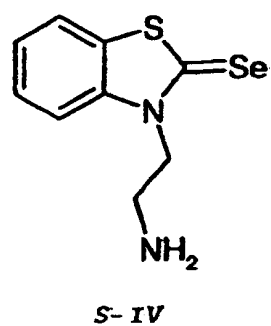
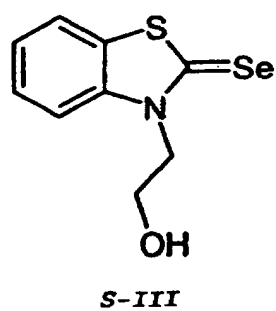
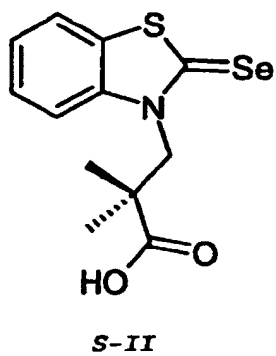


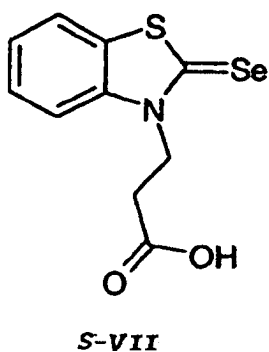
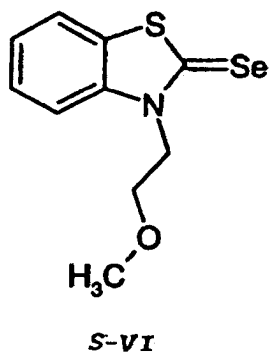
wherein:

- Nu represents a nucleophilic group which brings a free electron pair in heterolytic bond formation,
- L represents a divalent linking group representing each group which can position an electron donating atom in any of the 5th to the 8th position vis-a-vis the carbon atom of the carbon-chalcogen double bond,
- R¹ represents a hydrogen atom, a substituted or non-substituted alkyl-, aryl-, aralkyl- or heteroaryl-group, or one or more R¹-substituents together represent the atoms necessary to form a substituted or non-substituted, saturated or unsaturated ring,
- n equals 0, 1, 2, 3 or 4.

[0017] Typical, but not limiting examples of compounds useful in the present invention are given below.



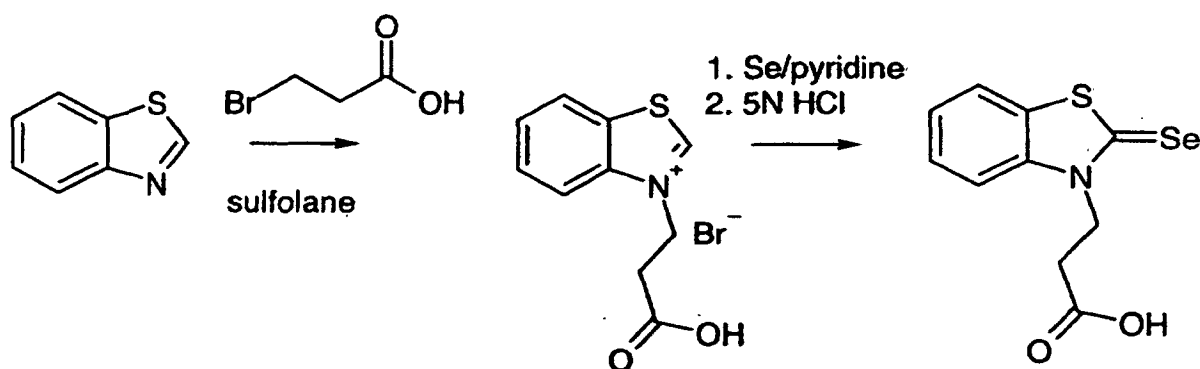




30 **[0018]** The compounds used in the present invention can be prepared by conventional synthetic strategies. As an illustration, the synthesis of sensitizer I and VI will be described.

The synthesis of sensitizer I:

35 **[0019]** General scheme:



50 (a) The alkylation of benzothiazole.

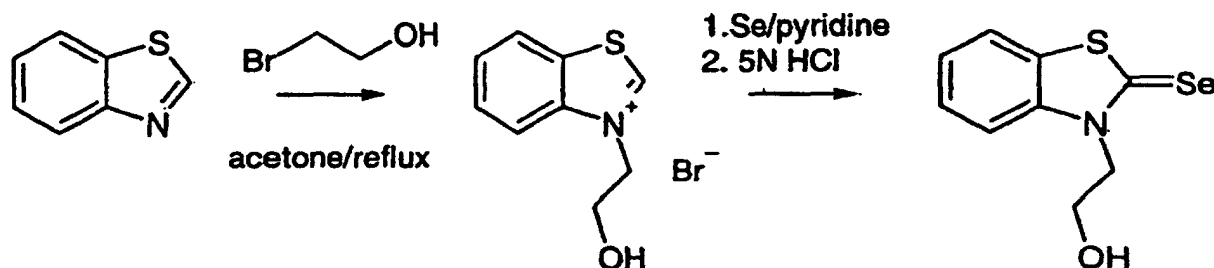
55 **[0020]** A mixture of 54 g (0.4 mol) benzothiazole, 61.2 g (0.4 mol) 2-bromopropionic acid and 40 ml sulfolane was heated to 130°C. After 90 minutes, the alkylated intermediate started to precipitate. On further alkylation the mixture solidified. The mixture was allowed to cool down and 100 ml acetone was added. The precipitated product was isolated by filtration, treated with 200 ml acetone, isolated by filtration for a second time, washed with 50 ml acetone and dried under reduced pressure (96 g, 83%)

(b) The synthesis of the selenone.

[0021] 14.4 g (0.05 mol) of the quaternary salt was dissolved in 100 ml pyridine. 3.95 g (0.05 mol) selenium was added. The mixture was refluxed for 6 hours and then allowed to cool down. After filtration to remove residual selenium, the mixture was poured into 600 ml of a 5 N solution of hydrochloric acid. After precipitation the product was isolated by filtration. The product was finally purified by preparative column chromatography (eluent $\text{CH}_2\text{Cl}_2/\text{MeOH}$ 95/5). (7.5 g, 52.5 %).

