The present invention relates to photography and, more particularly, to photographic products particularly adapted for employment in photographic diffusion transfer color processes.

The primary objects of the present invention are to provide photographic products particularly adapted for employment in diffusion transfer photographic color processes; to provide photographic products which comprise a photosensitive laminate which contains a plurality of essential layers including a dimensionally stable transparent layer, a photosensitive silver halide emulsion layer having dye image-forming material associated therewith which is soluble and diffusible, in alkaline, at a first pH, a permeable opaque layer, a permeable polymeric layer dyable by the dye image-forming material, a polymeric acidic layer containing sufficient acidifying groups to effect reduction of a selected processing solution having the first pH to a second pH at which the dye image-forming material is insoluble and nondiffusible, and a dimensionally stable transparent layer; to provide photographic diffusion transfer products comprising a film unit including a photosensitive laminate, of the last-identified type, in combination with a rupturable container retaining an alkaline processing composition having the first pH; to provide a diffusion transfer film unit, of the last-identified type, having the container fixedly positioned and extending transverse to the leading edge of the photosensitive laminate whereby to effect, upon application of compressive pressure, discharge of the alkaline processing composition intermediate the opposed surfaces of the dyable polymeric layer and the photosensitive silver halide emulsion next adjacent thereto; and to provide photographic diffusion transfer color processes employing such products.

Other objects of the invention will in part be obvious and will in part appear hereinafter.

The invention accordingly comprises the product possessing the features, properties and the relation of components and the process involving the several steps and the relation and order of one or more of such steps with respect to each of the others which are exemplified in the following detailed disclosure, and the scope of the application of which will be indicated in the claims.

For a fuller understanding of the nature and objects of the invention, reference should be had to the following detailed description taken in connection with the accompanying drawings wherein:

FIGURE 1 is a perspective view of a photographic film unit embodying the invention; FIGS. 2, 4 and 6 are diagrammatic enlarged cross-sectional views of the film unit of FIGURE 1, along section line 2—2, illustrating the association of elements during the three illustrated stages of the performance of a diffusion transfer process, for the production of a multicolor transfer image according to the invention, the thickness of the various materials being exaggerated, and wherein FIG. 2 represents an exposure stage, FIG. 4 represents a processing stage and FIG. 6 represents a product of the process; and FIGS. 3, 5 and 7 are diagrammatic, further enlarged cross-sectional views of the film unit of FIGS. 2, 4 and 6, along section lines 3—3, 5—5 and 7—7, respectively, further illustrating, in detail, the arrangement of layers comprising the photosensitive laminate during the three illustrated stages of the transfer process.

As disclosed in U.S. Patent No. 2,983,606, issued May 9, 1961, a photosensitive element containing a dye developer, that is, a dye which is a silver halide developing agent, and a silver halide emulsion may be exposed and wetted by a liquid processing composition, for example, by immersion, coating, spraying, flowing, etc., in the dark, and the exposed photosensitive element is superposed prior to, during, or after wetting, on a sheetlike support element which may be utilized as an image-receiving element. In a preferred embodiment, the liquid processing composition is applied to the photosensitive element in a substantially uniform layer as the photosensitive element is brought into superposed relationship with the image-receiving layer. The liquid processing composition, positioned intermediate the photosensitive element and the image-receiving layer, permeates the emulsion to initiate development of the latent image contained therein. The dye developer is immobilized or precipitated in exposed areas as a consequence of the development of the latent image. This immobilization is apparently, at least in part, due to a change in the solubility characteristics of the dye developer upon oxidation and especially as regards its solubility in alkaline solutions. It may also be due in part to a tanning effect on the emulsion by oxidized developing agent, and in part to a localized exhaustion of alkaline as a result of development. In unexposed and partially exposed areas of the emulsion, the dye developer is unreacted and diffusible and thus provides an imagewise distribution of unoxidized dye developer dissolved in the liquid processing composition, as a function of the point-to-point degree of exposure of the silver halide emulsion.

At least part of this imagewise distribution of unoxidized dye developer is transferred, by immersion, to a superposed image-receiving layer or element, said transfer substantially excluding oxidized dye developer. The image-receiving element receives a depositwise diffusion, from the developed emulsion, of unoxidized dye developer without appreciably disturbing the imagewise distribution thereof to provide the reversed or positive color image of the developed image. The image-receiving element may contain agents adapted to mordant or otherwise fix the diffused, unoxidized dye developer. If the color of the transferred dye developer is affected by changes in the pH of the image-receiving element, this pH may be adjusted in accordance with well-known techniques to provide a pH affording the desired color image and the photosensitive element at the end of a suitable immersion period.

The dye developers, as noted above, are compounds which contain, in the same molecule, both the chromophoric system of a dye and also a silver halide developing function. By "a silver halide developing function" is meant a grouping adapted to develop exposed silver halide. A preferred silver halide development function is a hydroquinonyl group. Other suitable developing functions include ortho-dihydroxyphenyl and ortho- and para-aminophenyl substituted hydroxyphenyl groups. In general, the
development function includes a benzenoid developing function, that is, an aromatic developing function which forms quinonoid or quinone substances when oxidized. Such quinones or quinonoids may be obtained using color image-forming components such as, for example, the previously mentioned dye developers, in diffusion transfer processes by several techniques. One such technique contemplate obtaining multicolor transfer images utilizing dye developers by employment of an integral multilayer photosensitive element, such as is disclosed in the aforementioned U.S. Patent No. 2,983,606, and particularly with reference to FIG. 9 of the patent's drawing, wherein at least two selectively sensitized photosensitive strata, superposed on a single support, are processed, simultaneously and without separation, with a single, common image-receiving layer. A suitable arrangement of this type comprises a support carrying a red-sensitive silver halide emulsion stratum, a green-sensitive silver halide emulsion stratum and a blue-sensitive silver halide emulsion stratum, said emulsions having associated therewith, respectively, for example, a cyan dye developer, a magenta dye developer and a yellow dye developer. The dye developer may be utilized in the silver halide emulsion layer, for example, in the form of particles, or it may be employed as a layer behind the appropriate silver halide emulsion strata. Each set of silver halide emulsion and associated dye developer strata are disclosed to be optionally separated from other sets by suitable laterlayers, for example, by a layer of gelatin or polyvinyl alcohol. In certain instances, it may be desirable to incorporate a yellow filter in front of the green-sensitive emulsion and such yellow filter may be incorporated in an interlayer. However, where desirable, a yellow dye developer of the appropriate spectral characteristics and particle size, in a state capable of functioning as a yellow filter may be employed. In such instances, a separate yellow filter may be omitted.

The dye developers are preferably selected for their ability to provide colors that are useful in carrying out subtractive color photography, that is, the previously mentioned cyan, magenta and yellow. The dye developers employed may be incorporated in the respective silver halide emulsion or, in the preferred embodiment, in a separate layer behind the respective silver halide emulsion. Specifically, the dye developer may, for example, be in a coating layer behind the respective silver halide emulsion and such a layer of dye developer may be applied by use of a coating solution containing about 0.5 to 8%, by weight, of the respective dye developer distributed in a film-forming natural, or synthetic, polymer, for example, gelatin, polyvinyl alcohol, and the like, adapted to be permeated by the chosen diffusion transfer fluid processing composition.

As disclosed in my aforementioned copending U.S. application Ser. No. 234,864 now U.S. Patent 3,362,819, image-receiving elements, particularly adapted for employment in diffusion transfer processes of the type disclosed in aforementioned U.S. Patent No. 2,983,606, wherein the image-receiving elements are separated from contact with a superposed photosensitive element, subsequently to substantial transfer image formation, preferably comprise a support layer possessing on one surface thereof, in sequence, a polymeric acid layer and an image-receiving layer adapted to provide a visible image upon transfer to said layer of diffusable dye image-forming substance, and most preferably include an inert timing or spacer layer intermediate the polymeric acid layer and the image-receiving layer.

As set forth in my last-mentioned application, the polymeric acid layer comprises polymers such as carboxylic acid groups, such as carboxylic acid and sulfonic acid groups, which are capable of forming salts with alkali metals, such as sodium potassium etc., or with organic bases, particularly quaternary ammonium bases, such as tetramethyl ammonium hydroxide, or potentially acid-yielding groups, such as amphotrides or lactones, or other groups which are capable of reacting with bases to capture and retain them. The acid-reacting group is, of course, non-diffusible from the acid polymer layer. In the preferred embodiments disclosed, the acid polymer contains free carboxyl groups and the transfer processing composition employed contains a large concentration of sodium and/or potassium ions. The acid polymers stated to be most useful are characterized by containing free carboxyl groups, being insoluble in water in the free acid form, and by forming water-soluble sodium and/or potassium salts. One may also employ polymers containing carboxylic acid anhydride groups, at least some of which preferably have been converted to free carboxyl groups prior to imbibition. While the most readily available polymeric acid derivatives of cellulose or of vinyl polymers, polymeric acids from other classes of polymers may be used. As examples of specific polymeric acids set forth in the application, mention may be made of dibasic acid half-ester derivatives of cellulose which derivatives contain free carboxyl groups, e.g., cellulose acetate hydroxyphthalate, cellulose acetate hydroxyglutarate, cellulose acetate hydroxy succinate, ethyl cellulose hydroxy succinate, ethyl cellulose acetate hydroxy succinate, cellulose acetate hydroxy glutarate phthalate; ether and ester derivatives of cellulose modified with sulfoanhydrides, e.g., with ortho-sulfobenzoic anhydride; poly-styrene sulfonic acid; carboxymethyl cellulose; polyvinyl hydroxy acetate hydroxy phthalate, polyvinyl acetate hydroxy phthalate, polyacrylic acid; acetics of polyvinyl alcohol with carboxy or sulfo substituted aldehydes, e.g., o-, m-, or p-benzaldehyde sulfonic acid or carboxylic acid; partial esters of ethylene/maleic anhydride copolymers; partial esters of methyl-vinyl ether/maleic anhydride copolymers; etc.

