REPUBLIC OF SOUTH AFRICA PATENTS ACT, 1978

PUBLICATION PARTICULARS AND ABSTRACT

(Section 32(3)(a) - Regulation 22(1)(g) and 31)

21 0		LODGING DATE			
•	2006/05581	22	21 JAN 2005	43 174-09-2007	
INTERNATIONAL CLASSIFIC ATION NOT FOR PUBLICATION					
51	D2 1H		CLASSIFIED B	r: WIPO	
FULL NAMES OF APPLICANT					
71	NALCO COMPANY				
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EARLIEST PRIORITY CLAIMED					
	COUNTRY		NUMBER	DATE	
33	US	31	10/764,935	32 26 JAN 2004	
TITLE OF INVENTION					
54	METHOD OF USIENG ALDEHYDE-FUNCTIONALIZED POLYMERS TO ENHANCE PAPER MACHINE DEWATERING				
57	ABSTRACT (NOT MOTRE THAT 150 V	VORDS)	NU	MBER OF SHEETS 33	

If no classification is finished, Forn— P.9 should accompany this form. The figure of the drawing to which—the abstract refers is attached.

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ABSTRACT

A method of enhancing the dewatering of a paper shaeet on a paper machine comprising adding to the paper sheet about 0.05 lb/ton to about 15 lb/ton, based on dry fiber, of one or more aldehyde functionalized polymers comprising amino or amido groups wherein at least about 15 mole percent of the amino or amido groups are functionalized by reacting with one or more aldehydes and wherein the aldeh yde functionalized polymers have a molecular weight of at least about 100,000.

METHOD OF USING ALDEHYD E-FUNCTIONALIZED POLYMERS TO ENHANCE PAPER IMACHINE DEWATERING

TECHNICAL FIELD

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This is a method of enhancing praper machine dewatering using aldehydefunctionalized polymers having a specific level of functionalization.

BACKGROUND OF THE INVENTION

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Papermaking comprises taking a slurry of papermaking raw materials at a consistency (weight percent solids) in the range 0.1 to 1.0 weight percent and dewatering it to form a sheet with a final consistency of about 95 weight percent. Paper machines accomplish this dewate ring through a series of different processes which include from the beginning to end: 1) gravity or inertial dewatering (early forming section of the machine); 2) vacuum dewatering (late forming section of the machine); 3) press dewatering (press section of the machine); and 4) thermally evaporating the water (dryer section of the machine). The cost of dewatering increases in going from 1 to 4, which makes it advantageous to remove as much water as possible in the earlier stages. The rate of paper production or, equivalently, the machine speed is dictated by the rate at which the water can be removed, and consequently, any chemical treatment which can increase the rate of water removal has value for the papermaker. Many grades of paper require the use of retention aid chemicals for their manufacture in order to retain the fine particles found in the raw materials used to make the paper. It is well known in the paper industry that these retention aids can also enhance the rate of gravity, inertial, and vacuum dewatering or drainage, as it is often called. Such retention chemicals include the well known flocculants, coagulants, and microparticles used in the industry. Existing laboratory free and vacuum drainage tests can reactily identify the drainage effects of these retention aid chemicals.

The production rate for the vast majority of paper machines is limited by the drying capacity of the machine's dryer section. Consequently, the consistency of the paper sheet leaving the press section and going into the dryer section is most often

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critical in determining the paper machine speed or production rate. The effects of chemical additives on press dewatering are unclear with little information available on this topic. The effect of retention aid chemicals on press dewatering is often reported to be detrimental as a cornsequence of the decreased consistency entering the press as a result of increased water retention or reduction in press efficiency resulting from a loss in sheet formation. Both these factors arise from the flocculation of the paper making particles by the retention chemicals. Because the consistency of the sheet leaving the press section is most often the most critical factor in determining machine speed, any treatment capable of increasing this consistency would obviously be highly describle. Currently, no chemical treatments are known to be marketed as commercial press dewatering aids, although anecdotal reports suggest that some polymers can favorably effect out going press consistency. Accordingly, there is an ongoing need to develop compositions having effective press dewatering activity.

Glyoxylated poly-vinylamides prepared from glyoxal and polyvinylami_de in a mole ratio of 0.1 to 0.2 are disclosed as wet strength resins in U.S. Pat. No. 3, ..556,932.

Low molecular weight glyoxylated cationic polyacrylamides prepared from glyoxal and cationic polyvinylamide in a ratio of 0.1-0.5:1 are disclosed as termporary wet strength resins in U. S. Patent No. 4,605,702.

A method of imparting strength to paper by adding to a pulp slurry a maximal resin comprising an ami-nopolyamide-epichlorohydrin resin and a glyoxylated acrylamide-dimethyl diallyl ammonium chloride resin prepared from glyoxal and acrylamide-dimethyl diallyl ammonium chloride copolymer in a molar ratio of about 2-0.5:1 is disclosed in U.S. Patent No. 5,674,362.

25 SUMMARY OF THE INVENTION

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This invention is a method of enhancing the dewatering of a paper sheet on a paper machine comprising adding to the paper sheet about 0.05 lb/ton to about 15 lb/ton, based on dry fiber, of one or more aldehyde functionalized polymers comprising amino or amido groups wherein at least about 15 mole percent of the amino or amido groups are functionalized by reacting with one or more aldehydes and wherein the

aldehyde functionalized polymers have a weight average molecular weight of at least about 100,000 g/mole.

DETAILED DESCRIPTION OF THE UNVENTION

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"Acrylamide monomer" means a monomer of formula

$$R_1 O = C - CNHR_2$$

wherein R₁ is H or C₁-C₄ alkyl and R₂ \(\bar{a}\)s H, C₁-C₄ alkyl, aryl or arylalkyl. Preferred acrylamide monomers are acrylamide a.nd methacrylamide. Acrylamide is more preferred.

