APPLICATION FOR A PATENT
(Combined Form – Convention and Non-Convention)

X/We C. H. DEXTER LIMITED, a corporation organized under the Laws of the United Kingdom having a place of business at Chirnside, Duns, Berwickshire, Scotland TD11 3JU hereby apply for the grant of a Patent for an invention entitled PAPER HAVING GOOD ABSORBENCY AND ALKALI RESISTANCE which is described in the accompanying Provisional Specification.

2. This application is a Convention Application and is based on the application(s) for a patent or similar protection made—
in
UNITED KINGDOM
on June 25, 1976, numbered 26531/76, and
on June 25, 1976, numbered 26530/76,

3. My/Our address for service is: Care of COWIE, THOMSON & CARTER Patent Attorneys, of Suite 5, 65 Queens Road, Melbourne, Australia, 3004

DATED this 14th day of June 1977

COWIE, THOMSON & CARTER.

PATENT ATTORNEYS FOR C. H. DEXTER LIMITED.

To the Commissioner of Patents, COMMONWEALTH OF AUSTRALIA

COWIE, THOMSON & CARTER Patent Attorneys, Suite 5, 65 Queens Road, Melbourne, Australia, 3004
Declarations in Support of an Application for a Patent

(Combined Form - Convention and Non-Convention)

*Strike out for non-convention

In support of the Convention application made for a patent for an invention entitled, PAPER HAVING GOOD ABSORBENCY AND ALKALI RESISTANCE

I, Dr. Newlyn Jones, Technical Manager and Director of C. H. Dexter Limited,

do solemnly and sincerely declare as follows:-

1. I am the applicant(s) for the patent of addition.

(or in the case of an application by a body corporate)

1. I am authorised by C. H. Dexter Limited, the applicant for the patent of addition, to make this declaration on its behalf.

2. The basic application(s) as defined by section 141 of the Act was made at

in United Kingdom

on the 25th day of June 1976 No. 26531/76

by Alistair C. Stewart & Newlyn Jones

and

on the 25th day of June 1976 No. 26530/76

by Newlyn Jones, Alistair C. Stewart & J. F. Ward

and

on the day of 19. No.

3. I am the actual inventor(s) of the invention.

(or where a person other than the inventor is the applicant)

3. Newlyn Jones, a British Subject of Forrest Lodge, of Foulcon Rford, Berwick-upon-Tweed, England,

Alistair Charles Stewart, a British Subject, of Newtonlea, Chirnside, Duns, Berwickshire, Scotland,

James F. Ward, a British Subject, of Hedworth, Todlaw Road, Duns, Berwickshire, Scotland

are the actual inventor(s) of the invention and the facts upon which I am the said Corporation is entitled to make the application are as follows:- the said Corporation is the assignee of the invention from the said actual inventors

4. The basic application(s) referred to in paragraph 2 of this Declaration was the first application(s) made in a Convention country in respect of the invention the subject of the application.

DECLARED AT Chirnside this 24th day of June 1977.

Signature of Declarant
PAPER HAVING GOOD ABSORBENCY AND ALKALI RESISTANCE

Apparatus is also claimed

CLAIM

1. A process for the production of paper, which process comprises forming a fibrous paper web and treating the fibres with a water-soluble cationic, thermosetting, epihalohydrin-containing resin, a non-viscose film-forming material and a polyalkylene imine.
COMPLETE SPECIFICATION

(ORIGINAL)

FOR OFFICE USE

Class:

Int. Cl:

Application Number:
Lodged:

Complete Specification—Lodged:
Accepted:
Published:

Priority:

Related Art:

TO BE COMPLETED BY APPLICANT

Name of Applicant: C.H. DEXTER LIMITED, a corporation organized under the Laws of the United Kingdom having a place of business at Chirnside, Duns, Berwickshire, Scotland TD11 3JU.

Address of Applicant:

Actual Inventor: NEWLYN JONES
ALISTAIR C. STEWART
JAMES F. WARD

Address for Service: c/- COWIE, THOMSON & CARTER,
PATENT ATTORNEYS,
SUITE 5, 65 QUEENS ROAD,
MELBOURNE, VICTORIA 3004.