The acid polymer layer is disclosed to contain at least sufficient acid groups to effect a reduction in the pH of the image layer from a pH of about 13 to 14 to a pH of at least 11 or lower at the end of the imbibition period, and preferably to a pH of about 5 to 8 within a short time after imbibition. As previously noted, the pH of the processing composition preferably is of the order of at least 13 to 14.

It is, of course, necessary that the action of the polymeric acid be so controlled as not to interfere with either development of the negative or image transfer of unoxidized dye developers. For this reason, the pH of the image layer is kept as a level of pH 12 to 14 until the dye image has been formed after which the pH is reduced very rapidly to at least pH 11, and preferably about pH 9 to 10, before the transfer image is separated and exposed to air. Unoxidized dye developers containing hydroquinonyl developing radicals diffuse from the negative to the positive as the sodium or other alkali salt. The diffusion rate of such dye image-forming components thus is at least partly a function of the alkali concentration, and it is necessary that the pH of the image layer remain on the order of 12 to 14 until transfer of the necessary quantity of dye has been accomplished. The subsequent pH reduction, in addition to its desirable effect upon image light stability, serves a highly valuable photographic function by substantially terminating further dye transfer. The processing technique thus effectively minimizes changes in color balance as a result of longer imbibition times in multicolor transfer processes using multilayer negatives.

In order to prevent premature pH reduction during transfer processing, as evidenced, for example, by an undesired reduction in positive image density, the acid groups are so placed on the acid polymer layer that the rate of their availability to the alkali is controllable, e.g., as a function of the rate of swelling of the polymer layer which rate in turn has a direct relationship to the diffusion rate of the alkali ions. The desired distribution of the acid groups in the acid polymer layer may be effected by mixing the acid polymer...
with a polymer free of acid groups, or lower in concentration of acid groups, and compatible therewith, or by using only the acid polymer but selecting one having a relatively lower proportion of acid groups. These embodiments are illustrated, respectively, in the cited copending application, by (a) a mixture of cellulose acetate and cellulose acetate hydrogen phthalate and (b) a cellulose acetate hydrogen phthalate polymer having a much lower percentage of phthalyl groups than the first-mentioned cellulose acetate hydrogen phthalate.

It is also disclosed that the layer containing the polymeric acid may contain a water-insoluble polymer, preferably a cellulose ester, which acts to control or modulate the pH reduction by the alkali, that is, when the polymer acid is formed. As examples of cellulose esters contemplated for use, mention is made of cellulose acetate, cellulose acetate butyrate, etc. The particular polymers and combinations of polymers employed in any given embodiment are, of course, selected so as to have adequate wet and dry strength and when necessary or desirable, suitable subcoats may be employed to help the various polymeric layers adhere to each other during storage and use.

The inert spacer layer of the aforementioned copending application, for example, an inert spacer layer comprising polyvinyl alcohol or gelatin, acts to "time" the pH reduction by the polymeric acid layer. This timing is disclosed to be a function of the rate at which the alkali diffuses through the inert spacer layer. It was stated to have been found that the pH does not drop until the alkali has passed through the spacer layer, i.e., the pH is not reduced to any significant extent by the mere diffusion into the interlayer, but the pH drops quite rapidly once the alkali diffuses through the spacer layer.

As disclosed in last-mentioned copending application Ser. No. 234,864, the presence of an inert spacer layer was found to be effective in evening out the various reaction rates over a wide range of temperatures, for example, by preventing premature pH reduction when the inhibition is effected at temperatures above room temperature, for example, at 95° to 100° F. By providing an inert spacer layer, that application discloses that the rate at which alkali is available for capture in the polymeric acid layer becomes a function of the alkali diffusion rates.

However, as disclosed in copending U.S. application Ser. No. 447,100, filed Apr. 9, 1965, now abandoned in the names of Leonard C. Farney, Howard G. Rogers and Richard W. Young, preferably the aforementioned rate at which the cations of the alkaline processing composition, i.e., alkali ions, are available for capture in the polymeric acid layer becomes a function of the alkali diffusion rates in order to provide diffusion transfer color processes relatively independent of positive transfer image variations over an extended range of ambient temperatures.

Slightly, it is there stated to have been found that the diffusion rate of alkali through a permeable inert polymeric spacer layer increases with increased processing temperature to the extent, for example, that at relatively high transfer processing temperatures, that is, transfer processing temperatures above approximately 80° F., a premature decrease in the transfer processing composition occurs due, at least in part, to the rapid diffusion of alkali from the dye transfer environment and its subsequent neutralization upon contact with the polymeric acid layer. This was stated to be especially true of alkali transferring an inert spacer layer possessing permeability to alkali to the extent of the effective curing range of optimum transfer processing. Conversely, at temperatures below the optimum transfer processing range, for example, temperatures below approximately 40° F., the last-mentioned inert spacer layer was disclosed to provide an effective diffusion barrier timely preventing effective transfer of the inert spacer layer having temperature depressed diffusion rates and to result in maintenance of the transfer processing environment's high pH for such an extended time interval as to facilitate formation of transfer image stain and its resultant degradation of the positive transfer images' color definition.

It is further stated in the last-mentioned copending application Ser. No. 447,100 to have been found, however, that if the inert spacer layer of the print-receiving element is replaced by a spacer layer which comprises a permeable polymeric layer exhibiting permeability inversely dependent on temperature, that is, a polymeric film-forming material which exhibits decreasing permeability to solubilized alkali derived cations such as alkali metal and quaternary ammonium ions under conditions of increasing temperature results in less transfer image defects resultant from the aforementioned pH maintenance and/or premature pH reduction are obviated.

As examples of polymers which were disclosed to exhibit inverse temperature-dependent permeability to alkali, mention may be made of: hydroxypropyl polyvinyl alcohol, polyvinyl methyl ether, polyethylene oxide, polyvinyl oxazolidinone, hydroxypropyl methyl cellulose, isopropl cellulose, partial acetics of polyvinyl alcohol such as partial polyvinyl butyral, partial polyvinyl formal, partial polyvinyl acetal, partial polyvinyl propional, and the like.

The last-mentioned specified acetics of polyvinyl were stated to generally comprise saturated aliphatic hydrocarbon chains of a molecular weight of at least 1000, preferably of about 1000 to 50,000, possessing a degree of acetalation within about 10 to 30%, 10 to 30%, 20 to 80%, and 10% to 40% of the polyvinyl alcohol's theoretical polymeric hydroxyl groups, respectively, and including mixed acetics where desired.

Where desired, a mixture of the polymers is to be employed, for example, a mixture of hydroxypropyl methyl cellulose and partial polyvinyl butyral.

As examples of materials, for use as the image-receiving layer, may be made of solution dyeable polymers such as nylons as, for example, N-methoxy-methyl polyhexamethylene adipamide; partially hydrolyzed polyvinyl acetate; polyvinyl alcohol with or without plasticizers; cellulose acetate with filler as, for example, one-half cellulose acetate and one-half oleic acid gelatin; and other materials of a similar nature. Preferred materials comprise polyvinyl alcohol or gelatin containing a dye mordant such as poly-4-vinylpyridine, as disclosed in U.S. Patent Nos. 3,148,611, issued Sept. 8, 1964.

As disclosed in the previously cited patents, the liquid processing composition comprising transfer processing temperatures, color diffusion transfer processes comprises at least an aqueous solution of an alkaline material, for example, diethylamine, sodium hydroxide or sodium carbonate and the like, and preferably possessing a pH in excess of 12, and most preferably includes a viscosity-increasing compound constituting a film-forming material of the type which, when the composition is spread and dried, forms a relatively firm and relatively stable film. The preferred film-forming materials disclosed comprise high molecular weight polymers such as polymeric, water-soluble ethers which are inert to an alkaline solution such as, for example, hydroxyethyl cellulose or sodium carboxymethyl cellulose. Additionally, film-forming materials or thickening agents whose ability to increase viscosity is substantially unaffected if left in solution for a long period of time are also disclosed to be capable of utilization. As stated, the film-forming material is preferably contained in the photographic composition in quantities as to impart to the composition a viscosity in excess of 100 cps. at a temperature of approximately 24° C. and preferably in the order of 100,000 cps. to 200,000 cps. at that temperature.

In accordance with the aforementioned U.S. Patent No. 2,983,606, an imaging receiving layer of the type disclosed in that patent need not be separated from its superposed
contact with the photosensitive element, subsequent to transfer image formation, if the image-receiving element is transparent and a processing composition containing a substrate is applied to the image-receiving element, the dried image remains opaque whereas the image-developing composition is spread between the image-receiving layer and the silver halide emulsion or emulsions.