"Aldehyde" means a compound containing one or more aldehyde (-CHO) groups, where the aldehyde groups are capable of reacting with the amino or amido groups of a polymer comprising amino or amido groups as described herein.

Representative aldehydes include form aldehyde, paraformaldehyde, glutaraldehyde, glyoxal, and the like. Glyoxal is preferred.

"Alkyl" means a monovalent group derived from a straight or branched chain saturated hydrocarbon by the removal of a single hydrogen atom. Representative alkyl groups include methyl, ethyl, n- and is o-propyl, cetyl, and the like.

"Alkylene" means a divalent group derived from a straight or branched chain saturated hydrocarbon by the removal of two hydrogen atoms. Representative alkylene groups include methylene, ethylene, propylene, and the like.

"Amido group" means a group of formula -C(O)NHY₁ where Y₁ is selected from H, alkyl, aryl and arylalkyl.

"Amino group" means a groups of formula -NHY2 where Y2 is selected from H, alkyl, aryl and arylalkyl.

"Amphoteric" means a polymer derived from both cationic monomers and anionic monomers, and, possibly, other non-ionic monomer(s). Representative amphoteric polymers include copolymers composed of acrylic acid and

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DMAEA_•MCQ, terpolymers composed of acrylic ancid, DADMAC and acrylamide, and the like.

"Aryl" means an aromatic monocyclic or multicyclic ring system of about 6 to about 10 carbon atoms. The aryl is optionally substituted with one or more C_1 - C_{20} alkyl, alknowy or haloalkyl groups. Representative aryl groups include phenyl or naphthyl, or substituted phenyl or substituted naphthyl.

"_Arylalkyl" means an aryl-alkylene- group where aryl and alkylene are defined herein. Representative arylalkyl groups include benzyl, phenylethyl, phenylpropyl, 1-naphthy methyl, and the like. Benzyl is preferred.

"Diallyl-N,N-disubstituted ammonium halade monomer" means a monomer of formula

$(H_2C=CHCH_2)_2N^+$ $\mathbb{R}_3R_4X^-$

wherein_ R₃ and R₄ are independently C₁-C₂₀ alkyl, aryl or arylalkyl and X is an anionic counterion. Representative anionic counterions irrelude halogen, sulfate, nitrate, phosphante, and the like. A preferred anionic cournterion is halogen. Halogen is preferred. A preferred diallyl-N,N-disubstituted a mmonium halide monomer is diallyld-imethylammonium chloride.

"'Halogen" means fluorine, chlorine, brommine or iodine.

"Monomer" means a polymerizable allylic, vinylic or acrylic compound. The monomer may be anionic, cationic, nonionic or z-witterionic. Vinyl monomers are preferred, acrylic monomers are more preferred.

Representative non-ionic, water-soluble nonomers include acrylamide, methaczrylamide, N,N-dimethylacrylamide, N,N-diethylacrylamide, N-isopropylacrylamide, N-vinylformamide, N-vinylformamide,

butylac=rylamide, N-methylolacrylamide, vinyl ac-etate, vinyl alcohol, and the like.

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Representative anionic monormers include acrylic acid, and it's salts, including, but not limited to sodium acrylate, and ammonium acrylate, methacrylic acid, and it's salts, including, but not limited to socilium methacrylate, and ammonium methacrylate, 2-acrylamido-2-methylpropanesulformic acid (AMPS), the sodium salt of AMPS, sodium vinyl sulfonate, styrene sulformate, maleic acid, and it's salts, including, but not limited to the sodium salt, and ammonium salt, sulfonate, itaconate, sulfopropyl acrylate or methacrylate or other water-soluble forms of these or other polymerisable carboxylic or sulphonic acids. Sulformethylated acrylamide, allyl sulfonate, sodium vinyl sulfonate, itaconic acid, acrylamidomethylbutanoic acid, fumaric acid, vinylphosphonic acid, vinylsulfonic acid, allylphosphonic acid, sulfomethylated acrylamide, phosphonomethylated acrylamide, itaconic anhydride, and the like.

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Representative cationic monomers include allyl amine, vinyl amine, dialkylaminoalkyl acrylates and methacrylates and their quaternary or acid salts, including, but not limited to, dimethy laminoethyl acrylate methyl chloride quaternary salt (DMAEA·MCQ), dimethylaminoethyl acrylate methyl sulfate quaternary salt, dimethyaminoethyl acrylate benzyl c hloride quaternary salt, dimethylaminoethyl acrylate sulfuric acid salt, dimethylar ninoethyl acrylate hydrochloric acid salt, dimethylaminoethyl methacrylate methyl chloride quaternary salt, dimethylaminoethyl methacrylate methyl sulfate quaterna_ry salt, dimethylaminoethyl methacrylate benzyl chloride quaternary salt, dimethylam-inoethyl methacrylate sulfuric acid salt, dimethylaminoethyl methacrylate hychrochloric acid salt, dialkylaminoalkylacrylamides or methacrylamides and their quaternary or acid salts such as acrylamidopropyltrimethylammonium chloride, dimethylaminopropyl acrylamide methyl sulfate quaternary salt, dimet hylaminopropyl acrylamide sulfuric acid salt, dimethylaminopropyl acrylamide hyclrochloric acid salt, methacrylamidopropyltrimethylamm onium chloride, dimethylaminopropyl methacrylamide methyl sulfate quate=rnary salt, dimethylaminopropyl methacrylamide sulfuric acid salt, dimethylaminopropyl methacrylamide hydrochloric acid salt, diethylaminoethylacrylate, diethylaminoethylmethacrylate, diallyldiethylammonium

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chloride and diallyldimethyl ammonium chloride (DADMAC). Alkyl groups are generally C₁₋₄ alkyl.