The synthesis of sensitizer III:

[0022] General scheme :



(a) The alkylation of benzothiazole.

[0023] A mixture of 67.5 g (0.5 mol) benzothiazole and 62.5 g (0.5 mol) bromoethanol in 100 ml acetone was refluxed for four hours. The mixture was allowed to cool down. The product precipitated on diluting with methylenechloride. The product was isolated by filtration, washed with ether and dried under reduced pressure (25g, 19%).

(b) The synthesis of the selenone.

[0024] 13 g (0.05 mol) of the quaternary salt was dissolved in 75 ml pyridine. 3.95 g (0.05 mol) selenium was added and the mixture was refluxed for four hours. The residual selenium was removed by filtration and 225 ml water was added to the mixture. The product crystallized from the medium with some difficulty. The crude product was isolated by filtration and recrystallized from ethanol (8g, 62%).

[0025] The amount of the chalcogenic compound of formula (1) which can be used in the material of the present invention can vary depending on the type of compound, the type of silver halide grain, the conditions of chemical sensitization, etc. The amount of the said chalcogenic sensitizer is usually in the range of 10^{-9} to 10^{-3} but preferably in the range of 10^{-8} to 10^{-4} .

The introduction of the chemical chalcogenic sensitizer represented by formula (1) can be done in various ways before starting or during the chemical sensitization procedure. Said chalcogenic sensitizers which are more or less water soluble can be added to the dispersion of silver halide crystals as an aqueous solution if desired mixed up with a water soluble organic solvent. Said sensitizers which are not water soluble can be introduced after solving the compound in a water soluble organic solvent which is normally used for the introduction of water insoluble products in a photographic silver halide emulsion. An alternative way for the introduction of an insoluble chemical chalcogenic sensitizer is using an 'oil-in-water' dispersion or if desired dispersion as disclosed in EP-A 0 703 492 by T.Yoshida et al. In the last named patent the said compound is therefore added to the silver halide emulsion before chemical sensitization in the form of a solid partial dispersion soluble in water.

[0026] Combinations of the chalcogenic compounds of the present invention can be used but can also be combined with other chalcogenic or other more "classical" sensitizers. The chemical sensitization itself can be carried out in the presence of a sulphur compound, a chalcogenic compound and if desired in the presence of a noble metal (e.g. gold). The sensitization in the presence of the chalcogenic compound represented by formula (1) and if desired together with any other chemical sensitizer, can be used in combination with a reduction sensitization too.

[0027] In the noble metal sensitization a salt of a noble metal (e.g., gold, iridium, palladium, platinum) is used but gold salts are preferably used as noble metal sensitizer. Examples of gold sensitizers which are often used include chloroauric acid, goldsulfide, chloroaurate salts, aurithiocyanate and goldselenide. The amount of the noble metal sensitizer can vary between 10^{-7} and 10^{-2} mole per mole silver halide.

[0028] Sulfur sensitization can be carried out with sulphur compounds like thiosulphates, thioureas, rhodamines, etc. The sulphur sensitizer can also be used in an amount of about 10^{-7} to 10^{-2} mole per mole silver halide.

[0029] In reduction sensitization a reducing compound is used like thiourea dioxide, hydrazine derivatives, sulphinic acid, polyamine compounds, stannous chloride, borane compounds, reductones like ascorbic acid, etc. Reduction sensitization can also be carried out at a low pAg or at a high pH or at both and if desired at elevated temperature. This kind of sensitization is referred to silver ripening'. More information can be found in Research Disclosure, Vol307,307105 and in P.Glaftides "Chimie et Physique Photographique.", P.Montel - Paris, 5th Ed., 1987.

[0030] The chemical sensitization with compounds of the present invention satisfying formula (1) should be carried out in the presence of a silver solvent which preferably should be thiocyanate. This should be added as a sodium, a potassium or an ammonium salt but it is not limited thereto. The thiocyanate salt can be added before or during the addition of said compound of the invention and before, during or after the addition of any other chemical sensitizer which can be or should be used together with the said selenium sensitizer. The amount of the thiocyanate which should be present together with said chemical sensitizer represented by formula (1) is limited between 10^{-6} and 10^{-1} mole per mole silver halide and preferably situated between 10^{-5} and 10^{-2} mole per mole silver halide.

[0031] The chemical sensitization has also to be carried out in the presence of a spectral sensitizer. This can be a dye out of different classes including polymethine dyes comprising cyanines, merocyanines, tri-, tetra- and polynuclear cyanines and merocyanines, oxanols, hemioxanols, styryls, merostyryls and so on. Many representative examples of these dye-classes can be found in EP-A 0 618 492, EP-A 0 638 841, US-A 5,308,748, US-A 5,310,645 and 5 US-A 5,338,656. Some types of spectral sensitizers are preferably used for this invention. It concerns the symmetrical or the unsymmetrical cyanine dyes containing a benzoxazole, benzthiazole or benzimidazole nucleus.

[0032] Sometimes more than one spectral sensitizer may be used in the case that a larger part of the spectrum has to be covered. Combinations of several spectral sensitizers are sometimes used to get supersensitization, which means that in a certain region of the spectrum the sensitization is greater than that from any concentration of one of the dyes alone or that which would result from the additive effect of the dyes. Generally supersensitization can be attained by using selected combinations of spectral sensitizing dyes and other addenda such as stabilizers, development accelerators or inhibitors, brighteners, coating aids, and so on. Many of the possibilities in spectral sensitization which can be important with respect to this invention are described in Res.Discl., 38957(1996) section V.

[0033] The chemical sensitization described in the present invention should be carried out within limited chemical and physical conditions. Therefore the pAg should be in the range of 6 till 11 but preferably between 7 and 10. Further the pH should be in the range of 3 to 10, preferably 4 to 8.5 while the temperature should be situated in the range between 40 to 95 degrees C, preferably between 45 to 85 degrees C, though there is no particular limitation to any of these conditions.