Complete Specification for the invention entitled:
PAPER HAVING GOOD ABSORBENCY AND ALKALI RESISTANCE.

The following statement is a full description of this invention, including the best method of performing it known to me:

*Note: The description is to be typed in double spacing, pica type face, in an area not exceeding 250 mm in depth and 160 mm in width, on tough white paper of good quality and it is to be inserted inside this form.
The present invention relates to a process for the production of paper containing certain additives, which paper exhibits a good alkaline wet strength and a good water absorbency, to the paper so produced and to products comprising such paper.

One field in which the present invention finds application is in the production of casing paper, which may be used for the manufacture of packaging for meat products such as sausage.

Casing paper is commonly manufactured from paper webs of relatively strong, high-tenacity natural fibres, such as abaca, sisal or flax. The paper web is saturated with a dilute viscose solution, for example a solution obtained by diluting a solution containing 7% by weight of cellulose (as cellulose xanthate) and 6% by weight of sodium hydroxide to a 1% cellulose content. The viscose-saturated web is dried and the cellulose in the viscose is then regenerated by passing the web through an acidic regenerating bath containing, for example, a 1 - 8% aqueous sulphuric acid solution. The web is then washed free of acid and dried to produce a paper web impregnated with acid-regenerated cellulose. This casing paper is then generally formed into rolls ("master rolls").

Casings for the packaging of processed meats, e.g. sausage, may be manufactured from the casing paper by cutting it into strips which are then folded to form tubes. The tubes are saturated with an alkaline viscose solution, containing, for example, 7% by weight of cellulose and 6% by weight of sodium hydroxide. The cellulose in the viscose is then regenerated by means of an acidic regenerating bath containing, for example, dilute sulphuric acid and possibly such salts as sodium sulphate or ammonium sulphate. The tube is then passed through one or more baths in order to
wash out the acid and the salts.

If desired, the tube may be passed through an aqueous bath which contains a plasticizer, e.g. glycerine, for the regenerated cellulose. The tube is dried by passing it through a heated chamber (the tube being in an inflated state) to give a cellulosic tubing which has embedded therein a paper web. This tubing may then be stuffed with a processed meat product under pressure. A process of this type is described in detail in United States Patent No. 3,135,613.

The purpose in treating the initial paper web with dilute viscose solution, followed by regeneration, is to provide the web with strength and structural integrity so that it may withstand the treatment with the highly caustic viscose solution used in the formation of the casing tubes. The amount of acid-regenerated cellulose in the casing paper is, in fact, comparatively low; thus, the casing paper may have a typical basis weight (weight per unit area) of 20 grams/m², of which the acid-regenerated cellulose accounts for 0.6 g/m², compared with the material of the casing tubes which may have a typical basis weight of 70 - 80 g/m², of which 50 - 60 g/m² may be accounted for by the acid-regenerated cellulose. However, despite the initial treatment with viscose, the treatment with the highly caustic viscose solution used in the formation of the casing tubes will inevitably entail a certain degree of softening and weakening of the web. This imposes a limit on the production speeds if difficulty in handling the webs and possible break-downs in production are to be avoided. There is accordingly a need in the art for casing paper having an improved alkali resistance in order to permit higher production speeds in the manufacture of the casing tubes.
In United States Patent 3,378,379, there is disclosed a tubular regenerated-cellulose casing for dry sausage, which casing is provided with a coating comprising a cationic thermosetting resin bonded to the inside wall thereof. The patent suggests that polyethylene imine may also be employed for this coating, although this material is not, in fact, a thermosetting resin. The purpose of the inner coating is to improve the adhesion of the sausage casing to a dry sausage product despite any shrinkage which may occur when the dry sausage product is processed and dried in the casing over a prolonged period of time. It should be noted, however, that in the aforesaid process it is not the casing paper as such which is treated with the thermosetting resin, but the tubular casing material. In the embodiment illustrated in U. S. Patent 3,378,379, the cationic thermosetting resin is applied to the inner surface of the casing tube after the application of glycerine and before the casing is dried, in an inflated state, in a heated chamber.