"Zwitterionic monomer" means a polymerizable molecule containing cat-ionic and anionic (charged) functionality in equal proportions, so that the molecule is met neutral overall. Representative zwitterionic monomers include N,N-dimethyl-N-5 acryloyloxyethyl-N-(3-sulfop-ropyl)-ammonium betaine, N,N-dimethyl-Nacrylamidopropyl-N-(2-carboxymethyl)-ammonium betaine, N,N-dimethyl-N-acrylamidopropyl-N-(3-sulfopropyl)-ammonium betaine, N,Ndimethyl-N-acrylamidopropyl-N-(2-carboxymethyl)-ammonium betaine, 2-(methylthio)ethyl methacryloyl-S-(sulfopropyl)-sulfonium betaine, 2-[(2-10 acryloylethyl)dimethylammonio]ethyl 2-methyl phosphate, 2-(acryloyloxyethyl)-2'-(trin-nethylammonium)ethyl phosphate, [(2-acryloylethyl)dimethylarmmonio]methyl phosphonic acid, 2-methacryloylox yethyl phosphorylcholine (MPC), 2-[(3-acrylamidopropyl)dimethylammonio]ethyl 2'isopropyl phosphate (AAPI) → 1-vinyl-3-(3-sulfopropyl)imidazolium hydroxide, <2-15 acryloxyethyl) carboxymethyl methylsulfonium chloride, 1-(3-sulfopropyl)-2vinylpyridinium betaine, N-C4-sulfobutyl)-N-methyl-N, N-diallylamine ammonitum betaine (MDABS), N,N-dial Iyl-N-methyl-N-(2-sulfoethyl) ammonium betaine, and the like.

"Papermaking process" means a method of making paper products from pulp comprising forming an aque-ous cellulosic papermaking furnish, draining the furnish to form a sheet and drying the sheet. The steps of forming the papermaking furnish, draining and drying may be carried out in any conventional manner generally known to those skilled in the art. Conventional microparticles, alum, cationic starch or a combination thereof may be utilized as adjuncts with the polymer treatment of this invention, though it must be emphasized that no adjunct is required for effective dewatering activity.

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Preferred Embodiments

The aldehyde-functionalized polymers according to this invention are preparred by reacting a polymer comprising armino or amido groups with one or more aldehydles. The polymer comprising amino or a mide groups can have various architectures including linear, branched, star, block, graft, dendrimer, and the like.

Preferred polymers comprising amino or amido groups include polyamines and polyamides. The polyamides and polyamides may be prepared by copolymerizing monomers under free radical forming conditions using any number of techniques including emulsion polymerization, dispersion polymerization and solution polymerization.

Polyamines may also be premared by modification of a pre-formed polyami-de, for example by hydrolysis of acrylamide-vinylformamide copolymer using acid or base as described in U.S. Patent Nos. 6,6 10,209 and 6,426,383.

Polyaminoamides may also be prepared by direct amidation of polyalkyl carboxylic acids and transamidation of copolymers containing carboxylic acid and (meth)acrylamide units as described in U.S. Patent No. 4,919,821.

"Emulsion polymer" and "lætex polymer" mean a polymer emulsion comprising an aldehyde-functionalized polymer according to this invention in the aqueous phæse, a hydrocarbon oil for the oil phase arad a water-in-oil emulsifying agent. Inverse emulsion polymers are hydrocarbon continuous with the water-soluble polymers dispersed within the hydrocarbon matrix. The inverse emulsion polymers are therm "inverted" or activated for use by releasing the polymer from the particles using slaear, dilution, and, generally, another surfactant. See U.S. Pat. No. 3,734,873, incorpor ated herein by reference. Representative preparations of high molecular weight inverse emulsion polymers are described in U.S. Patent nos. 2,982,749; 3,284,393; and 3,734,873. See also, Hunkeler, et al., "Mechanism, Kinetics and Modeling of the Inverse-Microsuspension Homopolymerization of Acrylamide," Polymer, vol. 30(1), pp 127-42 (1989); and Hunkeler et al., "Mechanism, Kinetics and Modeling of Inverse-Microsuspension Polymerization: 2. Copolymerization of Acrylamide with Quaternary Ammonium Cationic Monomers," Polymer, vol. 32(14), pp 2626-40 (1991).

The aqueous phase is prepared by mixing together in water one or more water-soluble monomers, and any polymerization additives such as inorganic salts, chelants, pH buffers, and the like.

The oil phase is prepared by mixing together—an inert hydrocarbon liquid with one or more oil soluble surfactants. The surfactant mixture should have a low hydrophilic-lypophilic balance (HLB), to ensure the formation of an oil continuous emulsion. Appropriate surfactants for water-in-oil emulsion polymerizations, which are commercially available, are compiled in the North American Edition of McCutcheon's Emulsifiers & Detergents. The oil pshase may need to be heated to ensure the formation of a homogeneous oil solution.

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The oil phase is then charged into a reactor equipped with a mixer, a thermocouple, a nitrogen purge tube, and a condenser. The aqueous phase is added to the reactor containing the oil phase with vigorous stirring to form an emulsion. The resulting emulsion is heated to the desired temperature, purged with nitrogen, and a free-radical initiator is added. The reaction mixture is stirred for several hours under a nitrogen atmosphere at the desired temperature. Upon completion of the reaction, the water-in-oil emulsion polymer is cooled to room temperature, where any desired post-polymerization additives, such as antioxidants, or a high HLB surfactant (as described in U.S. Patent 3,734,873) may be added.

The resulting emulsion polymer is a free-flowing liquid. An aqueous solution of the water-in-oil emulsion polymer can be generated by adding a desired amount of the emulsion polymer to water with vigorous miximg in the presence of a high-HLB surfactant (as described in U.S. Patent 3,734,873).