[0034] The silver halide emulsion can be prepared in various ways by conventional methods. These start always with a nucleation phase followed by a grain growth phase. In this last phase of the emulsion preparation reactants can be added to the reaction vessel in the form of solutions of silver and halide salts or in the form of preformed silver halide nuclei or fine grains which easily dissolve in the precipitation medium.

[0035] The individual reactants can be added through surface or subsurface delivery tubes by hydrostatic pressure or by an automatic delivery system for maintaining the control of pH and/or pAg in the reaction vessel and of the rate of the reactant solutions introduced in it. The reactant solutions or dispersions can be added at a constant rate or a constantly increasing, decreasing or fluctuating rate, if desired in combination with stepwise delivery procedures. More details about the possible ways in making a silver halide emulsion which can be principally used in practicing this invention are summarized in Res.Discl.,38957 (1996)591-639 section I-C.

[0036] Beside the individual reactants necessary to form silver halide crystals additional chemical metal salts can be added for occlusion in the crystal lattice. Such compound is replacing an appropriate of silver and halide ions in the silver halide lattice. These so-called dopants can be distinguished from the metal complexes which are added just before coating as an additive by EPR- or ENDOR-technique. These dopants can be used to modify the crystal structure or the crystal properties and can therefore be employed to influence many photographic properties like sensitivity, reciprocity failure, gradation, pressure sensitivity, fog, stability, etc. When coordination complexes or even oligomeric coordination complexes are used the different ligands bound at the central metal ion can be occluded in the crystal lattice too and in this way influence the photographic properties of the silver halide materials as well. Dopants which are introduced in emulsions of the present invention are those which can act as a permanent or as a non-permanent electron trap.

[0037] The doping procedure itself can normally be executed at any stage during the grain growth phase of the emulsion preparation. It is important to know that the dopants can also be added in an indirect way by the addition of a dispersion containing very fine soluble silver halide grains or nuclei comprising the dopant. More additional information about the introduction and the use of dopants in the emulsion crystals of this invention can be found in Research Disclosure, 38957 (sept. 1996), section I-D.

[0038] The photographic emulsions prepared in this way contain silver halide crystals comprising chloride, bromide

or iodide alone or in combinations thereof. Other silver salts which can be incorporated in a limited amount in the silver halide lattice are silver phosphate, silver thiocyanate, silver citrate and some other silver salts. The chloride and bromide halide can be combined in all ratios to form a silverchlorobromide salt. Iodide ions however can be coprecipitated with chloride and/or bromide ions in forming a iodohalide with an iodide amount which depends on the saturation limit of iodide in the lattice with the given halide composition; this means up to a maximum amount of about 40 mole percent in silver iodobromide and up to at most 13 mole percent in silver iodochloride both based on silver.

[0039] The composition of the halide can change in the crystal in a continuous or discontinuous way. Emulsions containing crystals composed of various sections with different halide compositions are used for several photographic applications. So a structure with a difference in halide composition between the center and the rest of the crystal (what is called 'core-shell'-emulsion) or with more than two crystal parts differing in halide composition (called a 'band'-emulsion) may occur. The changes in halide composition can be realised by direct precipitation or in an indirect way by conversion where fine silver halide grains of a certain halide composition are dissolved in the presence of the so-called host grains forming a 'shell' or 'band' on the given grain.

[0040] The crystals formed by the methods described above have a morphology which can be tabular or non-tabular. In tabular crystals the aspect ratio (ratio of equivalent circular diameter to thickness) can vary from low (<2) over 'medium' (2 till 8) to high (>8) where specially in the case of the ultra thin tabular crystals high aspect ratios can be realised. The major faces of the formed tabular grains can have a {111} or a {100}-habitus the structure of which is (respectively) stable or has to be stabilised (for instance by a 'habitus modifying agent'). In the class of non-tabular grains there are a lot of possibilities which can be divided in the more regular shaped crystals or the crystals with a mixed crystal habit.

[0041] For certain applications it can be important to apply a well-defined amount of iodide on the crystal surface under controlled conditions in order to get reproducible sensitometric results after image-wise exposure and subsequent processing. This can be done by using an iodide releasing agent as described in EP-A 0 561 415 and in EP-A 0 563 708 and applied on emulsions before, during or after the chemically sensitization in addition to the method and the conditions of the present invention as described hereinbefore.

[0042] The photographic emulsions of the present invention can be comprised of silver chloride, silver bromide and silver iodide, silver chlorobromide, silver chloroiodide, silver bromoiodide, silver chlorobromoiodide or mixtures thereof.

[0043] A hydrophilic colloid is used as a binder or a protective colloid for the emulsion or any other layer of the photographic material of the invention. Gelatin is an advantageous hydrophilic colloid. The preparation of conventional lime-treated or acid-treated gelatin has been described in e.g. "The Science and Technology of Gelatin", edited by A. G. Ward and A. Courts, Academic Press 1977, page 295 and further. The gelatin can also be enzyme-treated as described in Bull. Soc. Sci. Phot. Japan, Nr 16, page 30 (1966).

[0044] Gelatin may, however, be replaced in part or integrally by synthetic, semi-synthetic, or natural polymers. Synthetic substitutes for gelatin are e.g. polyvinyl alcohol, poly-N-vinyl pyrrolidone, polyvinyl imidazole, polyvinyl pyrazole, polyacrylamide, polyacrylic acid, and derivatives thereof, in particular copolymers thereof. Natural substitutes for gelatin are e.g. other proteins such as zein, albumin and casein, cellulose, saccharides, starch, and alginates. In general, the semi-synthetic substitutes for gelatin are modified natural products e.g. gelatin derivatives obtained by conversion of gelatin with alkylating or acylating agents, by grafting of polymerisable monomers on gelatin or prehardened gelatins with blocked functional groups as a consequence of this prehardening treatment, cellulose derivatives such as hydroxyalkyl cellulose, carboxymethyl-cellulose, phthaloyl cellulose, and cellulose sulphates and even potato starch.