However, it has been found, if the image-receiving element is maintained in contact with the photosensitive element, subsequent to dye developer transfer image formation, that the image-developing composition includes a transfer image composition, necessarily having a pH at which dye developer is reduced, for example, in reduced form, diffuses to form the dye transfer image, intermediate the elements, the transfer image thus formed is unstable over an extended period of time.

The dye image instability is due, at least in part to the presence of what is, in general, a relatively high pH alkaline composition in intimate contact with the dye or dyes forming the image. This contact itself provides instability to the molecular structure of dye by, for example, catalyzing degradation and undesirable structural shifts effecting the spectral absorption characteristics of the image dye. In addition, the presence of an alkaline composition, possessing a pH at which the dye, for example, in reduced form, diffuses, also provides an integral dynamic system wherein oxidized dye, immobilized in areas of the photosensitive element, as a function of its development, with the passage of time attains a state of equilibrium, in such areas, of the alkaline composition, in favor of additional formation of reduced dye. As a function of the efficiency of the image-receiving layer, as a dye sink, such nonimage-wise dyeing of the image-carrying layer still further balances the equilibrium in favor of the additional formation of dye as reduced, diffusible form. Under such circumstances, the transfer image definition, originally carried by the image-receiving layer, will suffer a continuous decrease in the density between the image's maximum and minimum densities and may, ultimately, result in the image-receiving element's loss of all semblance of image definition, merely becoming a polymeric stratum carrying a relatively uniform overall dyeing.

Any attempt to decrease the dye sink capacity of the image-carrying layer, for example, by reducing its mordant capacity, in order to alleviate, at least to an extent, the reaction of the image carrying layer with the image-developing composition, which may be the opposite of the alkaline composition, will result in a continuous decrease in the density between the image's maximum and minimum densities and may, ultimately, result in the image-receiving element's loss of all semblance of image definition.

As previously discussed, my co-inventors and I, in U.S. Patent No. 3,415,645, disclosed certain image-receiving elements particularly adapted for employment in diffusion transfer color processes, wherein the image-receiving element is separated from contact with a superposed photosensitive element, subsequent to substantially complete image formation, and which provide particular protection of transfer images from the effects of aerial oxidation of dyes forming the color transfer image.

It has now unexpectedly been discovered that the problems inherent in fabricating a film unit of the type wherein the image-receiving element, the alkaline processing composition and the photosensitive element are maintained in contiguous contact subsequent to dye transfer image formation, for example, a film unit of the type described hereinafter with reference to aforementioned U.S. Patent No. 3,415,645, may be simply and effectively obviated by fabrication of the film unit in accordance with the physical parameters hereinafter specifically set forth.

Specifically, it has now been unexpectedly discovered that an integral photographic film unit particularly adapted for the production of a dye transfer image, by a color diffusion transfer process described hereinafter will be constructed to include a photosensitive element comprising a laminate having, in sequence, as essential layers, a first dimensionally stable transparent layer; a photosensitive silver halide emulsion layer having associated therewith dye image-providing material which is soluble and diffusible, in alkali, at first pH; an alkaline solution permeable opaque layer; an alkaline solution permeable polymeric layer dyeable by the dye image-providing material; an alkaline solution permeable polymeric acid layer containing sufficient acidifying groups to effect reduction, subsequent to substantial transfer dye image formation, of a selected processing solution having the first pH to a second pH at which said dye image-providing material is insoluble and nondiffusible; and a second dimensionally stable transparent layer. In combination with the laminate, a rupturable container retaining an aqueous alkaline processing composition having the first pH is fixedly extended and extends transverse a leading edge of the laminate whereby to effect unidirectional discharge of the container's contents between the alkaline solution permeable and dyeable polymeric layer and the photosensitive silver halide emulsion layer next adjacent thereto, upon application of compressive force to the container.

The film unit preferably additionally contains an opaque sheet fixedly positioned extending transverse a leading edge of the unit and adapted to be superposed on and coextensive with the surface of the first transparent layer opposite the silver halide emulsion layer next adjacent thereto.

In a preferred embodiment of the present invention, the film unit is specifically adapted to provide for the production of a multicolor dye transfer image and the photosensitive laminate comprises, in order of essential image layers, the dimensionally stable transparent layer; at least two selectively sensitized silver halide emulsion strata each having dye image-providing materials of predetermined color associated therewith which are soluble and diffusible, in alkali, at a first pH; the alkaline solution permeable polymeric layer; the alkaline solution permeable polymeric acid layer containing sufficient acidifying groups to effect reduction, subsequent to substantial multicolor transfer dye image formation, of a processing composition having the first pH to a second pH, at which the dye image-providing material is insoluble and nondiffusible; and the dimensionally stable transparent layer.

The silver halide emulsions comprising the multicolor photosensitive laminate preferably possess predominant spectral sensitivity to separate regions of the spectrum and each has a regulated processing zone, wherein the silver halide developing agent and is, most preferably, substantially soluble in the reduced form only at the first pH, possessing a spectral absorption range substantially complementary to the predominant sensitivity range of its associated emulsion subsequent to exposure of the film unit. Specifically, preferably, the film unit will exhibit major spectral absorption outside of the primary region of the spectrum to which the associated silver halide emulsion is sensitive and a spectral transmission substantially complementary to that absorption, during exposure of the emulsion, and major spectral absorption within the spectral range to which the associated silver halide emulsion is sensitive and spectral transmission sub-
stantially complementary to said absorption, subsequent to exposure of said emulsion, for example, of the type disclosed in copending U.S. application Ser. No. 418,628, filed Dec. 16, 1964, now U.S. Patent No. 3,307,947 in the names of Elbert M. Idelson and Howard G. Rogers.

In the preferred embodiment, each of the emulsion strata, and its associated dye, is separated from the remaining emulsion strata, by separate alkaline solution permeable polymeric interlayers and the dyeable polymeric layer is separated from the polymeric acid layer by an alkaline solution permeable polymeric spacer layer, most preferably a polymeric spacer layer having decreasing permeability to alkaline solution with increasing temperature.

In such preferred embodiment of the invention, the silver halide emulsion comprises photosensitive silver halide dispersed in gelatin and is about 0.6 to 6 microns in thickness; the dye itself is dispersed in an aqueous alkaline solution polymeric binder, preferably gelatin, as a separate layer about 1 to 7 microns in thickness; the alkaline solution permeable polymeric interlayers, preferably gelatin, are about 1 to 5 microns in thickness; the alkaline solution permeable and dyeable polymeric layer is transparent and about 0.25 to 0.41 mil in thickness; the alkaline solution polymeric spacer layer intermediate the dyeable polymeric layer and the polymeric acid layer is transparent and about 0.15 to 0.7 mil in thickness; the alkaline solution permeable polymeric acid layer is transparent and about 0.25 to 1.5 mils in thickness; and each of the dimensionally stable transparent layers are alkaline solution permeable and about 2 to 6 mils in thickness. It will be specifically recognized that the relative dimensions recited above may be appropriately modified, in accordance with the desires of the operator, with respect to the specific product to be ultimately prepared.

In view of the fact that the preferred dye image-providing materials comprise dyes which are silver halide developer agents, as stated above, the present invention will be further described hereinafter in terms of such dyes without limitation of the invention to the illustrative dyes.

In the preferred embodiment of the present invention's film unit for the production of a multicolor transfer image, the respective silver halide/dye developer units of the photosensitive element will be in the form of a tripack configuration which will ordinarily comprise a yellow dye developer/blue-sensitive emulsion unit contiguous a dimensionally stable transparent layer, the cyan dye developer/red-sensitive emulsion unit most distant from the same transparent layer and developer/green-sensitive emulsion unit intermediate those units, recognizing that the relative order of such units may be varied in accordance with the desires of the operator.

Employment of the detailed film unit of the present invention, according to the hereinafter described color diffusion transfer photographic process, specifically, provides for the production of a color transfer image accomplished by a diffusion transfer process which includes in process adjustment of the environmental pH of the film unit, from a pH at which transfer processing is operative to a pH at which dye transfer is inoperative, subsequent to substantial transfer image formation, in order to obviate the previously discussed disadvantages of the prior art products and processes. The color transfer image is thus obtained and maintained irrespective of the fact that the film unit comprises an integral laminate unit during exposure processing and subsequent viewing, and storage of the unit. Accordingly, by means of the present invention, color transfer images and specifically multicolor images may be obtained, employing an integral laminate film unit, which exhibit desired maximum and minimum dye densities; yellow, magenta and cyan dye saturation; red, green and blue hues; and color separation, over an extended period of time, as compared with prior art disclosed integral laminates. These unexpected advantages are in addition to the manufacturing advantages, obtained by reason of the present invention's integral color transfer film unit and which will be readily apparent from examination of the unit's parameters, that is, for example, advantages in more efficient utilization of manufacturing materials and components, enhanced simplicity of film manufacture and camera design and construction, and more simplified and effectively controlled customer utilization of the unit.