"Dispersion polymer" polymer means a warter-soluble polymer dispersed in an aqueous continuous phase containing one or more -organic or inorganic salts and/or one or more aqueous polymers. Representative examp les of dispersion polymerization of water-soluble polymers in an aqueous continuous phase can be found in U.S. Patent Nos. 5,605,970; 5,837,776; 5,985,992; 4,929,655; 5,006,590; 5,597,859; and 5,597,858 and in European Patent Nos. 183,466; 657,478; and 630,909.

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In a typical procedure for preparing a dispersion polymer, an aqueous solution containing one or more inorganic or organic salts, one or more water-soluble monomers, any polymerization additives such as processing aids, chelants, pH buffers and a water-soluble stabilizer polymer is charged to a reactor equipped with a mixer, a thermocouple, a nitrogen purging tube, and a water condenser. The monomer solution is mixed vigorously, heated to the desired temper-ature, and then a free radical initiator is added. The solution is purged with nitrogen w hile maintaining temperature and mixing for several hours. After this time, the mixture is cooled to room temperature, and any post-polymerization additives are charged to the reactor. Water continuous dispersions of water-soluble polymers are free flowing liquids with product viscosities generally 100-10,000 cP, measured at low shear.

In a typical procedure for preparing solution polymers, an aqueous solution containing one or more water-soluble monomers and any additional polymerization additives such as chelants, pH buffers, and the liesce, is prepared. This mixture is charged to a reactor equipped with a mixer, a the mocouple, a nitrogen purging tube and a water condenser. The solution is mixed vigorously, heated to the desired temperature, and then one or more free radical polymerization initiators are added. The solution is purged with nitrogen while maintaining temperature and mixing for several hours. Typically, the viscosity of the solution increases during this period. After the polymerization is complete, the reactor contents are cooled to room temperature and then transferred to storage. Solution polymer viscosities vary widely, and are dependent upon the concentration and molecular weight of the active polymer component.

The polymerization reactions are initiated by any means which results in generation of a suitable free-radical. Thermally derived radicals, in which the radical species results from thermal, homolytic dissociation of an azo, peroxide, hydroperoxide and perester compound are preferred. Especially preferred initiators are azo compounds including 2,2'-azobis(2-amidinopropane) dihydrochloride, 2,2'-azobis[2-(2-imidazolin-2-yl)propane] dihydrochloride, 2,2'-azobis(isobutyronitrile) (AINN), 2,2'-azobis(2,4-dimethylvaleronitrile) (AIVN), and the like.

The polymerization processes can be carried out as a batch process or in steps. In a batch process, all of the reactive monomers are reacted together, whereas in a step or semi-batch process, a portion of the reactive monomer is withheld from the main reaction and added over time to affect the compositional drift of the copolymer or the formation of the dispersion particles.

The polymerization and/or post polymerization reaction conditions are selected such that the resulting polymer comprising amino or annido groups has a molecular weight of a t least about 1,000 g/mole, preferably about 2,000 to about 10,000,000 g/mole.

The polymer comprising amino or amido groups is then functionalized by reaction wi_th one or more aldehydes. Suitable aldehydess include any compound containing at least one aldehyde

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(-CHO) fu⊐nctional group having sufficient reactivity to react with the amino or amido groups of t⊐he polymer. Representative aldehydes inclu-de formaldehyde,

15 paraformal_dehyde, glutaraldehyde, glyoxal, and the like. Glyoxal is preferred.

The aldehyde-functionalized polymer is prepared by reacting the polyamide or polyamine with aldehyde at a pH between 4 to 12. The total concentration of polymer backbone plus aldehyde is between about 5 to about 35 weight percent. Generally, an aqueous solution of the polymer backbone is prepared for better reaction rate control and increased product stability. The pH of the aqueous polymer backbone solution is increased to between about 4 to about 12. The reaction temperature is generally about 20 to about 80 °C preferably about 20 to about 40 °C. An aqueous aldehyde solution is added to the aqueous polymer backbone solution with good mixing to prevent gel formation. After the addition of aldehyde the pH is adjusted to about 4 to about 12 to achieve the desired reaction rate. After the adjustment of the pH generally the amount of monore acted amide/amine is optimum for the given. ratio of aldehyde to amide/amine and the amount of direacted amide/amine is low. The rate of viscosity increase is monitored during the reaction using a Brookfield viscometer. A viscosity increase of 0.5 cps indicates an increase in polymer medicular weight and an increase in the amount of direacted amide/amine. The amount of monoreacted amide/amine is

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generally maint ained during the viscosity increase but the amount of direacted amide/amine in creases with viscosity. Generally, the desired viscosity increase corresponds to a desired level of monoreacted amide/amine, di_reacted amide/amine and molecular veight. The rate of reaction depends on the terriperature, total concentration of polymer and aldehyde, the ratio of aldehyde to amide/amine functional groups and pH. Higher rates of glyoxylation are expected when the temperature, to tal concentration of polymer and aldehyde, the ratio of aldehyde to amide/amine functional groups or pH is increased. The rate of reaction can be slowed down by decreasing the total concentration of polymer and aldehyde, temperature, the ratio of aldehyde to amide/amine functional groups or pH (to between about 2 to about 3.5). The amount of unreacted aldehyde at the end of the reaction increases as the ratio of aldehyde to amide/amine functional groups is increased. However, the total amount of monoreacted and direacted amide/amine becomes larger.

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For example, reaction of a 95/5 mole percent diallyldimethylammonium chloride/acrylamide copolymer with glyoxal in a molar ratio of 0.4 to 1 glyoxal to acrylamide results in a 95/5 mole percent acrylamide/DADM_AC copolymer with about 15 to 23 mole percent monoreacted and direacted acrylamide and with about 60 to 70 mole percent total unreacted glyoxal at the target product viscosity and molecular weight. A molar ratio of 0.8 to 1 glyoxal to acrylamide results in a 95/5 mole percent acrylamide/DADMAC copolymer with about 22 to 30 mole percent monoreacted and direacted acrylamide and with about 70 to 80 mole percent total unreacted glyoxal at the target product viscosity and molecular weight.