[0045] Further synthetic high molecular compounds described in JP-B-52-16365, Journal of The Society of Photographic Science and Technology of Japan, Vol. 29(1), 17, 22(1966), *ibid.*, Vol. 30(1), 10, 19(1967), *ibid.*, Vol. 30(2), 17 (1967), and *ibid.*, Vol. 33(3), 24(1967) may be used as a dispersion medium. Also the crystal habit restraining agent described in EP-A 0 534 395 may be used.

[0046] Part of gelatin may further be replaced with a synthetic or natural high-molecular material.

[0047] An interesting substitute for gelatin may be silica as has been described in the published EP-A's 0 392 092, 0 517 961, 0 528 476 and 0 649 051 and 0 704 749. As has been set forth in EP-A 0 528 476 a method of preparing a silver halide light-sensitive photographic material incorporating layers of silver halide precipitated in colloidal silica serving as a protective colloid is given. In this document the silver halides are prepared in colloidal silica, leading to emulsion crystals that are stable at the end of the precipitation, without however having a predictable mean crystal diameter and crystal size distribution. These problems have been overcome as has been described in EP-A 0 682 287, for the preparation of crystals rich in silver chloride, wherein clearly defined circumstances wherein such crystals can be prepared: during the precipitation stage of regular silver chloroiodide crystals amounts of silica sol and of stabilising onium compound(s), should be optimised in order to avoid uncontrolled formation and growth of aggregates.

[0048] After precipitation the emulsions can be coagulated and washed in order to remove the excess soluble salts. These procedures are together with different alternative methods like dia- or ultrafiltration and ion-exchange described in Res. Discl., 38957(1996), section III. The silver halide emulsions of this invention which are prepared in one of the ways described hereinbefore contain crystals which have a spherical equivalent diameter (SED) of not more than 1.5

μm but preferable less than $1.0\ \mu\text{m}$. The spherical equivalent diameter (SED) of the crystal represents the diameter of the sphere which has the same volume as the average volume of the silver halide crystals of the said 0 emulsion.

[0049] The emulsions can be surface-sensitive emulsions which form latent images primarily on the surface of the silver halide grains or they can be emulsions forming their latent-image primarily in the interior of the silver halide grain. Further the emulsions can be negative-working emulsions such as surface sensitive emulsions or unfogged internal latent image-forming emulsions. However direct-positive emulsions of the unfogged, latent image-forming type which are positive-working by development in the presence of a nucleating agent, and even pre-fogged direct-positive emulsions can be used in the present invention.

[0050] The photographic elements comprising the said silver halide emulsions can include various compounds which should play a certain role in the material itself or afterwards in the processing, 5 finishing or warehousing the photographic material.

[0051] These products can be stabilizers and anti-foggants. The antifoggant prevents occurrence of a fog while a stabilizer has a function of stabilizing the photographic property. The antifoggant and the stabilizer are used in preparation, storage or processing stage of the photographic material. The antifogging agent and the stabilizers can be azoles, mercaptopyrimidines, mercaptotriazines, azaindenes, etc. (see Res.Discl., 38957 (1996) section VII).

[0052] The hydrophilic colloidal layer (silver halide emulsion layer, backing layer, antihalo-layer, etc.) of the photographic material can contain an inorganic or an organic hardening agent (see Res.Discl.,38957(1996) section IIB), brighteners (see Res.Discl.,38957(1996) section VI), light absorbers and scattering materials (see Res.Discl.,38957 (1996) section VIII), coating aids (see Res.Discl.,38957(1996) section IXA), antistatic agents (see Res.Discl.,38957 (1996) section IXC), matting agents (see Res.Discl.,38957(1996) section IXD) and development modifiers (see Res. Discl.,38.957(1996) section XVIII).

[0053] The photographic element may further comprise various other additives such as e.g. compounds improving the dimensional stability of the photographic element, ultraviolet absorbers and spacing agents. Suitable additives for improving the dimensional stability of the photographic element are e.g. dispersions of a water-soluble or hardly soluble synthetic polymer e.g. polymers of alkyl(meth)acrylates, alkoxy(meth)acrylates, glycidyl (meth)acrylates, (meth)acrylamides, vinyl esters, acrylonitriles, olefins, and styrenes, or copolymers of the above with acrylic acids, methacrylic acids, α - β -unsaturated dicarboxylic acids, hydroxyalkyl (meth)acrylates, sulfoalkyl (meth)acrylates, and styrene sulphononic acids.

[0054] Suitable UV-absorbers are e.g. aryl-substituted benzotriazole compounds as described in US-A 3,533,794, 4-thiazolidone compounds as described in US-A 3,314,794 and 3,352,681, benzophenone compounds as described in JP-A 56-2784, cinnamic ester compounds as described in US-A's 3,705,805 and 3,707,375, butadiene compounds as described in US-A 4,045,229, and benzoxazole compounds as described in US-A 3,700,455 and those described in Res.Discl.,38957 (sept. 1996) Section VI, wherein also suitable optical brighteners are mentioned.

[0055] Spacing agents may be present of which, in general, the average particle size is comprised between 0.2 and $10\ \mu\text{m}$. Spacing agents can be soluble or insoluble in alkali. Alkali-insoluble spacing agents usually remain permanently in the photographic element, whereas alkali-soluble spacing agents usually are removed therefrom in an alkaline processing bath. Suitable spacing agents can be made e.g. of polymethyl methacrylate, of copolymers of acrylic acid and methyl methacrylate, and of hydroxypropylmethyl cellulose hexahydrophthalate. Other suitable spacing agents have been described in US-A 4,614,708.

[0056] Prior to coating any thickening agent may be used in order to regulate the viscosity of the coating solution, provided that they do not particularly affect the photographic characteristics of the silver halide emulsion in the coated photographic material. Preferred thickening agents include aqueous polymers such as polystyrene sulphononic acid, dextran, sulphuric acid esters, polysaccharides, polymers having a sulphononic acid group, a carboxylic acid group or a phosphoric acid group as well as colloidal silica. Polymeric thickeners well-known from the literature resulting in thickening of the coating solution may even be used in combination with colloidal silica. Patents concerning thickening agents are e.g. US-A 3,167,410; Belgian Patent No. 558.143 and JP-A's 53-18687 and 58-36768. Negative effects on physical stability possibly resulting from the addition of polymeric compounds can be avoided by exclusion of those compounds and by restricting extra additions of colloidal silica.