Reference is now made to FIGS. 1 through 7 of the drawings wherein there is illustrated a preferred film unit of the present invention and wherein like numbers, appearing in the various figures, refer to like components. As illustrated in the drawings, FIGURE 1 sets forth a perspective view of the film unit, designated 10, and each of FIGS. 2 through 7 illustrate diagrammatic cross-sectional views of film unit 10, along the stated section lines 2--2, 3--3, 5--5, and 7--7, during the various depicted stages in the performance of a photographic diffusion transfer process as detailed hereinafter.

Film unit 10 comprises rupturable container 11, retaining, prior to processing, aqueous alkaline solution 12, photosensitive laminate 13 including, in order, dimensionally stable transparent layer 14, preferably an acetic radiation transmissive flexible sheet material; blue-sensitive silver halide emulsion layer 15 containing yellow dye developer; interlayer 16; green-sensitive silver halide emulsion layer 17 containing magenta dye developer; interlayer 18; red-sensitive silver halide emulsion layer 19 containing cyan dye developer; opaque layer 20; image-receiving layer 21; spacer layer 22; neutralizing layer 23; and dimensionally stable transparent layer 24, also preferably an acetic radiation transmissive flexible sheet material, and opaque cover sheet 25, preferably an acetic radiation opaque flexible sheet material.

The structural integrity of laminate 13 may be maintained, at least in part, by the adhesive capacity exhibited between the various layers comprising the laminate at their opposed surfaces. However, the adhesive capacity exhibited at an interface intermediate image-receiving layer 21 and the silver halide emulsion layer next adjacent thereto, for example, between the opposed surface of opaque layer 20 and either of the layers next adjacent thereto, that is, red-sensitive silver halide emulsion layer 19 or image-receiving layer 21, as illustrated in FIGS. 2 through 7, should be less than that exhibited at the interface between the opposed surfaces of the remainder of the layers forming the laminate, in order to facilitate the distribution of processing solution 12 intermediate the stated image-receiving layer 21 and the silver halide emulsion layer next adjacent thereto. The laminate's structural integrity may also be enhanced or provided, in whole or in part, by providing a binding member extending around, for example, the edges of laminate 13, and maintaining the layers comprising the laminate intact, except at an interface between layers 19 and 21 during distribution of alkaline solution 12 intermediate those layers. As illustrated in the figures, the binding member may comprise a pressure-sensitive tape 26 securing and/or maintaining the layers of laminate 13 together at its respective edges. Tape 26 will also act to maintain processing solution 12 intermediate image-receiving layer 21 and silver halide emulsion layer next adjacent thereto, upon application of compressive pressure to pod 11 and distribution of its contents intermediate the stated layers. Under such circumstances, binder tape 26 will act to prevent leakage of processing composition from the film unit's laminate during and subsequent to photographic processing.
two walls 27 which are sealed to one another along their longitudinal and end margins to form a cavity in the container 12 of the container by the application of compressive pressure to walls 27 of the container.

As illustrated in FIGS. 1, 2, and 4, container 11 is fixedly positioned and extends transverse a leading edge of photosensitive laminate 13 whereby to effect unidirectional discharge of the container's contents 12 between image-receiving layer 21 and the silver halide emulsion layer 24. Opaque sheet 25 is sealed to laminate 13 by extension 30 of tape 26 extending over a portion of one wall 27 of the container, in combination with a separate retaining member such as illustrated retaining tape 31 extending over a portion of the other wall 27 of the container and a portion of laminate 13's surface generally equal in area to about that covered by tape 26.

Opaque sheet 25, as illustrated in FIGS. 1, 2 and 4, is fixedly attached to laminate 13 extending the leading edge of film unit 10 and is adapted to be superposed, subsequent to exposure of photosensitive laminate 13, on and coextensive with exposure surface 32 of dimensional transparent layer 14, in order to occlude actinic radiation from effecting further exposure of photosensitive silver halide emulsion layers 15, 17 and 19, during processing in the presence of such radiation.

As illustrated in FIG. 6, extension flap 30 of tape 26 is preferably of such area and dimensions that when, for example, a manual separation of container 11 and tape 31, subsequent to distribution of processing composition 12, from the film unit 10, flap 30 may be folded over the edge of laminate 13, previously covered by tape 31, in order to facilitate maintenance of the laminate's structural integrity, for example, during the flexations inevitable in storage and use of the processed film unit, and to provide a suitable mask or frame, for viewing of the transfer image through the picture viewing area of transparent layer 24.

In general, the opacity of opaque layer 25 will preferably be sufficient to prevent further exposure of the film unit's silver halide emulsion or emulsions, that is, the opacity of the layer will be such as to prevent actinic radiation incident on opaque layer 25, during processing of the unit in the presence of radiation actinic to the emulsion or emulsions. Accordingly, the film unit may be processed, subsequent to exposure, in the absence of such radiation, in view of the fact that the silver halide emulsion or emulsions of the laminate are appropriately protected by incident radiation, at one major surface by the opaque layer 20 composition and at the remaining major surface by the dimensionally stable opaque sheet 25 as described hereinafter. If the illustrated binder tapes are opaque, edge leakage of actinic radiation incident on the emulsion or emulsions will also be prevented. The selected opaque layer 25, however, should be one providing a background suitable for viewing the dye developer transfer image formed in the dycable polymeric layer. In general, while substantially any alkali solution may be used as the opaque layer, it is preferred that a layer be selected that will not interfere with the color integrity of the dye transfer image, as viewed by the observer, and, most preferably, a layer which is aesthetically pleasing to the viewer and does not provide background noise signaling, or detracting from, the information content of the image. Particularly desirable opaque layers will be those providing a white background for viewing the transfer image, and specifically those adapted to be employed to provide background for photographic prints and, especially, those layers possessing the optical properties desired for reflection of incident radiation.

The opaque layer may comprise substantially any opacifying agent compatible with the photographic system, such as, for example, barium titanate, titanium oxide, barium stearate, silver flake, silicates, alumina, zirconium oxide, zirconium acetyl acetate, sodium zirconium sulfate, kaolin, mica, and the like, which may be distributed in an alkaline solution permeable polymeric matrix or binder such as, for example, gelatin, polyvinyl alcohol, or equivalent.

A particularly preferred opaque layer comprises titanium dioxide due to its highly effective reflection properties. In general, based upon percent titanium dioxide (weight/volume), a coating composition containing about 4, 10, 20, 30, and 40% titanium dioxide will provide a percent reflectance of about 58, 76, 85, 90 and 99%, respectively. In the most preferred embodiments, the percent reflectance desired will be in the order of about 85%.

Where it may be desired to increase the opacifying capacity of a layer containing, for example, titanium dioxide, beyond that ordinarily obtained, it may also be desirable to provide the layer with an additional opacifying agent such as carbon black, for example, in a concentration of about 1 part carbon black to 100 to 500 parts titanium dioxide.

In the performance of a diffusion transfer multicolor process employing film unit 10, the unit is exposed to radiation, actinic to photosensitive laminate 13, incident on the laminate's exposure surface 32, as illustrated in FIG. 2.

Subsequent to exposure, as illustrated by FIGS. 2 and 4, film unit 10 is processed by being passed through opposed suitably gapped rolls 33 in order to apply compressive pressure to the film unit's silver halide emulsion or emulsions, and to effect destruction of longitudinal seal 28 and distribution of alkaline processing composition 12, having a pH at which the cyan, magenta and yellow dye developers are soluble and diffusible, intermediate opaque layer 25 and red-sensitive silver halide emulsion layer 19, and to effect superposition of opaque sheet 25 on and coextensive with exposure surface 32 of dimensionally stable transparent layer 14.

Alkaline processing solution 12 permeates emulsion layers 15, 17 and 19 to initiate development of the latent images contained in the respective emulsions. The cyan, magenta and yellow dye developers, of layers 15, 17 and 19, are immobilized, as a function of the development of their respective associated silver halide emulsions, preferably substantially as a result of their conversion from the reduced form to their relatively insoluble and non-diffusible oxidized form, thereby providing imagewise distribution of mobile, soluble and diffusible cyan, magenta and yellow dye developer, as a function of the point-to-point degree of their associated emulsions' exposure. At least part of the imagewise distributions of mobile cyan, magenta and yellow dye developer transfers, by diffusion, to aqueous alkaline solution permeable polymeric layer 21, to provide background reflecting photographic prints, and to permeable polymeric layer 23 whereby alkaline solution 12 is employed. The developed surface of opaque layer 25 is then exposed to a pH at which the cyan, magenta and yellow dye developers, in the reduced form, are insoluble and non-diffusible, to provide thereby a stable multicolor dye transfer image viewable through dimensionally stable transparent layer 24.

Opaque sheet 25 may be maintained in superposed relationship with exposure surface 32 of transparent layer.
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During processing, by any suitable fastening or retaining means such as, for example, a pressure-sensitive contact adhesive disposed on contact surface 35, exposure surface 32 and/or the contact surface of binding tape 26, mechanical interengaging members, and the like, adapted to maintain the respective elements in contiguous relationship.

Subsequent to distribution of processing solution 12, container 11 may be manually dissociated from the remainder of the film unit, as described above. In addition, subsequent to substantial transfer image formation, opaque sheet 25 may be manually dissociated, such as, by stripping when the stated surface contact is, at least in part, maintained. By, for example, a pressure-sensitive contact adhesive retained on surface 34 of sheet 25 such as to engage the portions of binding tape 26 extending over the areas of transparent layer's 14 exposure surface 32, to provide the product illustrated in FIG. 6.