Another problem recognised in U. S. Patent No. 3,378,379 is the variation in extensibility in the transverse direction exhibited by casing paper strips cut from different parts of the master roll. This can cause variation in the properties of the final casing tubes, which may therefore be unsatisfactory to the meat packager, for whom dimensional stability in the product is of commercial importance. To meet this problem it is suggested in the aforesaid U. S. Patent that a cationic thermosetting resin (e.g. a reaction product of epichlorohydrin and a polyamide, a modified melamine-formaldehyde resin or a modified urea-formaldehyde resin) may be employed as the bonding agent in the casing paper, instead of the commonly
employed acid-regenerated viscose. The cationic thermosetting resin is employed in an amount of at least 0.5% by weight based upon the dry weight of the impregnated fibrous web. The resin may be incorporated into the fibrous web by adding the resin to the fibrous slurry prior to forming the fibrous web. Alternatively, the formed fibrous web may be impregnated with the resin by passage through an aqueous solution of said resin. The said U.S. Patent also discloses the use of the thermosetting resin in combination with viscose which is not regenerated with an acid; however, in the latter case the viscose may be auto-regenerated by storing the web for a sufficient period.

U.S. Patent 3,484,256 discloses a process for the production of casing paper, in which a fibrous web is bonded with a cationic thermosetting resin, and a polyacrylamide resin. An interesting point is that in Table III of that Patent, there is a comparison of the tensile strengths, under various conditions (dry, wet and 6% aqueous NaOH), of paper webs that are bonded with regenerated viscose or with polyamide-epichlorohydrin resin and polyacrylamide or with just polyamide-epichlorohydrin resin. The results might at first suggest that the alkaline wet strength of the web bonded with just polyamide-epichlorohydrin resin is at least as good as that of the viscose-bonded web. However, these results require careful qualification and are considered in greater detail hereinafter in Example 5.

In United Kingdom Patent Specification 1,091,105 there is described a process in which casing paper is produced by incorporating into a paper web an alkaline-curing resin such as polyethylene imine or a polymeric reaction product of
epichlorohydrin and a polyamide. The use of the alkaline-curing resin in place of the customary treatment with dilute viscose is said to result in a casing paper which has more uniform characteristics across its width and which results in a casing having improved burst strengths. However, it has since been found that the casing paper so produced has an alkali resistance which is insufficient to permit sufficiently high production speeds for the commercial manufacture of the casing tubes.

In general, the alkali resistance of regenerated viscose casing paper is merely adequate: this means that in order to avoid numerous breaks in the web on the manufacturing equipment, the operation speeds must be limited. It has also been found that the absorbency of viscose-treated casing paper webs deteriorates with ageing. This will also serve to restrict the rate of production of the casing tubes; it is not uncommon for a period of three months or more to elapse between the manufacture of the casing paper and its subsequent use.

Another field in which the present invention finds application is in the production of paper for the manufacture of infusion pouches, for example teabags and spice-bags.

Teabags and the like are commonly formed as pouches of a material ("teabag paper") that is permeable to water and to the beverage formed by infusion i.e. by the dissolution of soluble solids in the contents of the pouch, upon the application of hot water thereto. Desirable characteristics of the material are cleanliness, good absorbency, high wet strength, a sheet structure to permit rapid diffusion of the tea extract, and ability to perform satisfactorily in high-speed packaging equipment with which the teabags are fabricated and filled.
The strength of the pouch is determined, by and large, by three main factors: firstly, the fibre composition of the paper and the chemical treatments, if any, that the paper has undergone; secondly, the nature of the contents (e.g. tea) of the pouch; and thirdly, the interaction, if any, between the solids dissolved during the infusion and the chemicals with which the paper fibres have been treated.