[0057] In order to reach a high hardening degree the layer binder should of course dispose of an acceptably high number of functional groups, which by reaction with an appropriate hardening agent can provide a sufficiently resistant layer. Such functional groups are especially the amino groups, but also carboxylic groups, hydroxy groups, and active methylene groups. Hardeners may be added to the antistress layer, covering one or more light-sensitive silver halide emulsion layers rich in chloride before or during the coating procedure, or to one or more of the said emulsion layers. The binders of the photographic element, especially when the binder used is gelatin, can be hardened with appropriate hardening agents such as those of the epoxide type, those of the ethylenimine type, those of the vinylsulfone type e.g. 1,3-vinylsulphonyl-2-propanol, chromium salts e.g. chromium acetate and chromium alum, aldehydes e.g. formaldehyde, glyoxal, and glutaraldehyde, N-methylol compounds e.g. dimethylolurea and methyloldimethylhydantoin, dioxan derivatives e.g. 2,3-dihydroxy-dioxan, active vinyl compounds e.g. 1,3,5-triacryloyl-hexa-hydro-s-triazine, active

halogen compounds e.g. -2,4-dichloro-6-hydroxy-s-triazine, and mucohalogenic acids e.g. mucochloric acid and mucophenoxy-chloric acid. These hardeners can be used alone or in combination. The binders can also be hardened with fast-reacting hardeners such as carbamoylpyridinium salts. Formaldehyde and phloroglucinol can e.g. be added respectively to the protective layer(s) and to the emulsion layer(s). Further suitable possibilities for hardening can be found in Res.Discl.,38957(sept. 1996) section IIB.

[0058] The photographic elements can be coated on a variety of supports which can be flexible or rigid. The flexible materials include plastic films and papers while the rigid materials include glass, metals, etc. The surface of the support is generally subjected to undercoating treatment (like corona discharge, irradiation with ultraviolet rays, etc) to enhance the adhesion with the silver halide emulsion layer (see Res.Discl.,38957(1996) section XV and the references cited therein). The photographic elements can be exposed to actinic radiation, specially in the visible, near-ultraviolet and near-infrared region of the spectrum, to form a latent image (see Res.Discl., 38957(1996) section XVI). This latent-image can be processed in order to form a visible image (see Res.Discl.,38957 (1996) section XIX).

[0059] Various exposure means can be employed for exposure of the photographic material of the invention. As the light source, any optional light source releasing a radiation corresponding to the sensitivity wavelength of the photographic material can be employed. Examples of the light sources generally used include natural light, incandescent lamp, halogen lamp, mercury lamp, fluorescent lamp and all types of flash light sources. Light sources that emit light in the ultraviolet to infrared region can be also used as recording light sources. Photographic materials can for example also be exposed to gas lasers, semiconductor lasers, light emitting diodes or plasma light sources. In the same way the material can be exposed to a LCD light source or to a fluorescent surface given by a phosphor stimulated with electron rays. This latent-image formed in the silver halide crystals after exposure can be processed in order to form a visible image. Therefore various methods are known and many developing, fixing and stabilizing agents are described for the formation of photographic silver image. These know-how for processing photographic silver halide materials which principally can be used in relation with the present invention is described in Res.Discl.,176043(dec.1978) sections XIX to XXIV and in Res.Discl.,38957 (1996) section XIX)

[0060] In the conventional way of processing most of the materials are developed by means of a liquid containing hydroquinone as main developing agent usually in combination with a so-called auxiliary developer. In an alternative way of processing hydroquinone is incorporated in the photographic material itself while the processing liquid is an mere alkaline solution. However it is important to realize that hydroquinone is suspect in various ways specially in an ecological and medical point of view. The present invention is also related with a more ecological way of processing where hydroquinone is at least partly replaced by ascorbic acid as developing agent. Ascorbic acid should be interpreted in a broad sense and includes ascorbic acid isomers, derivatives, salts and analogous compounds (including some reductones and reductic acid derivatives). The most preferred compounds are ascorbic acid, isoascorbic acid and their salts. Useful combinations of developers containing an ascorbic acid developing agent which should be preferably used in the scope of the present invention is described for many applications (in graphics, radiography, etc) in Res. Discl., 37152 (March 1995) 185-224.

[0061] The photographic emulsions according of the present invention can also be used in multi-layered multicolor materials. These materials comprise a support and two or more silver halide emulsion layers that have different spectral sensitivities. The multi-layered color photographic material generally comprises at least one red-sensitive emulsion layer, at least one green-sensitive emulsion layer and at least one blue-sensitive emulsion layer on a support. A non-light sensitive layer may be provided between two or more emulsion layers having the same color sensitivity. Otherwise, another emulsion layer having a different color sensitivity can be provided between two or more emulsion layers having the same color sensitivity. A light-reflecting layer such as a layer of silver halide grains can be provided under a high sensitive layer, particularly under a high blue-sensitive layer in order to enhance the sensitivity.

[0062] The silver halide material can also contain different types of couplers which can be incorporated in color photographic materials. The red-sensitive emulsion layer contains generally a cyan coupler, the green-sensitive layer generally contains a magenta coupler and the blue-sensitive emulsion layer generally contains a yellow coupler. All the information which can be important for the application of the present invention in these type of materials is extensively described in Res.Discl., 38957 (sept. 1996) section X. More information in relation with the various color applications which belong also to the scope of the present invention, can be found in US-P 05,532,120 too.

[0063] Processing to form a visible dye image for colour materials means contacting the element with a colour developing agent in order to reduce developable silver halide and to oxidize the colour developing agent which in turn normally reacts with a coupler to form a dye (see Res. Discl.,38957(1996)section XX).

[0064] The invention is described hereinafter by way of the following example.

EXAMPLE

[0065] This example demonstrates the possibilities of a silver halide. emulsion comprising tabular AgBr(I)-crystals which have been chemically sensitized with a chalcogenic chemical sensitizer of the present invention compared with

a chemical sensitizer which lack the neighbouring group participating action. It also illustrates the influence of other chemical sensitizers like sulphur and gold salts.