As previously stated, the multicolor dye transfer image is viewable through dimensionally stable transparent layer 24 both during and subsequent to transfer image formation. In addition, subsequent to dissociation of opaque sheet 25 from the remainder of the film unit, a multicolor dye image, negative with respect to the transfer image, may be viewed through dimensionally stable transparent layer 14.

The pH of the alkaline processing solution initially employed must be an alkaline pH at which the dye developers employed are soluble and diffusible. Although it has been found that the specific pH to be employed may be readily determined empirically for any dye developer, or group of dye developers, most particularly desirable dye developers are soluble at pH's above 9 and relatively insoluble at pH's below 9. In reduced form, and relatively insoluble at substantially any alkaline pH, in oxidized form, and the system can be readily balanced accordingly for such dye developers. In addition, although as previously noted, the processing composition, in the preferred embodiment, will include the stated film-forming viscosity-increasing agent, or agents, to facilitate spreading of the composition and to provide maintenance of the spread composition as a structurally stable layer of the laminate, subsequent to distribution, it is not necessary that such agent(s) be employed by the composition. In the latter instance, however, it will be preferred that the concentration of solvent, that is, water, etc., comprising the composition be the minimum amount necessary to conduct the desired transfer process, in order not to adversely affect the structural integrity of the laminate and that layers formed by the laminate can readily accommodate and dissipate the solvent throughout during processing and drying without effecting undesirable dimensional changes in the layers forming the laminate. It will be noted that the liquid processing composition employed may contain an auxiliary or accelerating developing agent, such as p-methylnaphthenophenol, 2,4-diaminophenol, p-benzylaminophenol, hydroquinone, toluhydroquinone, phenylhydroquinone, 4'-methylphenylhydroquinone, etc. It is also contemplated to employ a plurality of auxiliary or accelerating developing agents, such as a 3-pyrazolodine developing agent and a benzimidazolodine developing agent, as disclosed in U.S. Patent No. 3,039,669, issued June 19, 1962. As examples of suitable combinations of auxiliary developing agents, mention may be made of 1-phenyl-3-pyrazolodine in combination with p-benzilaminophenol and 1-phenyl-3-pyrazolidone in combination with 2,5-bis-ethylidino-hydroquinone. Such auxiliary developing agents may be employed in the liquid processing composition or they may be initially incorporated, at least in part, in any one or more of the silver halide emulsion strata, the strata containing the dye developers, the interlayers, the overcoat layer, the image-receiving layer, or in any other auxiliary layer, or layers, of the film unit. It may be noted that at least a portion of the dye developers oxidized during development may be oxidized and immobilized as a result of a reaction, e.g., an energy-transfer reaction, with the oxidation product of an oxidized auxiliary developing agent, the latter developing agent being oxidized by the development of exposed silver halide. Such a reaction of oxidized developing agent with unoxidized developer would regenerate the auxiliary developing agent for further reaction with the exposed silver halide.

In addition, development may be effected in the presence of an onium compound, particularly a quaternary ammonium compound, in accordance with the processes disclosed in U.S. Patent No. 3,173,786, issued Mar. 16, 1965.

It will be apparent that the relative proportions of the agents of the diffusion transfer processing composition may be altered to suit the requirements of the operator. Thus, it is within the scope of this invention to modify the herein described developing compositions by the substitution of preservatives, alkalies, etc., other than those specifically mentioned, provided that the pH of the composition is initially at the pH as required. When desirable, it is also contemplated to include, in the developing composition, components such as restrainers, accelerators, etc. Similarly, the concentration of various components may be varied over a wide range and when desirable adaptable components may be disposed in the photosensitive element, prior to exposure, in a separate permeable layer of the photosensitive element and/or in the photosensitive emulsion.

The dimensionally stable layers and sheet referred to may comprise any of various types of conventional opaque and transparent rigid or flexible materials, for example, glass, paper, metal, and polymeric films of both synthetic types and those derived from naturally occurring products. Suitable materials include alkaline solution impermeable materials such as paper; polyacrylic acid, methyl and ethyl esters; vinyl chloride polymers; polyvinyl acetal; polyamides such as nylon; polyesters such as polymeric films derived from ethylene glycol terephthalic acid; and cellulose derivatives such as cellulose acetate, triacetate, nitrate, propionate, butyrate, acetate-propionate, or acetate-butyrate. It will be recognized that one or more of the designated layers may not be required where the remaining layers of the laminate are such as to provide the functions of these layers in the absence of same, for example, where the remaining layers of the laminate provide the requisite dimensional stability and radiation filtering properties.

In all examples of this specification, percentages of components are given by weight unless otherwise indicated.

An extensive compilation of specific dye developers particularly adapted for employment in photographic diffusion transfer processes is set forth in aforementioned U.S. Patent No. 2,983,606 and in the various capping U.S. applications referred to in that patent, especially in the table of U.S. applications incorporated by reference into the patent as detailed in column 27. As examples of additional U.S. patents detailing specific dye developers for photographic transfer processes, mention may also be made of U.S. Patents Nos. 2,983,606; 2,992,106; 3,076,386; 3,076,808; 3,076,820; 3,077,402; 3,126,280; 3,131,061; 3,134,762; 3,134,765; 3,135,604; 3,135,605; 3,135,606; 3,135,734; 3,141,772; 3,142,555; and the like.

As additional examples of synthetic, film-forming, permeable polymers particularly adapted to retain dispersed dye developer, mention may be made of nitrile-grafted polyethylene cellulose, as disclosed in U.S. Patent No. 2,992,106; an acylaminobenzene sulfon ester of a partial sulfobenzyl of polyvinyl alcohol, as disclosed in U.S. Patent No. 3,043,692; polymers of N-alkyl-α,ω-unsaturated carboxamides and copolymers of N-alkyl-α,ω-carboxamides with α-hydroxyalkyl-α,ω-unsaturated carboxamides, as disclosed in U.S. Patent No. 3,069,263; copolymers of vinylphthalimide and α,ω-unsaturated carboxylic acids, as...
disclosed in U.S. Patent No. 3,061,428; copolymers of N-vinylpyrrolidones and α,β-unsaturated carboxylic acids and terpolymers of N-vinylpyrrolidones, α,β-unsaturated carboxylic acids and alkyl esters of α,β-unsaturated carboxylic acids, as disclosed in U.S. Patent No. 3,044,873; copolymers of N,N-diacyl-α,β-unsaturated carboxamides with α,β-unsaturated carboxylic acids, the corresponding amides of such acids, and copolymers of N-arylyl- and N-cycloalkyl-α,β-unsaturated carboxamides with α,β-unsaturated carboxylic acids, as disclosed in U.S. Patent No. 3,069,264; and the like.

In addition to conventional techniques for the direct dispersion of a particulate solid material in a polymeric, or colloidial, matrix such as ball-milling and the like techniques, the preparation of the dye developer dispersion may also be obtained by dissolving the dye in an appropriate solvent, or mixture of solvents, and the resultant solution distributed in the polymeric binder, with optional subsequent removal of the solvent, or solvents, employed, as, for example, by vaporization where the selected sol-vent, or solvents, possesses a sufficiently low boiling point or washing where the selected solvent, or solvents, possesses sufficiently high differential solubility in the wash medium, for example, water, when measured against the solubility of the remaining composition components, and/or obtained by dissolving both the polymeric binder and dye in a common solvent.

For further detailed treatment of solvent distribution systems of the types referred to above, and for an extensive compilation of the conventional solvents traditionally employed in the art to effect distribution of photographic color-providing materials in polymeric binders, specifically for the formation component layers of photographic film units, reference may be made to U.S. Patents Nos. 2,269,158; 2,322,027; 2,304,939; 2,384,940; 2,801,171; and the like.

Although the invention has been discussed in detail throughout employing dye developers, the preferred dye image-providing materials, it will be readily recognized that other, less preferred, dye image-providing materials may be substituted in replacement of the preferred dye developers in the practice of the invention. For example, there may be employed dye image-forming materials such as those disclosed in U.S. Patents Nos. 2,647,049, issued July 28, 1953; 2,661,293, issued Dec. 1, 1953; 2,698,244, issued Jan. 28, 1954; 2,698,798, issued Jan. 4, 1955; and 2,802,735, issued Aug. 13, 1957, wherein color diffusion transfer processes are described which employ color coupling techniques comprising, at least in part, reacting one or more color developing agents and one or more color formers or coformers to provide a dye transfer image to a superposed image-receiving layer and those disclosed in U.S. Patent No. 2,774,668, issued Dec. 18, 1956, wherein color diffusion transfer processes are described which employ the image-wise differential transfer of complete dyes by the mechanisms therein described to provide a transfer dye image to a contiguous image-receiving layer.