Hitherto, the chemical treatment of fibrous webs for tea-bag paper has, in general, been effected by either of two methods. In the first method, the fibrous sheet is saturated with viscose and the cellulose in the viscose is then regenerated with a dilute acid. This method produces a fibrous web with an adequate strength in aqueous alkaline conditions. However, the product has a disadvantage, in that it can impart an undesirable taste to tea or other beverages.

The second method comprises saturating the fibrous sheet with a mixture of a polycarboxylic chlorohydrin resin and carboxy methyl cellulose, as described, for instance, in U.K. Patent Specification 1,111,165. This method produces a fibrous web which is somewhat weaker in alkaline, aqueous media than the fibrous webs produced by the first method. This product, however, is neutral as regards taste.

The drawbacks of these prior-art methods are particularly noticeable when the papers are used for the production of herbal teas; herbal teas are alkaline. Thus the viscose-treated paper can be formed into a bag having an adequate alkaline wet strength, but which gives rise to a beverage of poor taste or flavour. On the other hand, the other type of paper will not affect the taste of the beverage, but the beverage...
formed therefrom will have a low alkaline wet strength.

Mention should also be made of U.S. Patent No. 2,698,793 which discloses a process for producing a sized cellulosic sheet having resistance to water and ink. A water-insoluble, hydrophobic sizing agent, in the form of a water-dispersible salt, and an alkyleneimine resin are added separately to an aqueous suspension of the cellulosic fibres, which suspension is allowed to stand after each addition, and the fibres are subsequently formed into a sheet. The sheet is then heated to 105° to 150°C to develop the sizing properties of the sizing agent. Additionally, other sizing or filling agents, e.g. starch, can also be employed.

The said U.S. Patent, it should be stressed, is concerned with giving the sheet a resistance to water by rendering it water-repellant. Thus, water and water-based inks are prevented from penetrating the interstices of the paper by the increased surface tension at the surfaces of the paper matrix. It should be noted that the sizing agents specified in U.S. Patent 2,698,793 must be water-insoluble, hydrophobic compounds. The "wet strength" exhibited by the sheets treated in accordance with the process of the said U.S. Patent is attributable primarily to the fact that the fibres remain in a dry state below the water-repellant surface of the matrix. It has, in fact, been known for a long time that water-resistance can be imparted to paper webs by treatment with chemicals such as rosin size. However, once water has been driven into the interstices of the web, either by mechanical forces or by use of surfactants, the apparent wet strength of the sheets would be lost.

In contrast, the present invention is directed to the
very different problem of producing a paper web that has alkaline wet strength (a property with which U.S. Patent 2,698,793 is not concerned) yet at the same time is absorbent. Clearly, a water-repellancy mechanism, as employed in U.S. Patent 2,698,793, would be of no use in the field of casing papers and teabag papers: the former are required to absorb aqueous viscose solutions and the latter must not present any impediments to the passage of water and of the aqueous infusion.

It is an object of the present invention to provide a process for the production of paper having a good alkaline wet strength and a good absorbency.

It is a further object to provide such a process whereby casing paper can be produced which retains its absorbency even after ageing, thus permitting casing tubes to be produced using high-speed commercial processes.

It is another object of the present invention to provide a process whereby teabag paper can be produced that combines taste neutrality with a good alkaline wet strength, whilst retaining a good absorbency.

It is yet a further object to provide such processes that avoid the use of viscose and its consequent acid regeneration.

According to the present invention, a process for the production of paper comprises forming a fibrous paper web and treating the fibres with a water-soluble cationic, thermosetting, epichlorohydrin-containing resin, a non-viscose film-forming material and a polyalkylene imine.

The present invention also provides paper, for example casing paper and teabag paper, when prepared by the afore-
said process.

The invention also provides casing material (casing tubes or skins) prepared from the said casing paper by a process comprising the application of a caustic viscose solution and subsequent acid regeneration, as well as infusion pouches, e.g. teabags (especially herbal teabags) and spice bags, manufactured from the said teabag paper.