The product shelf stability depends on the storage temperature, product viscosity, total amount of reacted amide/amine, total concentration of polymer and aldehyde, the ratio of aldehyde to amide/amine functional groups and pH. Generally, the pH of the product is maintained at a low pH (2 to 3.5) and the total concentration of polymer and aldehyde is optimized to extend shelf stability.

The reaction conditions are selected such that at least about 15 mole percent, preferably at least about 20 mole percent of the amino or am ido groups in the polymer react with the aldehyde to form the aldehyde-functionalized polymer. The resulting

aldehyde-functionalized polymers have a weight average molecular weight of at least about 100,000 g/mole, preferably at least about 300,000 g/mole.

In a preferred aspect of this invention, the aldehyde functionalized polymer is an aldehyde functionalized polyamide.

In another preferred aspect, the aldehyde functionalized polyamide is an aldehyde-functionalized polymper comprising 100 mole percent of one or more nonionic monomers.

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In another preferred as pect, the aldehyde functionalized polyamide is an aldehyde functionalized copol_ymer comprising about 5 to about 99 mole percent of one or more acrylamide monomers and about 95 mole percent to about 1 mole percent of one or more cationic, anionic or zwitterionic monomers, or a mixture thereof.

Copolymers prepared from nonionic monomers and cationic monomers preferably have a cationic charge of about 1 to about 50 mole percent, more preferably from about 1 to about 30 mole percent.

Copolymers prepared from nonionic monomers and anionic monomers preferably have an anionic charge of about 1 to about 50 mole percent, more preferably from about 1 to about 30 mole percent.

Amphoteric polymers preferably have an overall positive charge. Preferred amphoteric polymers are composed of up to about 40 mole percent cationic monomers and up to about 20 mole percent anionic monomers. More preferred amphoteric polymers comprise about 5 to about 10 mole percent cationic monomers and about 0.5 to about 4 mole percent anionic monomers.

Zwitterionic polymers preferably comprise 1 to about 95 mole percent, preferably 1 to about 50 mole percent zwitterionic monomers.

In a preferred aspect of this invention the aldehyde-functionalized polyamide is an aldehyde functionalized copolymer comprising about 1 to about 99 mole percent of one or more acrylamide monomers and about 99 mole percent to about 1 mole percent of one or more cationic, anionic or zwitterionic monomers, or a mixture thereof.

In another preferred espect, the aldehyde functionalized polyamide is an aldehyde functionalized copolymer comprising about 50 to about 99 mole percent of

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one or more acrylamide monomers and about 50 to about 1 mole percent of one or more cationic monomers.

In another preferred aspect, the aldehyde functionalized polymer is a coportymer comprising about 50 to about 99 ranole percent of one or more acrylamide monomers and about 50 to about 1 mole percent of one or more cationic monomers wherein the copolymer is functionalized with glyoxal.

In another preferred aspect, the cationic monomer is a diallyl-N,N-disubstatuted ammonium halide monomer.

In another preferred aspect, about 20 to about 50 mole percent of the amidee groups of the copolymer have reacted with glyoxal.

In another preferred aspect, the nonionic monomer is acrylamide and the diallyl-N,N-disubstituted ammonāum halide monomer is diallyldimethylammoniu_m chloride.

In another preferred aspect, the functionalized polymer is a copolymer comprising about 70 to about 99 mole percent of acrylamide and about 1 to about 30 mole percent of diallyldimethylar monium chloride functionalized with glyoxal.

In another preferred aspect, about 20 to about 26 mole percent of the amide groups of the copolymer have reacted with glyoxal.

The aldehyde-functionalized polymers are useful for dewatering all grade_s of paper and paperboard with board grades and fine paper grades being preferred.

Recycle board grades using OCC (old corrugated containers) with or without mixed waste have been particularly responsive.

Useful increases in dewattering can be achieved with aldehyde-functional zed polymer doses in the range 0.05 to 15.0 lb polymer/ton of dry fiber with best results normally achieved in the range 0.5 to 3.0 lb/ton depending on the particular papermaking circumstances (pap ermachine equipment and papermaking raw materials used).

The aldehyde functionalized polymers of the invention can be added in traditional wet end locations used for conventional wet end additives. These include to thin stock or thick stock. The actual wet end location is not considered to be critical,

but the aldehyde-functionalized polymers are preferably added prior to the addition of other cationic additives. Because the aldehyde-functionalized polymers are believed to act as pressing aids, the raddition to the wet end is not necessary, and the option of adding them just prior to the press section after the formation of the sheet can also be practiced. For example, the polymer can be sprayed on the wet web prior to entering the press section, and their can be a preferred mode of addition to reduce dosages or the effects of interferences which might occur in the wet end. Other traditional wet end additives can be used ir combination with the aldehyde functionalized polymers.

These include retention aids, strength additives such as starches, sizing agents, and the like.

When using ald hyde-functionalized polymers as described herein having net anionic charge, a method of fixing the polymer to the fiber is needed. This fixing is typically accomplished by using cationic materials in conjunction with the polymers. Such cationic materials are most frequently coagulants, either inorganic (e.g. alumn, polyaluminum chloride s, iron chloride or sulfate, and any other cationic hydrolyzing salt) or organic (e.g. p-DADMACs, EPI/DMAs, PEIs, modified PEIs or any other high charged density low to medium molecular weight polymers). Additionally, cationic materials added for other purposes like starch, wet strength, or retention additives can also serve to fix the ani-onic polymer. No additional additives are need to fix cationic aldehyde-functionalized polymers to the filler.