For the production of the photosensitive gelatin silver halide emulsions employed to provide the film unit, the silver halide crystals may be prepared by reacting a water-soluble silver salt, such as silver nitrate, with at least one water-soluble halide, such as ammonium, potassium or sodium bromide, preferably together with a corresponding iodide, in aqueous solution of a precipitating agent such as a colloidal gelatin solution; digesting the dispersion at an elevated temperature, to provide increased crystal growth; washing the resultant dispersion to remove undesirable reaction products and residual water-soluble salts by chilling the dispersion, noodling the set dispersion, and washing the nodules with cold water, or alternatively, employing any of the various flocc systems, or procedures, adapted to effect removal of undesired components, for example, the procedures described in U.S. Patent Nos. 2,614,928; 2,614,929; 2,728,662; and the like; after-ripening the dispersion at an elevated temperature in combination with the addition of gelatin and various adjuncts, for example, chemical sensitizing agents of U.S. Patents Nos. 1,174,944; 1,263,499; 2,410,689; 2,597,856; 2,597,915; 2,487,830; 2,518,698; 2,521,926; and the like; all according to the traditional procedures of the art, as described in Neblette, C. B., Photography, Its Materials and Processes, 1946, pages 101-102.

Optical sensitization of the emulsion's silver halide crystals may be accomplished by contact of the emulsion composition with an effective concentration of the selected optical sensitizing dyes dissolved in an appropriate dispersing solvent such as methanol, ethanal, acetone, water, and the like; all according to the traditional procedures of the art, as described in Hammer, F. M., The Cyanine Dyes and Related Compounds.

Additional optical additives, such as coating aids, hardeners, viscosity-increasing agents, stabilizers, preservatives, and the like, for example, those set forth hereinafter, also may be incorporated in the emulsion formulation, according to the conventional procedures known in the photographic emulsion manufacturing art.

The photoreactive material of the photographic emulsion will, as previously described, preferably comprise a crystal of silver; for example, a halide such as silver chloride, silver iodide, silver bromide, or mixed silver halides such as silver chlorobromide or silver iodobromide, of varying halide ratios and varying silver concentrations.

The emulsions may include the various adjuncts, or addenda, according to the techniques disclosed in the art, such as speed-increasing compounds of the quaternary ammonium type, as described in U.S. Patents Nos. 2,271,623; 2,288,226; and 2,334,864; or of the polyethylene glycol type, as described in U.S. Patent No. 2,708,162; or of the preceding combination, as described in U.S. Patent No. 2,886,437; or the thiopropolymers, as described in U.S. Patents Nos. 3,046,129 and 3,046,134.

The emulsions may also be stabilized with the salts of the noble metals such as rhenium, rhodium, palladium, iridium and platinum, as described in U.S. Patents Nos. 2,566,245 and 2,566,263; the mercury compounds of U.S. Patents Nos. 2,728,663; 2,728,664 and 2,728,665; the triazoles of U.S. Patent No. 2,444,608; the azlindines of U.S. Patents Nos. 2,444,605; 2,444,606; 2,444,607; 2,450,397; 2,444,609; 2,713,541; 2,743,181; 2,716,062; 2,735,769; 2,756,147; 2,772,164; and those disclosed by Burr in U.S. Patents Nos. 2,886,437; and 2,350,317; the benzothiazolium compounds of U.S. Patents Nos. 2,131,038 and 2,694,716; the zinc and cadmium salts of U.S. Patent No. 2,839,405; and the mercapto compounds of U.S. Patent No. 2,819,965.

Hardeners agents such as inorganic agents providing polyvalent metallic ions, specifically polyvalent aluminum or chromium ions, for example, potash alum [K₂Al(SO₄)₃·2H₂O] and chrome alum [K₂Cr₂(SO₄)₃·2H₂O] and inorganic agents of the aldehyde type, such as formaldehyde, glyoxal, mucorol, etc.; the ketone type such as diacetyl; the quinone type; and the specific agents described in U.S. Patents Nos. 2,080,019; 2,725,294; 2,725,295; 2,725,305; 2,726,162; 2,732,316; 2,950,197; and 2,870,013, may be incorporated, where desired and compatible, in the selected coating solution compositions.

Coating solution compositions for fabricating the respective strata of the film unit may contain one or more coating aids, where desired and compatible, such as saponin; a polyethylene glycol of U.S. Patent No. 2,831,766; a polyethylene glycol ether of U.S. Patent No. 2,719,887; a taurate of U.S. Patent No. 2,739,891; a malonate of U.S. Patent No. 2,733,931; and amino acid of U.S. Patent No. 3,038,804; a sulfosuccinate of U.S. Patent No. 2,992,108; or a polyether of U.S. Patent No. 2,600,831; or a gelatin plasticizer such as glycerin;

As the binder for the respective emulsion strata, the aforementioned gelatin may be, in whole or in part, replaced with some other colloidal material such as albumin; casein; or zein; or resins such as a cellulose derivative, as described in U.S. Patents Nos. 2,327,806; 2,327,808; 2,451,472; vinyl polymers such as described in U.S. Patents Nos. 2,253,078; 2,276,322; 2,276,323; 2,281,703; 2,310,223; 2,311,058; 2,311,059; 2,414,208; 2,461,023; 2,484,456; 2,538,257; 2,579,016; 2,614,931; 2,624,674; 2,632,704; 2,642,420; 2,678,884; 2,691,582; 2,725,296; 2,753,264; and the like.

Although the preceding description of the invention has been couched in terms of the preferred photosensitive component construction wherein at least two selectively sensitized photosensitive strata are in contiguous coplanar relationship and, specifically, in terms of the preferred tripack type structure comprising a red-sensitive silver halide emulsion stratum, a green-sensitive silver halide emulsion stratum and a blue-sensitive silver halide emulsion stratum having associated therewith, respectively, a cyan dye developer, a magenta dye developer and a yellow dye developer, the photosensitive component of the film unit may comprise at least two sets of selectively sensitized minute photosensitive elements arranged in the form of a photosensitive screen wherein each of the minute photosensitive elements has associated therewith, for example, an appropriate dye developer in a reflective silver halide emulsion portion. In general, a suitable photosensitive screen will comprise minute red-sensitized emulsion elements, minute green-sensitized emulsion elements and minute blue-sensitized emulsion elements arranged in side-by-side relationship in a screen pattern and having associated therewith, respectively, a cyan, a magenta and a yellow dye developer.

The present invention also includes the employment of a black dye developer and the use of a mixture of dye developers adapted to provide a black and white transfer image, for example, the employment of dye developers of the three subtractive colors in an appropriate mixture in which the quantities of the dye developers are proportioned such that the colors combine to provide black. Wherein in the specification, the expression "positive image" has been used, this expression should not be interpreted in a restrictive sense since it is used primarily for purposes of illustration, in that it defines the image produced on the image-carrying layer as being reversed, in the positive-negative sense, with respect to the image in the photosensitive emulsion layers. As an example of an alternative meaning for "positive image," assume that the photosensitive element is exposed to actinic light through a negative transparency. In this case, the latent image in the photosensitive emulsion layers will be a positive and the dye image produced on the image-carrying layer will be a negative. The expression "positive image" is intended to cover such an image produced on the image-carrying layer.

In addition to the described essential layers, it will be recognized that the film unit may also contain one or more subcoats or layers, which, in turn, may contain one or more additives such as plasticizers, intermediate essential layers for the purpose, for example, of improving adhesion, and that any one or more of the described layers may comprise a composite of two or more strata of the same, or different, components and which may be contiguous, or separated from, each other, for example, two or more neutralizing layers or the like, one of which may be disposed intermediate the cyan dye image-forming component retaining layer and the dimensionally stable opaque layer.

Since certain changes may be made in the above product and process without departing from the scope of the invention herein involved, it is intended that all matter contained in the above description or shown in the accompanying drawings shall be interpreted as illustrative and not in a limiting sense.

What is claimed is:

1. A photographic film unit which is adapted to be processed by passing said unit between a pair of juxtaposed pressure-applying members and which comprises, in combination:

a photosensitive element comprising a composite structure containing, as essential layers, in sequence, a first dimensionally stable transparent layer, a photosensitive silver halide emulsion layer having associated therewith a dye image-forming material which is soluble and diffusible in alkaline solution permeable opaque layer, an alkaline solution permeable polymeric layer dyeable by said dye image-providing material, an alkaline solution permeable polymeric acid layer containing sufficient acidifying groups to effect reduction of a processing solution having said first pH to a second pH at which said dye image-providing material is substantially insoluble and nondiffusible, a second dimensionally stable transparent layer, and means securing said layers in fixed relationship, and

a rupturable container retaining an alkaline processing solution, having said first pH, fixedly positioned and extending transversely a leading edge of said photosensitive element to effect unidirectional discharge of the container's contents between said alkaline solution permeable and dyeable polymeric layer and the photosensitive silver halide emulsion layer next adjacent thereto upon application of compressive force to said container.

2. A photographic film unit as defined in claim 1, including a separate dimensionally stable opaque sheet fixedly positioned extending transversely a leading edge of said film unit and adapted to be superposed during processing on the surface of said first dimensionally stable transparent layer opposite said silver halide emulsion layer and wherein said opaque layer possesses sufficient opacifying capacity to prevent exposure of silver halide emulsion during processing in the presence of radiation actinic thereto.

3. A photographic film unit as defined in claim 1, wherein said dye image-providing material is a dye which is a silver halide developing agent.

4. A photographic film unit as defined in claim 3, wherein said dye is substantially soluble and diffusible in only the reduced form at said first pH and is substantially insoluble and nondiffusible in said reduced form at said second pH.