The fibrous web may be formed by papermaking techniques that are conventional in the art. The choice of the actual method and indeed of the composition of the fibres will depend upon the intended use of the paper.

For example, the fibrous webs used in making casing paper are conventionally composed of natural vegetable fibres of pure cellulose and are preferably composed of long lightweight nonhydrated fibres of the Musa type, particularly hemp fibres of the Manila or abaca hemp variety. Webs made from this material are generally soft porous papers of uniform texture and thickness and have for some time found wide acceptance as the primary fibre component of the fibrous base webs used in casing manufacture.

In the case of teabag paper, a preferred fibrous material is a lightweight permeable paper made from any of abaca pulp, sisal pulp, regenerated rayon, esparto grass pulp and long-fibred chemical wood pulp. Of course, in order to permit the fabrication of a heat-sealed pouch the fibrous material may comprise heat-sealable fibres of such materials as polyolefins, e.g. polyethylene or polypropylene, or vinyl chloride and vinyl acetate polymers. The basis weight of the paper of the invention is usually from 8 to 12 pounds (214 and 36480).
It will be appreciated that the invention lies in the selection of the three specified treating agents; these have been found capable of acting in an unexpected, synergistic manner to impart an excellent alkaline wet strength and good absorbency to paper webs.

It should be noted that the three treating agents - namely the water-soluble, cationic, thermosetting, epichlorohydrin-containing resin, the non-viscose film-forming material and the polyalkylene imine - may be applied to the fibres at any stage of the paper-making process, even prior to the formation of the web, e.g. they may be added to the fibre slurry. However, the said agents are preferably applied to the fibres after the formation of the fibrous web or sheet. Preferably, the three treating agents are applied simultaneously; this can be effected conveniently by passing the fibrous web through an aqueous bath containing all three agents, or by spraying the agents on the web.

Preferably, the water-soluble, cationic, thermosetting resin is an epichlorohydrin-containing resin.

A preferred class of cationic, thermosetting resins are the water-soluble polyamide-epichlorohydrin resins (PAE), such as those prepared by the reaction of epichlorohydrin with polyamides derived from polyalkylene polyamines (e.g. diethylene triamine or tetra-ethylene pentamine) and saturated or unsaturated aliphatic or aromatic polycarboxylic acid containing 3 to 10 carbon atoms (e.g. adipic, itaconic, or succinic acid). Resins of this type are described in U. S. Patents No. 2,926,116, No. 2,926,154 and No. 3,125,552.

In U. S. Patent 7,535,288 there are disclosed suitable resins that are derived from epichlorohydrin and polyamides.
that are the reaction products of certain amino poly-
carboxylates and derivatives thereof, especially ethylene-
diamine-tetraacetic acid or diethylenetriamine pentaaetate
cid, and polyalkylene polyamines, especially penta-
ethylenehexamine.

Suitable polyamide-epichlorohydrin resins are also
disclosed in U.S. Patent 3,526,608. Here the polyamides
are obtained from the reaction of polyalkylene polyamines,
and either iminodiacetic acid, N,N'-piperazinediacetic
acid, N-alkyliminodiacetic acids, N,N'-dialkylethylenediacetic
amine-N,N'-diamine-N,N'-diacetic acids, or their corresponding lower
alkyl esters. Preferred polyalkylene polyamines are
diethylenetriamine, triethylenetetramine, and tetra-
ethylenehexamine and mixtures of these polyamines.

The resins disclosed in U.S. Patent 3,565,754 are
provided by reacting epichlorohydrin with polyamides
derived from nitrilotriacetic acid and polyalkylene poly-
amines containing two primary amine groups and at least
one secondary amine group in which the nitrogen atoms are
linked together by groups having the formula \(-C_nH_{2n}\)
where \(n\) is a small integer generally having a value of 2
and the number of such groups in the molecule ranges from
2 to 19 and preferably up to 6. The nitrogen atoms may
be attached to adjacent carbon atoms in the group \(-C_nH_{2n}\)
or to carbon atoms further apart, but not to the same carbon
atom.