The foregoing remay be better understood by reference to the following examples, which are presented for purposes of illustration and are not intended too limit the scope of the invention.

25 Example 1

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Preparation of 95/5 mole % Acrylamide/DADMAC copolymer.

To a 1500-mL reaction flask fitted with a mechanical stirrer, thermocoupLe, condenser, nitrogen purge tube, and addition port is added 116.4 g of deionized o-r soft water, 26.3 g of phosphnoric acid, 63.8 g of a 62% aqueous solution of diallyldimethyl ammonium chloride (N alco Company, Naperville, IL), 7.6 g of sodium formate, and

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0.09 g of ethylenediaminetetraacetic acid, tetra sodium salt. The reaction mixture is stirred at 400 rpm and the pH adjusted to 4.7 to 4.9 using 1 7.3 g of aqueous 50 % sodium hydroxide solution. The resulting mixture is heate d to 100 °C and purged with nitroge at 50 mL/min. Upon reaching 100 °C, 17.6 g of a 25.0% aqueous solution of ammorphium persulfate is added to the reaction mixture over a period of 135 minutes. Five mainutes after starting the ammonium persulfate addition, 750.9 g of a 49.5% aqueous solution of acrylamide is added to the reaction mixture over a period of 120 minutes. The reaction is held at 100 °C for 180 minutes after ammonium persulfate addition. The reaction mixture is then cooled to ambient temperature and the pH is adjusted to 5.2 to 5.8 using 50 % aqueous sodium hydroxide solution or concentrated sulfuric acid. The product is a viscous, clear to amber solution. The product has a molecular weight of about 20,000 g/mole.

Example 2

Glyox_ylation of 95/5 mole % Acrylamide/DADMAC cop-olymer with 0.8 to 1 glyoxal to acr_ylamide mole ratio at 9.0 % actives (total glyoxal ar_id polymer).

To a 2000-mL reaction flask fitted with a mechan-ical stirrer, thermocouple, conde=nser, addition port and sampling valve at the bottorm of the reactor is added 238.0 g of a. 41% aqueous solution of 95/5 mole % acrylamide/DADMAC copolymer, preparred as in Example 1, and 1304.0 g of deionized or s. oft water. The polymer solution is stirred at 400 rpm. The pH of the solution is adjusted to 8.8 to 9.1 by addin_g 5.8 g of 50% aqueous sodium hydroxide solution. The reaction temperature is set at 24 to 26 °C. Glyoxal (143.0 g of a 40% aqueous solution) is added to the reaction mixture over 20 to 30 minutes. The Brookfield viscosity (Brookfield Programmable LVDV-II+ Viscometer, LV # 1 spindle at: 60 rpm, Brookfield Engineering Laboratories, Inc, Middleboro, MA) of the reaction mixture is about 4 to 5 cps a fter glyoxal addition. The pH of the reaction mixture is adjusted to 7.5 to 8.8 using 10 % aqueous sodium hydroxide (25 g) added over 20 to 30 minutes. The Brookfield viscosity (Brookfield Programmable LVDV-II+ Viscometer, LV # 1 spindle at 60 rpm, Brookfield Engineering Laboratories, Inc, Mi ddleboro, MA) of the reaction

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mixture is about 4 to 5 cps after sodium hydroxide addition. The TH of the reaction mixture is maintained at about 7.0 to 8.8 at about 24 to 26 °C with good mixing. The Brook_field viscosity is monitored and upon achieving the desired wiscosity increase of greate=r than or equal to 1 cps (5 to 200 cps, >100,000 g/mole) the mpH of the reaction mixture is decreased to 2 to 3.5 by adding sulfuric acid (93 %) to substantially decrease the reaction rate. The rate of viscosity increase is dependent on the reaction pH and temperature. The higher the pH of the reaction mixture the faster 7the rate of viscosity increa_se. The rate of viscosity increase is controlled by decreasing the pH of the reaction mixture. The product is a clear to hazy, colorless to amber, fluid with a Brook=field viscosity greater than or equal to 5 cps. The resulting product is more stable upon storage when the Brookfield viscosity of the product iss less than 40 cps and when the product is diluted with water to lower percent actives. The product can be prepared at higher or lower percent total actives by adjusting the d_esired target product viscosity. NMR analysis of the samples prepared indicate that about 70 to 80 % of the glyox all is unreacted and 15 to 35 mole percent of the acrylamide units reacted with glyox al to from monoreacted and direacted acrylamide.

Exam_ple 3

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Dewa tering effectiveness of representative aldehyde-functionalize d polymers.

The dewatering effects of glyoxalated DADMAC/Acrylam_ide polymers preparted with glyoxal to acrylamide mole ratios (here after referred to as the G/A ratio) of 0.1 = 0.2, 0.4 and 0.8 are evaluated through paper machine trials. The relative performance of polymers prepared using the 0.1, 0.2, and 0.8 G/A ratios are compared to the polymer prepared with the 0.4 mole ratio. The trials are run on a dual headbox Fourd_rinier papermachine using 100% OCC furnish manufacturing recycle linerboard and corrugating medium. Actual papermachine conditions varied depending on the speciffic grade of paperboard being made. In all cases, a retention program of polyal_uminum chloride fed to the thick stock and a cationic floccu_lant fed to the thin stock is used. For linerboard grades, ASA sizing fed to the thin stock is also present. The glyoxalated acrylamide polymers are applied through a spray loom to the

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underside of the top ply prior to meshing with the bottom ply, although earlier trials demonstrated the dewatering effect could also be achieve d by wet-end thick or thin stock addition.