5. A photographic film unit as defined in claim 3, including an alkaline solution permeable polymeric spacer layer positioned intermediate said alkaline solution permeable and dyeable polymeric layer and said alkaline solution permeable polymeric acid layer.

6. A photographic film unit as defined in claim 5, wherein said alkaline solution permeable polymeric spacer layer possesses decreasing alkaline solution permeability with increasing temperature.

7. A photographic film unit as defined in claim 1, wherein said aqueous alkaline solution additionally contains a film-forming polymeric material.

8. A photographic film unit as defined in claim 7, wherein said film-forming polymeric material is hydroxyethyl cellulose.

9. A photographic film unit as defined in claim 1, wherein said opaque layer is actinic radiation reflective.

10. A photographic film unit as defined in claim 9, wherein said opaque layer comprises titanium dioxide dispersed in an alkaline solution permeable polymeric binder.

11. A photographic film unit as defined in claim 10, wherein said opaque layer contains carbon black.

12. A photographic film unit as defined in claim 3,
wherein said dye is dispersed in a separate alkaline solution permeable polymeric layer intermediate said associated photosensitive silver halide emulsion layer and said first dimensionally stable transparent layer.

13. A photographic film unit, as defined in claim 3, which is adapted to be processed by passing said unit between a pair of juxtaposed pressure-applying members and which comprises, in combination:
   a photosensitive element comprising a composite structure containing, as essential layers, in sequence,
   a first dimensionally stable alkaline solution impermeable transparent layer, at least two selectively sensitized silver halide emulsion layers each having a dye, which dye is a silver halide developing agent, of predetermined color associated therewith,
   each of said dyes soluble and diffusible, in alkali, at a first pH, an alkaline solution permeable opaque layer,
   an alkaline solution permeable polymeric layer dyebale by said dyes, an alkaline solution permeable transparent polymeric acid layer containing sufficient acidifying groups to effect reduction of a processing solution having said first pH to a second pH at which said dyes are substantially insoluble and nondiffusible, a second dimensionally stable alkaline solution permeable transparent polymeric layer, and means securing said layers in fixed relationship and a rupturable container retaining an aqueous alkaline processing solution, having said first pH, fixedly positioned and extending transverse a leading edge of said photosensitive element to effect unidirectional discharge of the container's contents between said alkaline solution permeable and dyebale polymeric layer and the photosensitive silver halide emulsion layer next adjacent thereto.

14. A photographic film unit as defined in claim 13, wherein each of said selectively sensitized photosensitive silver halide emulsion layers has predominant spectral sensitivity to separate regions of the spectrum and the dye associated with each of said silver halide emulsion layers possesses a spectral absorption range substantially complementary to the predominant sensitivity range of its associated emulsion layer.

15. A photographic film unit as defined in claim 13, wherein each of said silver halide emulsion layers and its associated dye is separated from the next adjacent silver halide emulsion layer and its associated dye by an alkaline solution permeable polymeric interlayer.

16. A photographic film unit as defined in claim 13, wherein at least one of said dyes is disposed in a separate alkaline solution permeable polymeric layer adjacent its associated silver halide emulsion layer intermediate said emulsion layer and said first dimensionally stable transparent layer.

17. A photographic film unit as defined in claim 13, including an alkaline solution permeable transparent polymeric spacer layer positioned intermediate said alkaline solution permeable and dyebale polymeric layer and said alkaline solution permeable polymeric acid layer.

18. A photographic film unit as defined in claim 17, wherein said alkaline solution permeable polymeric spacer layer has decreasing alkaline solution permeability with increasing temperature.

19. A photographic film unit as defined in claim 13, wherein said opaque layer is actinic radiation reflective.

20. A photographic film unit as defined in claim 19, wherein said opaque layer comprises titanium dioxide dispersed in an alkaline solution permeable polymeric binder.

21. A photographic film unit as defined in claim 20, wherein said opaque layer contains carbon black as an additional opacifying agent.

22. A photographic film unit as defined in claim 13, including a separate dimensionally stable opaque sheet fixedly positioned extending transverse a leading edge of said film unit and adapted to be superposed during processing on the surface of said first dimensionally stable transparent layer opposite said emulsion layers and wherein said opaque layer possesses sufficient opacifying capacity to prevent exposure of said silver halide emulsion layers during processing in the presence of radiation actinic thereto.

23. A photographic film unit, as defined in claim 14, which is adapted to be processed by passing said unit between a pair of juxtaposed pressure-applying members and which comprises, in combination:
   a photosensitive element including a composite structure containing, as essential layers, in sequence,
   a first dimensionally stable alkaline solution impermeable transparent layer, a blue-sensitive silver halide emulsion layer having associated therewith a yellow dye, a green-sensitive silver halide emulsion layer having associated therewith a magenta dye, a red-sensitive silver halide emulsion layer having associated therewith a cyan dye, each of said dyes, a magenta and yellow dyes being silver halide developing agents and being soluble and diffusible, in alkali, at a first pH, an alkaline solution permeable opaque layer, and alkaline solution permeable polymeric layer dyebale by said dyes, an alkaline solution permeable transparent polymeric acid layer containing sufficient acidifying groups to effect reduction of a processing solution having said first pH to a second pH at which said dyes are substantially insoluble and nondiffusible, a second dimensionally stable alkaline solution permeable transparent polymeric layer, means securing at least the side edges of said opaque layers in fixed relationship, and a rupturable container retaining an aqueous alkaline processing solution, having said first pH, fixedly positioned and extending transverse a leading edge of said photosensitive element to effect unidirectional discharge of the container's contents between said alkaline solution permeable and dyebale polymeric layer and said red-sensitive silver halide emulsion layer upon application of compressive force to said container.

24. A photographic film unit as defined in claim 23, including an alkaline solution permeable transparent polymeric spacer layer positioned intermediate said alkaline solution permeable dyeable polymeric layer and said alkaline solution permeable polymeric acid layer.

25. A photographic film unit as defined in claim 23, wherein said aqueous alkaline solution additionally contains a film-forming polymeric matrix.

26. A photographic film unit as defined in claim 23, including a separate dimensionally stable opaque sheet fixedly positioned extending transverse a leading edge of said film unit and adapted to be superposed during processing on the surface of said first dimensionally stable transparent layer opposite said emulsion layers and wherein said opaque layer possesses sufficient opacity to prevent exposure of said silver halide emulsion layers during processing in the presence of radiation actinic thereto.

27. A photographic film unit as defined in claim 26, wherein said opaque layer is actinic radiation reflective.

28. A photographic film unit as defined in claim 27, wherein said opaque layer comprises titanium dioxide dispersed in an alkaline solution permeable polymeric matrix.

29. A photographic film unit as defined in claim 28, wherein said opaque layer contains carbon black as an additional opacifying agent.

30. A photographic film unit as defined in claim 23, wherein said first pH is above 9 and said second pH is below 9.

31. A photographic film unit as defined in claim 28, wherein said opaque layer and each of said silver halide emulsion layers comprises gelatin.

32. A photographic film unit as defined in claim 31, wherein said opaque layer contains sufficient titanium di-
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oxide to provide a percent reflectance of at least about 85% of incident visible radiation.

33. A process of forming transfer images in color which comprises the steps of:
(a) exposing a photographic film unit which is adapted to be processed by passing the unit between a pair of juxtaposed pressure-applying members and which includes, in combination, a photosensitive element comprising a composite structure containing, as essential layers, in sequence, a first dimensionally stable transparent layer, a photosensitive silver halide emulsion layer having associated therewith dye image-providing material which is soluble and diffusible, in alkali, at a first pH, an alkaline solution permeable opaque layer, an alkaline solution permeable polymeric layer dyeable by said image-providing material, an alkaline solution permeable polymeric acid layer containing sufficient acylating groups to effect reduction of a processing solution having said first pH to said second pH at which said dye image-providing material is substantially insoluble and nondiffusible, a second dimensionally stable transparent layer, and means securing said layers in fixed relationship, and a rupturable container retaining an aqueous alkaline processing solution, having said first pH, fixedly positioned and extending transverse a leading edge of said photosensitive element to effect unidirectional discharge of the container's contents between said alkaline solution permeable and dyeable polymeric material and said photosensitive silver halide emulsion layer next adjacent thereto upon application of compressive force to said container, said exposure accomplished by radiation actinic to said film unit incident on said first transparent layer;
(b) applying compressive force to said rupturable container to effect unidirectional discharge of the container's aqueous alkaline processing solution between said alkaline solution permeable and dyeable polymeric layer and said photosensitive silver halide emulsion layer next adjacent thereto;
(c) effecting thereby development of said silver halide emulsion;
(d) immobilizing the dye image-providing material associated with said emulsion, as a result of said development;
(e) forming thereby an imagewise distribution of mobile dye image-providing material, as a function of the point-to-point degree of emulsion exposure;
(f) transferring, by diffusion, at least a portion of said imagewise distribution of mobile dye image-providing material to said alkaline solution permeable polymeric layer dyeable by said dye image-providing material to provide a dye image thereto in terms of said imagewise distribution;
(g) transferring, by diffusion, subsequent to substantial transfer image formation, a sufficient portion of the ions of said alkaline solution to said alkaline solution permeable polymeric acid layer to thereby reduce the alkalinity of said aqueous alkaline solution to said second pH; and
(h) maintaining said composite structure intact subsequent to development.