As disclosed in U.S. Patent 3,816,556, water-soluble,
cationic, thermal-setting polyamide-epichlorohydrin resins
can be converted into polylsalts by reaction with certain
anionic, water-soluble polyacrylamides. Such polylsalts
are useful in this invention.

Other epichlorohydrin-containing resins which can be used include the reaction products of epichlorohydrin with polymers derived from polyacrylamide and polyamines such as ethylenediamine (U.S. Patent 3,507,857); with hydrolysed polyvinylimidazoline (J.S. Patent 3,640,936); with polyamine-polyamide polymers derived from certain heterocyclic dicarboxylic acids, aminocarbonyl compounds, di- or polyalkylene polyamines and epichlorohydrin (U.S. Patent 3,761,340); and with polyethyleneimine (U.S. Patent 3,520,774).

The preferred film-forming material is hydroxyethyl cellulose (HEC), although other cellulose derivatives, such as methyl cellulose, hydroxypropyl cellulose, and sodium carboxymethyl cellulose (CMC), could be used. Other film-forming agents include polyvinyl alcohol, starch, starch derivatives, natural gums and other, water-soluble, polymers.

Polyethyleneimine (PEI) - which is the preferred polyalkylene imine - is a water-soluble, cationic homopolymer which may be synthesised by acid-catalysed polymerisation of ethyleneimine. It is known in the paper-making art as a retention aid to prevent short fibres passing through the paper-making wire. It has also been suggested for us in certain specialist papers where it may slightly increase the wet strength.

The water-soluble, cationic, thermosetting epichlorohydrin-containing resin is usually applied in an amount to give 0.05% to 4%, by weight of the fibrous web, on a dry weight basis. For casing paper, the preferred amount is 0.3% to 4%, especially 1.2% to 2.4%. For teabag paper the preferred amount is 0.05% to 1%, especially 0.25% to 0.5%.
The non-viscose film-forming material is usually applied in an amount to give 0.5% to 8%, by weight of the fibrous web, on a dry weight basis.

For casing paper, the preferred amount is 0.6% to 8%, especially 2.4% to 4.8%. For teabag paper, the preferred amount is 0.5%, to 5%, especially 1.0% to 3.0%.

The polyalkylene imine is usually applied in an amount to give 0.05% to 2.0%, by weight of the fibrous web, on a dry weight basis.

This range is entirely suitable for casing paper, although the preferred amount is 0.4% to 8%. For teabag paper, the preferred amount is 0.1% to 1.0%, especially 0.2% to 0.3%.

Prior to its impregnation with a caustic viscose solution or the like in order to form the final casing material, the casing paper of the present invention may be treated in order to improve its absorbency yet further. One preferred method is to subject at least one side of the web to a corona discharge treatment at an energy density of at least 0.5 Watt-min/ft$^2$ of web surface. The usual level will exceed 1.5 Watt-min/ft$^2$ and is preferably 5 to 40 Watt-min/ft$^2$.

This treatment is described in B.W. Conway and J.P. Molinari's U.S. Patent Application 636,082 filed 28th November, 1975.

The present invention is illustrated by the following Examples.

Example 1

A quantity of casing fibrous sheets was manufactured without any chemical treatment. Various blends of the chemical combinations containing Kynene 557 (trade name of a watersoluble, polyamide-epichlorohydrin resin marketed by Hercules
Powder Company), Polymin P (trade name of a polyethyleneimine marketed by BASF, Germany) and Natrasol 250L (trade name of a commercially available hydroxy ethyl cellulose) were applied to the untreated sheet by means of a laboratory size-press.

A standard viscose-treated sheet was also manufactured on a pilot-scale papermaking machine for use as a standard. The sheets were then tested for alkaline tensile strength and absorbency. Eventually a suitable combination (Formulation A) was accepted for a full-scale production run.

This run was made by applying the chemical mixture by means of a size-press unit which was immediately followed by a drying apparatus consisting of a series of steam filled cylinders.

A second production run was also made where the Polymin P concentration was reduced by 50% in the chemical mixture (Formulation B).