The dewatering effect of the polymers is evaluated on the basis of steam press ure changes in the machine dryer section which are provided through the mills DCS (distributive control system) computer system. The sheet moisture at the reel is meas ured on-line and is maintained by adjustment of the steam pressure (a measure of steam usage or energy consumption). Lower sheet moistuire at the reel reflects a lower sheet moisture going into the dryer section or equivalently, better dewatering through the machine sections preceding the dryer section. The lower steam demand, as meas ured by pressure, then reflects improved dewatering. If the steam pressure in these sections drops to a level where the operator feels comfortable that normal swings in steam demand can be handled, then he will increase the machine speed manually. When changes in polymer type or dose are made, the stealm pressure from one of the steams sections is followed closely to see if any change oc curs, with proper consi deration given to changes in production rates when they occur. The initial effect of a derier sheet is observed by lower percent moisture det ected at the reel. However, this £ rop in percent moisture is short lived because of the automatic regulation leaving only the steam reduction as a permanent reminder of any dewatering effect produced. Many factors other than addition of the aldehyde-function alized polymer also affect the sheet moisture, but most, like stock changes, occur over a longer time frame than the steam reduction effect caused by the polymer additive, particularly when applied on the table through spray application. Consequently, the steam reduction is a better indicator of pollymer dewatering than the average production rate or machine speed, as these measures are more easily confounded with the other factors which effect machine speed.

Example 3a

Comparison of polymer with a 0.1 G/A ratio with polymer having a 0.4 G/A ratio.

Comparison of these two polymers is conducted on 42 lb linerboard in the absence of wet-end starch. After a baseline is established with the 0.4 G/A ratio

polymer at 2.0 1b/ton, the 0.1 G/A ratio polymer is substituted at 2.2 lb/ton. Almost immediately, the sheet at the reel is consistently observed to be wetter and the steam demand increases to maximum in about 1 hr which necessitates the re-introduction of the 0.4 G/A ratio polymer to prevent slowing down the paper machine. To regain control of the machine, 3 lb/ton of the 0.4 G/A ratio polymer is needed, and its addition results in a dramatic reduction in steam pressure, 12 psi in 15 min. Subsequently, a baseline with the 0.4 G/A ratio polymer is reestablished at 2 lb/ton whereupon substitution with 0.1 G/A ratio polymer at the higher dose of 3.4 lb/ton is initiated. At this much higher dose, the steam pressure progressively increases over a period of about an hour a gain to the point where it becomes necessary to revert b ack to the 0.4 G/A ratio polyrmer to prevent slowing the machine. Again, with the 0.4 G/A ratio polymer added at 3.0 lb/ton the steam pressure is quickly reduced, 12 p-si in 15 min. and this reduction could be maintained even when the 0.4 G/A ratio po lymer's dose is reduced to 2 lb/ton. The 0.1 G/A ratio polymer could not maintain the steam pressure, and therefore the machine speed, achieved by 0.4 G/A ratio polymer even at a dose 70% higher. No change in the strength specifications for this grade (Mullen and Scott bond) could be detected when the 0.1 G/A ratio polymer is substituted for the 0.4 G/A ratio polymer.

20 Example 3b

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Comparison of polymer with a 0.2 G/A ratio with polymer having a 0.4- G/A ratio.

Comparison of these two polymers is conducted on 35 lb linerboard with 5 lb/ton wet-end starch fed to the thick stock. After a baseline is established with the 0.4 G/A ratio polymer at 2.0 lb/ton, the 0.2 G/A ratio polymer is substituted at 2.2 lb/ton. At this dosage, a modest increase in steam pressure of 5 psi is measured over a period of about the hour. Reintroduction of the 0.4 G/A ratio polymer resulted in an immediate decrease in reel moisture and a quick decline in steam pressure of 3 psi in 10 min. Switching back to the 0.2 G/A ratio polymer at 2.2 lb/ton at this point keeps the steam reasonably constant for about an hour with only a 2 psi increase. Again, reintroduction of 2 lb of the 0.4 G/A ratio polymer results in a quick decline in steam

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pressure of 8 psi in 20 min. indecative of improve dewatering. Based on these results, the 0.2 G/A ratio polymer certaginly demonstrates dewatering ability, but even at a 10% increase in dosage, it could not maintain the pressure achievable with the 0.4 G/A ratio polymer. Additionally, unlike the 0.1 G/A ratio polymer, the 0.2 G/A ratio polymer is capable of keeping the machine running at the desired speed although at increased steam demand and dosage relative to the 0.4 G/A ratio polymer. The trial results with these three polymers indicated that the 0.4 G/A ratio polymer gives better dewatering than the 0.2 G/A ratio polymer and it in turn gives better dewatering than the 0.1 G/A ratio polymer. No change in the strength specifications for this grade (STFI) could be detected when the 0.2 G/A ratio polymer is substituted for the 0.4 G/A ratio polymer.

Example 3c

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Comparison of polymer with a 0.8 G/A ratio with polymer having a 0.4 G/A ratio.

Based on the discovery that increasing the G/A ratio in the preparation of the polymers increases dewatering , an even higher G/A ratio of 0.8 is prepared and evaluated on the same papermachine. Comparison of the 0.8 G/A ratio polymer with the 0.4 G/A ratio polymer is conducted on 33 lb corrugating medium in the absence of wet-end starch. Addition of the 0.4 G/A ratio polymer at 2.0 lb/ton results in a very good reduction in steam pressure of 21 psi after about 2 hours at which time 1.5 lb/ton of the 0.8 G/A ratio polymer replaces the 2 lb/ton of 0.4 G/A ratio polymer. Even with the 25% reduction in dose, the addition of the 0.8 G/A ratio polymer results in an even further reduction in steam pressure by 3 psi and a dramatic increase in steam pressure of 12 psi occurs in 0.5 hour when it is removed. Further trialing is conducted on 26 lb corrugating medium in the abs ence of wet-end starch. Starting again with 2.0 lb/ton of 0.4 G/A ratio polymer to estab lish the baseline, a substitution of 2.0 lb/ton of 0.8 G/A ratio polymer results in a drop in steam pressure of 7 psi in 60 min., which further decreases by 4 psi when the dosage is increased to 3 lb/ton in 10 min. Reducing the 0.8 G/A ratio polymer to only 1.0 lb/ton relative to the 3 lb/ton results in an increase in steam pressure, but it remains 8 psi below the 2.0 lb/ton 0.4 G/A ratio polymer value even with an increase in mach-ine speed of 30 ft/min. Based on these trial results the

0.8 G/A ratio polymer appears to yield equivalent dewatering at a dose 25 to 50% less than required by the 0.4 G/A ratio polymer. The strength specification for both medium grades (Concorra) made with the 0.8 G/A ratio polymer exhibit values equal to or greater than those obtained with for the 0.4 G/A ratio polymer even though the dosages are generally I ower.