[0066] For this example three solutions were used during the precipitation of emulsion (2):

- 5 Solution 1: 1.5 litre of an aqueous solution containing 500 grams of silver nitrate.
 Solution 2: 1.5 litre of an aqueous solution containing 350 grams of potassium bromide.
 Solution 3 : 1.5 litre of an aqueous solution containing 341 grams of potassium bromide and 7.5 grams of potassium iodide.

10 Preparation of Emulsion 1.

(a)Nucleation step:

- 15 **[0067]** 15.2ml of solutions 1 and 2 were introduced into a reaction vessel in 35 seconds using the double jet technique. Said reaction vessel initially contained 2127 ml of distilled water at 51°C, 12.5 grams of potassium bromide and 6 grams of gelatin. After one minute the reaction temperature of this mixture was raised to 70°C in 20 minutes and a solution of 47.5 grams of phthalated gelatin in 475 ml of distilled water were added. After 6 minutes the neutralisation step was started.

20 (b)Neutralisation step:

[0068] Solution 1 was added to the reaction vessel at a rate of 7.5 ml per minute to reach a pAg value of 8.99 (-10 mV measured against a saturated Ag/AgCl reference electrode), whereafter the first growth step was started.

25 (c)First growth step:

- 30 **[0069]** A double jet precipitation was started using solutions 1 and 2 which continued for 45 minutes 44 seconds. During this precipitation, the pAg value was kept constant at 8.99. The flow rate of solution 1 was 7.5 ml per minute at the start, linearly increasing to 14.5 ml per minute at the end of the precipitation. Thereafter the second neutralisation step was started.

(d)Second neutralisation step:

- 35 **[0070]** Solution 1 was added to the reaction vessel at a rate of 7.5 ml per minute in order to reach a pAg value of 7.38, whereafter the precipitation further continued with a second growth step.

(e)Second growth step:

- 40 **[0071]** An amount of 0,904 l of solution 1 was injected into the reaction vessel at a rate of 7.5 ml per minute at the start, linearly increasing to 22.5 ml per minute at the end of the precipitation. The pAg was kept constant at 7.38 using solution 3 for 60 minutes and 15 seconds.

[0072] The emulsion was flocculated after addition of a small amount of polystyrene sulphonic acid and the acidification to a pH value of 3.4. After sedimentation the mother liquid was removed, distilled water added and remaining salts were washed out after repeating this procedure.

- 45 The AgBr(I) crystals of the emulsion prepared in this way were containing 1 mol % of iodide and had a spherical equivalent diameter (SED) of 0.93 µm while the thickness was 0.22 µm.

Chemical sensitization of Emulsion 1.

- 50 **[0073]** The emulsion was redispersed and after addition of various compounds as described hereinafter chemically ripened to an optimal fog-sensitivity relationship. After positioning the emulsion at pH value of 5.5 and at the pAg value of 8.08 (at a temperature of 40 °C) the following solutions were added successively (per 500 gram of AgNO₃):

- 55 - 4 ml of a 1.03 molar solution of potassium thiocyanate, after 10 minutes followed by
 - 0.2 ml of a 0.00476 molar solution of sodium thiosulphonic acid, after 5 minutes by
 - 780 ml of a solution of hydro-5,5'-dichloro-3,3'-bis-(n-sulfobutyl)-9-ethyloxacarbo-cyanine hydroxide (2.5 gram per litre) as a green sensitizer and after 30 minutes
 - a solution of (0.00632 mole/l) sodiumthiosulphate (called "S-cpd"), a solution of (0.001456 mole/l) auric chloride

("Au-cpd") and a solution of a selenium compound ("Se-cpd").

Tabel 1.1

Chemical sensitization conditions for the tabular AgBr(I)-emulsion				
Exp Nr.	"Se-cpd"	"S-Cpd" (ml)	"Au-Cpd" (ml)	
2.1	S-I	2.5	6.5	Invention
2.2	S-III	2.5	6.5	Invention
2.3	Ref-1	2.5	6.5	Comparison
2.4	S-I	2.5	10	Invention
2.5	S-III	2.5	10	Invention
2.6	Ref-1	2.5	10	Comparison
2.7	S-I	2.5	15	Invention
2.8	S-III	2.5	15	Invention
2.9	Ref-1	2.5	15	Comparison

[0074] In this particular case several selenobenzthiazoles were used as chalcogenic compound which were added in an amount of 3 ml of an 0,0035 N solution per 500 grams of silver halide. The chemical sensitization of all the sample emulsions were carried out at 60°C in the presence of a sulphur and a gold compound (see respectively "S-cpd" and "Au-cpd" in the table) in amounts (number of ml of the named solutions) as indicated in Table 1.1 (always per 500 grams of silver halide).

Coating of Emulsion 1.

[0075] After chemical sensitization each emulsion was stabilized with 10 ml of 0.00375 mole/l 4-hydroxy-6-methyl-1,3,3a,7-tetra-azaindene and after addition of the normal coating additives the solutions were coated simultaneously together with a protective layer containing 0.7 g of gelatine per m² per side on both sides of a polyethylene terephthalate film support having a thickness of 175 µm. The resulting photographic material contained per side an amount of silver halide corresponding to 3.90 grams of AgN03 per m². Hardening of the layers was performed with a bis-vinylsulfon-methane compound.

Exposure and processing procedure.

[0076] Samples of these coatings were exposed with green light of 540 nm during 0.1 seconds using a continuous wedge and were processed during the 90 seconds cycle described below. The processing occurred in a glutaraldehyde containing hydroquinone/1-phenyl-3-pyrazolidinone developer marketed by Agfa-Gevaert N.V. under the trade name G138. Fixation was carried out in fixer G334, also marketed by Agfa-Gevaert N.V.. The processing was executed in a HT-300 processing machine (marketed by Agfa-Gevaert as well) with the following time (in seconds (sec.)) and temperature (in °C) characteristics:

loading	3.4 sec.
developing	23.4 sec./ 33°C high or low activity developer
cross-over	3.8 sec.
fixing	15.7 sec./ 33°C in fixer AGFA G334
cross-over	3.8 sec.
rinsing	15.7 sec./ 20°C.
drying	32.2 sec. (cross-over time included)
total time	98.0 sec.