34. A process as defined in claim 33 including the step of separating said container from said composite structure subsequent to substantial transfer image formation.

35. A process as defined in claim 33, wherein said photographic film unit includes an alkaline solution permeable polymeric spacer layer position intermediate said alkaline solution permeable and dyeable polymeric layer and said alkaline solution permeable polymeric acid layer.

36. A process as defined in claim 35, wherein said alkaline solution permeable polymeric spacer layer possesses decreasing alkaline solution permeability with increasing temperature.

37. A process as defined in claim 33, wherein said aqueous alkaline solution additionally contains a film-forming polymeric material.

38. A process as defined in claim 33, wherein said opaque layer is actinic radiation reflective.

39. A process as defined in claim 38, wherein said opaque layer contains titanium dioxide dispersed in an alkaline solution permeable polymeric binder.

40. A process as defined in claim 39, wherein said opaque layer contains carbon black as an additional opacifying agent.

41. A process as defined in claim 33, wherein said dye image-providing material is a dye which is a silver halide developing agent.

42. A process as defined in claim 41, wherein said dye is substantially soluble and diffusible in the reduced form only at said first pH and is substantially insoluble and nondiffusible in the reduced form at said second pH.

43. A process as defined in claim 41, wherein said photographic film unit includes a separate dimensionally stable opaque sheet fixedly positioned transverse a leading edge of said unit and adapted to be superposed subsequent to exposure of said photosensitive element co-extensive with the surface of said first dimensionally stable transparent layer opposite said emulsion layers and wherein said opaque layer possesses sufficient opacity to prevent exposure of said emulsion during processing in the presence of radiation theretofore including the step of superposing said opaque sheet on said surface subsequent to said exposure of said photosensitive element.

44. A process of forming transfer images in color, as defined in claim 33, which comprises, in combination, the steps of:
(a) exposing a photographic film unit which is adapted to be processed by passing said unit between a pair of juxtaposed pressure-applying members and which includes, in combination, a photosensitive element comprising a composite structure containing, as essential layers, in sequence, a first dimensionally stable transparent layer, and means securing said layers in fixed relationship, and a rupturable container retaining an aqueous alkaline processing solution having said first pH, fixedly positioned and extending transverse a leading edge of said photosensitive element to effect unidirectional discharge of the container's contents between said alkaline solution permeable and dyeable polymeric material and said photosensitive silver halide emulsion layer next adjacent thereto upon application of compressive force to said container, said exposure accomplished by radiation actinic to said film unit incident on said first transparent layer;
(b) applying compressive force to said rupturable container to effect unidirectional discharge of the container's aqueous alkaline processing solution between said alkaline solution permeable and dyeable polymeric layer and said photosensitive silver halide emulsion layer next adjacent thereto;
(c) effecting thereby development of said silver halide emulsion;
(d) immobilizing the dye associated with each of said emulsions as a result of development;
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(e) forming an imagewise distribution of mobile dye, as a function of the point-to-point degree of emulsion exposure;

(f) transferring, by imbibition, at least a portion of each of said imagewise distributions of mobile dye to said alkaline solution permeable polymeric layer dyeable by said dyes to provide thereto a dye image;

(g) transferring, by imbibition, subsequent to substantive transfer image formation, a sufficient portion of the ions of said aqueous alkaline solution to said alkaline solution permeable polymeric acid layer to thereby reduce the alkalinity of said solution to said second pH; and

(h) maintaining the thus-formed laminate intact subsequent to processing.

45. A process as defined in claim 44, wherein each of said selectively sensitized silver halide emulsion strata has predominant spectral sensitivity to a separate region of the spectrum and, subsequent to exposure, the dye associated with each of said emulsion strata has a spectral absorption range substantially complementary to the predominant sensitivity range of its associated emulsion.

46. A process as defined in claim 45, wherein each of said silver halide emulsion strata is separated from the silver halide emulsion strata next adjacent thereto by an alkaline solution permeable polymeric spacer layer.

47. A process as defined in claim 45, wherein said photographic film unit includes a separate dimensionally stable opaque sheet fixedly positioned transverse a leading edge of said unit and adapted to be superposed subsequent to exposure of said photosensitive element coextensive with the surface of said first dimensionally stable transparent layer opposite said emulsion layers and wherein said opaque layer possesses sufficient opacity to prevent exposure of said silver halide emulsions during processing in the presence of radiation actinic thereto, said process including the step of superposing said opaque sheet on said surface subsequent to said exposure of said photosensitive element and at least a portion of said developing and said transfer image formation is accomplished while said film unit is in the presence of said radiation.

48. A process as defined in claim 47, including the step of separating at least one of said container and said opaque sheet from said laminate subsequent to substantive transfer image formation.

49. A process as defined in claim 45, wherein said opaque layer is actinic radiation reflective.

50. A process as defined in claim 49, wherein said opaque layer comprises titanium dioxide dispersed in an alkaline solution permeable polymeric layer.

51. A process as defined in claim 50, wherein said opaque layer contains carbon black as an additional opacifying agent.

52. A process of forming transfer images in color, as defined in claim 45, which comprises, in combination, the steps of:

(a) exposing a photographic film unit which is adapted to be processed by passing said unit between a pair of juxtaposed pressure-applying members and which includes, in combination, a photosensitive element comprising a composite structure containing, as essential layers, in sequence, a first dimensionally stable alkaline solution impermeable transparent layer, a blue-sensitive silver halide emulsion layer having associated therewith a yellow dye, a green-sensitive silver halide emulsion layer having associated therewith a magenta dye, a red-sensitive silver halide emulsion layer having associated therewith a cyan dye, each of said cyan, magenta and yellow dyes being silver halide developing agents and being soluble and diffusible in alkalai, at a first pH, an alkaline solution permeable opaque layer, an alkaline solution permeable polymeric acid layer dyeable by each of said dyes, an alkaline solution permeable polymeric acid layer containing sufficient acidifying groups to effect reduction of a processing solution having said first pH to a second pH at which said dyes are substantially insoluble and nondiffusible, a second dimensionally stable alkaline solution impermeable transparent layer, and means securing said layers in fixed relationship, and a rupturable container retaining said alkaline processing composition having said first pH to mask said dyes, fixedly positioned and extending transverse a leading edge of said photosensitive element to effect unidirectional discharge of the container's contents between opposed surfaces of said alkaline solution permeable and dyeable polymeric layer and said red-sensitive silver halide emulsion layer upon application of compressive force to said container, said exposure effected by actinic radiation incident on said first dimensionally stable transparent layer;

(b) applying compressive force to said rupturable container to effect unidirectional discharge of the container's alkaline solution permeable and dyeable polymeric layer and said photosensitive silver halide emulsion layer next adjacent thereto;

(c) effecting thereby development of the latent image contained in each of said silver halide emulsions;

(d) immobilizing said yellow, magenta and cyan dye, as a result of development of their respective associated silver halide emulsions;

(e) forming thereby an imagewise distribution of mobile yellow, magenta and cyan dye, as a function of the point-to-point degree of exposure of their respective associated silver halide emulsions;

(f) transferring, by diffusion, at least a portion of each of said imagewise distributions of mobile dye to said alkaline solution permeable polymeric layer dyeable by said dyes to provide thereto a multicolor dye image;

(g) transferring, by diffusion, subsequent to substantive transfer image formation, a sufficient portion of the ions of said aqueous alkaline solution to said alkaline solution permeable polymeric acid layer to thereby reduce the alkalinity of said solution to said second pH; and

(h) maintaining said composite structure intact subsequent to processing.

53. A process as defined in claim 52 including the step of separating said container from said composite structure subsequent to substantive transfer image formation.

54. A process as defined in claim 52, wherein each of said alkaline solution permeable polymeric layers containing said dyes and each of said silver halide emulsion layers comprise gelatin.

55. A process as defined in claim 52, wherein said first pH is above 9 and said second pH is below 9.

56. A process as defined in claim 52, wherein said aqueous alkaline solution additionally contains a film-forming polymeric material.

57. A process as defined in claim 52, wherein said opaque layer is actinic radiation reflective.

58. A process as defined in claim 57, wherein said opaque layer comprises titanium dioxide dispersed in an alkaline solution permeable polymeric binder.

59. A process as defined in claim 58, wherein said opaque layer contains carbon black as an additional opacifying agent.

60. A laminate which comprises, as essential layers, in sequence, a first dimensionally stable transparent layer, a photosensitive silver halide emulsion layer having associated therewith a dye image-forming material which is soluble and diffusible, in alkalai, at a first pH, an alkaline solution permeable opaque layer, an alkaline solution permeable polymeric acid layer dyeable by said dye image-providing material, an alkaline solution permeable polymeric layer containing sufficient acidifying groups to effect reduction of a selected processing solution having a first
pH to a second pH at which said dye image-providing material is insoluble and nondiffusible, and a second dimensionally stable transparent layer, wherein the adhesion exhibited intermediate said dyeable polymeric layer and the silver halide emulsion layer next adjacent thereto is less than the adhesion at the remaining interfaces of the laminate.

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NORMAN G. TARCHIN, Primary Examiner.
A. T. SURO-PICO, Assistant Examiner.

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