Based on these trial results, increasing the G/A ratio in the preparation of the aldehyde-functionalized polymers is found to provide increased dewatering a_ctivity with the preferred ratio being greater than 0.4.

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Changes can be made in the composition, operation and arrangement of the method of the invention described herein without departing from the concept and scope of the invention as defined in the claims.

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CLAIMS

- 1. A method of enhancing the dewatering of a paper sheet on a paper machine comprising adding to the paper sheet about 0.05 lb/ton to about 15 lb/ton,

 5 based on dry fiber, of one or more aldeh yde functionalized polymers comprising amino or amido groups wherein at least about 1.5 mole percent of the amino or amido groups are functionalized by reacting with one or more aldehydes and wherein the aldehyde functionalized polymers have a weight a verage molecular weight of at least about 100,000 g/mole.
- 10 2. The method of claim 1 wherein the aldehyde functionalized polymers are selected from the group consisting of aldehyde functionalized polyamines and aldehyde functionalized polyamides.
 - 3. The method of claim 1 wherein the aldehydes are selected from formaldehyde, paraformaldehyde, glyoxal and glutaraldehyde.
 - 4. The method of claim 1 wherein the aldehyde functionalized polymer is an aldehyde functionalized polyamide.
 - 5. The method of claim 4 wherein the aldehyde functionalized polyamide is an aldehyde-functionalized polymer comprising 100 mole percent of one or more nonionic monomers.
 - 6. The method of claim 4 wherein the aldehyde functionalized polyamide is an aldehyde functionalized copolymer comprising about 5 to about 99 mole percent of one or more acrylamide monomers and about 95 mole percent to about 1 mole percent of one or more cationic, anionic or zwitterionic monomers, or a mixture thereof.
- 7. The method of claim 6 wherein the aldehyde functionalized polyamide is an aldehyde-functionalized copolymer c omprising 1 to about 50 mole percent of one or more anionic monomers and 99 to about 50 mole percent of one or more nonionic monomers.
- 8. The method of claim 6 whe rein the aldehyde functionalized polyamide
 30 is an aldehyde-functionalized copolymer comprising 1 to about 30 mole percent of one

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or more anionic monomers and 99 to about 70 mole percent of one or more nonionic monomers.

- 9. The method of claim 6 wherein the aldehyde functionalized copolymer is an aldehyde-functionalized amphoteric polymer comprising up to about 40 mole percent of one or more cationic monomers and up to about 20 mole percent of one or more anionic monomers.
- 10. The method of claim 6 wherein the al-dehyde functionalized copolymer is an aldehyde-functionalized amphoteric polymer comprising about 5 to about 10 mole percent of one or more cationic monomers and about 0.5 to about 4 mole percent of one or more anionic monomers.
- 11. The method of claim 6 wherein the alldehyde functionalized copolymer is an aldehyde-functionalized zwitterionic polymer comprising about 1 to about 95 mole percent of one or more zwitterionic monomers.
- 12. The method of claim 6 wherein the a_ldehyde functionalized copolymer

 1.5 is an aldehyde-functionalized zwitterionic polymer comprising about 1 to about 50 mole percent of one or more zwitterionic monomers.
 - 13. The method of claim 6 wherein the aldehyde functionalized polyamide is an aldehyde functionalized copolymer comprising about 50 to about 99 mole percent of one or more acrylamide monomers and about 50 to about 1 mole percent of one or more cationic monomers.
 - 14. The method of claim 13 wherein at Teast about 20 mole percent of the amide groups of the polyamide have reacted with a ldehyde.
 - 15. The method of claim 1 wherein the aldehyde functionalized polymer is a copolymer comprising about 50 to about 99 mole percent of one or more acrylamide monomers and about 50 to about 1 mole percent of one or more cationic monomers wherein the copolymer is functionalized with glyopal.
 - 16. The method of claim 15 wherein the cationic monomer is a diallyl-N,N-disubstituted ammonium halide monomer.
 - The method of claim 16 wherein about 20 to about 50 mole percent of the amide groups of the copolymer have reacted with glyoxal.

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- 18. The method of claim 16 wherein the nonionic monomer is acrylamide and the diallyl-N,N-disubstituted ammo nium halide monomer is diallyldimethylammonium chloride.
- 19. The method of claim 18 wherein the aldehyde-functionalized polymer has a molecular weight of at least 300,000 g/mole.
 - 20. The method of claim 19 wherein the aldehyde-functionalized polymer is a copolymer comprising about 70 to ab out 99 mole percent of acrylamide and about 1 to about 30 mole percent of diallyldimæthylammonium chloride functionalized with glyoxal.
- 10 21. The method of claim 20 wherein about 20 to about 26 mole percent of the amide groups of the copolymer have reacted with glyoxal.
 - 22. The method of claim 21 wherein about 0.5 lb/ton to about 3 lb/ton, based on dry fiber, of glyoxylated copolymer is added to the paper sheet.
- 23. The method of claim 1 wherein the aldehyde functionalized polymer is sprayed onto the paper sheet prior to press dewatering.