[0077] The results of the various experiments are summarized in Table 1.2 where the experiment number Exp Nr. corresponds with the same number in Table 1.1. The density which was realised after processing as a function of the light dose was measured and used to determine the following parameters:

- Fog level (with an accuracy of 0.001 density),
- the sensitivity S is given at a density of 1 above fog expressed in log (It): a decrease with a factor of 0.30 is indicative for a sensitivity increase with a factor of 2,
- the contrast g was calculated between densities 1.0 and 2.5 above fog.

Table 1.2

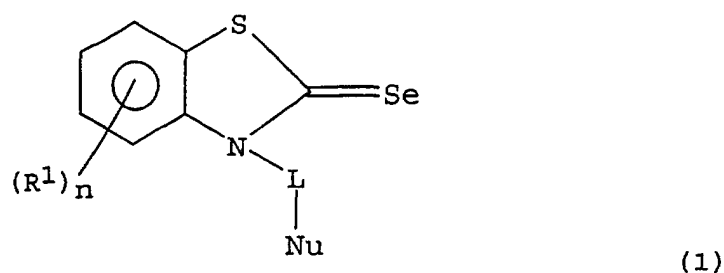
Sensitometric results of the emulsions after chemical sensitization under different ripening conditions.					
Exp Nr.	Se-compd	Fog 10^{-3}	Sensitivity S	γ	
2.1	S-I	19	1.76	1.74	Invention
2.2	S-III	14	1.94	1.96	Invention
2.3	Ref-1	11	2.02	2.03	Comparison
2.4	S-I	22	1.59	2.30	Invention
2.5	S-III	18	1.88	2.13	Invention
2.6	Ref-1	14	1.97	1.97	Comparison
2.7	S-I	24	1.66	2.34	Invention
2.8	S-III	17	2.04	1.95	Invention
2.9	Ref-1	16	2.09	1.99	Comparison

[0078] From these results it can be clearly seen that the chemical sensitization of the tabular AsBrl-emulsion with the chalcogenic compounds of this invention results in a substantial increase in sensitivity compared with the compounds which lacks neighbouring group participation. This effect can be optimized if the chemical sensitization is carried out under controlled conditions of the amount of sulfur and gold sensitizer.

[0079] Having described in detail preferred embodiments of the current invention, it will now be apparent to those skilled in the art that numerous modifications can be made therein without departing from the scope of the invention as defined in the following claims.

Claims

1. A photosensitive element comprising on a support, at least one silver halide emulsion layer and a non-light sensitive hydrophilic colloid layer **characterized in that** the silver halide emulsion layer comprises a chalcogenic compound satisfying formula (1) :



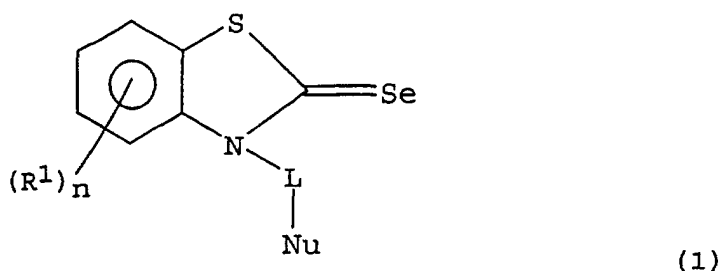
wherein:

- Nu is a nucleophilic group,
- L is a divalent linking group,
- R¹ represents a substituted or non-substituted alkyl-, aryl-, aralkyl- or heteroaryl-group, or one or more R¹ substituents together represent the atoms necessary to form a substituted or non-substituted saturated or unsaturated ring;
- n equals 0, 1, 2, 3 or 4.

2. A photosensitive element according to claim 1, wherein in the general formula (1) the nucleophilic group Nu is a nucleophilic group forming a heterolytic bond with its free electron pair.
3. A photosensitive element according to claim 1, wherein in the general formula (1) the divalent linking group L represents a group which can position an electron donating atom in any of the 5th to the 8th position vis-a-vis the carbon atom of the carbon-chalcogen double bond.
4. A photosensitive element according to any of the claims 1 to 3, wherein the silver halide emulsion is chemically sensitized with a chalcogenic compound satisfying formula (1) in an amount of 10^{-8} to 10^{-4} mole based on one mole of silver halide.
5. A method for chemically sensitizing a silver halide emulsion **characterized by** the step of adding a sulphur and/or a gold sensitizer and a chalcogenic compound as defined in any of the claims 1 to 4, **characterized by** the step of adding a silver halide solvent to the chalcogenic compound and by adding a spectral sensitizer, in following conditions:
 - a pAg in the range of from 6 to 11,
 - a pH in the range of from 3 to 10,
 - a temperature in the range of 40 up to 95 °C.
6. A method according to claim 5, wherein the silver halide solvent is a thiocyanate salt.
7. A method according to claim 5 or 6, wherein the silver halide solvent is present in an amount of 10^{-6} to 10^{-1} mole per mole of silver halide.
8. A method according to any of the claims 5 to 7, wherein the spectral sensitizer is a symmetrical or unsymmetrical cyanine dye containing a benzoxazole, benzthiazole or a benzimidazole nucleus.

Patentansprüche

1. Strahlungsempfindliches Element, das auf einem Träger wenigstens eine Silberhalogenid-Emulsionsschicht und eine nicht lichtempfindliche hydrophile Kolloidschicht umfasst, **dadurch gekennzeichnet, dass** die Silberhalogenid-Emulsionsschicht eine der Formel (1) entsprechende Chalkogenverbindung enthält:



in der bedeuten:

- Nu eine nukleophile Gruppe,
 - L eine zweiwertige verbindende Gruppe,
 - R¹ eine gegebenenfalls substituierte Alkyl-, Aryl-, Aralkyl- oder Heteroarylgruppe, oder ein oder mehrere Substituenten R¹ zusammen die zur Bildung eines gegebenenfalls substituierten, gesättigten oder ungesättigten Ringes benötigten Atome;
 - n eine Zahl gleich 0, 1, 2, 3 oder 4.
2. Strahlungsempfindliches Element nach Anspruch 1, **dadurch gekennzeichnet, dass** es sich bei der nukleophilen Gruppe Nu in der allgemeinen Formel (1) um eine nukleophile Gruppe, die mit ihrem freien Elektronenpaar eine heterolytische Bindung bildet, handelt.

3. Strahlungsempfindliches Element nach Anspruch 1, **dadurch gekennzeichnet, dass** die zweiwertige verbindende Gruppe L in der allgemeinen Formel (1) eine Gruppe, die ein elektronenlieferndes Atom in 5-, 6-, 7- oder 8-Stellung zu dem Kohlenstoffatom der Kohlenstoff-Chalkogen-Doppelbindung anordnen kann, bedeutet.

4. Strahlungsempfindliches Element nach einem der Ansprüche 1 bis 3, **dadurch gekennzeichnet, dass** die Silberhalogenid-Emulsion unter Verwendung einer der Formel (1) entsprechenden Chalkogenverbindung in einer Menge von 10^{-8} bis 10^{-4} mol, bezogen auf 1 mol Silberhalogenid, chemisch sensibilisiert wurde.

5. Durch die nachstehenden Schritte gekennzeichnetes Verfahren für die chemische Sensibilisierung einer Silberhalogenid-Emulsion : Zusetzen eines Sensibilisators aus Schwefel und/oder Gold sowie einer Chalkogenverbindung nach einem der Ansprüche 1 bis 4, Zusetzen eines Silberhalogenid-Lösungsmittels zu der Chalkogenverbindung und Zusetzen eines Spektralsensibilisators, unter folgenden Bedingungen:

- ein pAg-Wert im Bereich von 6 bis 11,
- ein pH-Wert im Bereich von 3 bis 10,
- eine Temperatur im Bereich von 40 bis 95°C.

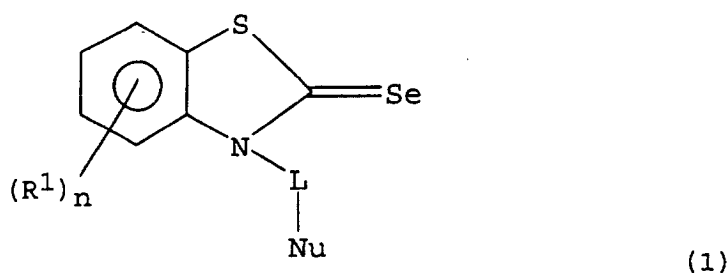
6. Verfahren nach Anspruch 5, **dadurch gekennzeichnet, dass** es sich bei dem Silberhalogenid-Lösungsmittel um ein Thiocyanatsalz handelt.

7. Verfahren nach Anspruch 5 oder 6, **dadurch gekennzeichnet, dass** das Silberhalogenid-Lösungsmittel in einer Menge von 10^{-6} bis 10^{-1} mol pro mol Silberhalogenid enthalten ist.

8. Verfahren nach einem der Ansprüche 5 bis 7, **dadurch gekennzeichnet, dass** es sich bei dem Spektralsensibilisator um einen symmetrischen oder unsymmetrischen Cyaninfarbstoff handelt, der einen Benzoxazol-, Benzthiazol- oder Benzimidazol-Ring enthält.

Revendications

1. Élément photosensible comportant sur un support au moins une couche d'émulsion aux halogénures d'argent ainsi qu'une couche colloïdale hydrophile non sensible à la lumière, **caractérisé en ce que** la couche d'émulsion aux halogénures d'argent comprend un composé chalcoène répondant à la formule (1) :



dans laquelle:

- Nu est un groupe nucléophile,
- L est un groupe de liaison bivalent,
- R¹ représente un groupe alkyle, aryle, aralkyle ou hétéroaryle substitués ou non, ou un ou plusieurs substituants R¹ représentent ensemble les atomes nécessaires pour constituer un noyau saturé ou insaturé substitué ou non;
- n est égal à 0, 1, 2, 3 ou 4.

2. Élément photosensible selon la revendication 1, **caractérisé en ce que** ledit groupe nucléophile Nu dans la formule générale (1) est un groupe nucléophile formant une liaison hétérolytique avec son doublet électronique libre.

3. Élément photosensible selon la revendication 1, **caractérisé en ce que** ledit groupe de liaison bivalent L dans la

formule générale (1) représente un groupe pouvant disposer un atome cédant des électrons en l'une quelconque des positions 5 à 8 par rapport à l'atome de carbone de la double liaison carbone-chalcogène.

4. Élément photosensible selon l'une quelconque des revendications 1 à 3, **caractérisé en ce que** ladite émulsion aux halogénures d'argent est sensibilisée par voie chimique au moyen d'un composé chalcogène répondant à la formule (1) en une quantité de 10^{-8} à 10^{-4} mole par rapport à 1 mole d'halogénure d'argent.
5. Procédé de sensibilisation chimique pour émulsions aux halogénures d'argent **caractérisé par** les étapes consistant à ajouter un sensibilisateur sulfuré et/ou aurique ainsi qu'un composé chalcogène tel que défini dans l'une quelconque des revendications 1 à 4, **caractérisé par** l'étape consistant à ajouter un solvant des halogénures d'argent au composé chalcogène et à ajouter un sensibilisateur spectral, dans les conditions suivantes:
 - un pAg compris entre 6 et 11,
 - un pH compris entre 3 et 10,
 - une température dans l'intervalle de 40 à 95 °C.
6. Procédé selon la revendication 5, **caractérisé en ce que** ledit solvant des halogénures d'argent est un sel de thiocyanate.
7. Procédé selon la revendication 5 ou 6, **caractérisé en ce que** ledit solvant des halogénures d'argent est contenu en une quantité de 10^{-6} à 10^{-1} mole par mole d'halogénure d'argent.
8. Procédé selon l'une quelconque des revendications 5 à 7, **caractérisé en ce que** le sensibilisateur spectral est un colorant cyanine symétrique ou asymétrique contenant un noyau benzoxazole, benzothiazole ou benzimidazole.