The resultant sheets were then tested and the results compared with those obtained with the standard casing sheet. The alkaline resistance was measured by recording the tensile strength on wet strips 25.4 mm wide after a 20 minute soak in 6% sodium hydroxide.

Absorbency of the sheet was also measured, this being measured as the time taken for water to climb 25.4 mm up a vertical strip of the paper. The greater absorbency, the less time it would take to climb the sheet.

Of course, the test actually measures the absorbency of the sheet in the plane of the sheet, whereas in the production of casing tubes by passing a tube of casing paper through a caustic viscose solution, it is the absorbency in the direction perpendicular to the plane of the sheet that
is important. However, it is postulated that the cited test correlates well with the absorbency in this direction. The penetration of an aqueous solution through a web — in whatever direction — is due to the same mechanism, i.e. capillary action, which in turn depends on the wettability of the capillary walls. It is furthermore thought that the wettability of the capillary walls may change with time in treated webs, particularly the in prior-art viscose-treated webs. Thus, the "water climb" test was made on samples shortly after their production and also on samples that had been aged for three months.

The formulations used are shown below

1. **Standard Viscose**

Viscose solution containing 1.75% cellulose.

Regenerated with sulphuric acid at a concentration of 0.4 Normal.

2. **Formulation A**

<table>
<thead>
<tr>
<th>Water (litres)</th>
<th>492.1 (37.8°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natrasol 250L (kg)</td>
<td>12</td>
</tr>
<tr>
<td>Mix Time (mins)</td>
<td>30</td>
</tr>
<tr>
<td>Volume Make up (litres)</td>
<td>To 613.2</td>
</tr>
<tr>
<td>Kymt 557 (litres)</td>
<td>60.6</td>
</tr>
<tr>
<td>Mix Time (mins)</td>
<td>5</td>
</tr>
<tr>
<td>Polymin P (litres)</td>
<td>5.46</td>
</tr>
<tr>
<td>Mix Time (mins)</td>
<td>10</td>
</tr>
<tr>
<td>Volume Make up (litres)</td>
<td>To 681.4</td>
</tr>
</tbody>
</table>

**Formulation B**

<table>
<thead>
<tr>
<th>Water (litres)</th>
<th>492.1 (37.8°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natrasol 250L (kg)</td>
<td>12</td>
</tr>
<tr>
<td>Mix Time (mins)</td>
<td>30</td>
</tr>
<tr>
<td>Volume Make up (litres)</td>
<td>To 613.2</td>
</tr>
<tr>
<td>Kymt 557 (litres)</td>
<td>60.6</td>
</tr>
<tr>
<td>Mix Time (mins)</td>
<td>5</td>
</tr>
<tr>
<td>Polymin P (litres)</td>
<td>2.73</td>
</tr>
<tr>
<td>Mix Time (mins)</td>
<td>10</td>
</tr>
<tr>
<td>Volume Make up (litres)</td>
<td>To 681.4</td>
</tr>
</tbody>
</table>

The results are shown in the table below
<table>
<thead>
<tr>
<th>Properties</th>
<th>Units</th>
<th>Standard Viscose</th>
<th>Formulation A</th>
<th>Formulation B</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sheet weight</td>
<td>gm</td>
<td>27.1</td>
<td>27.9</td>
<td>27.7</td>
</tr>
<tr>
<td>MD Dry Tensiles</td>
<td>gm/25.4mm</td>
<td>4366</td>
<td>4766</td>
<td>4404</td>
</tr>
<tr>
<td>MD Wet Tensiles</td>
<td>gm/25.4mm</td>
<td>940</td>
<td>738</td>
<td>590</td>
</tr>
<tr>
<td>Dry Elongation</td>
<td>%</td>
<td>3.2</td>
<td>2.16</td>
<td>2.5</td>
</tr>
<tr>
<td>Dry Toughness</td>
<td>cm/gm/sq.cm</td>
<td>95</td>
<td>85</td>
<td>75</td>
</tr>
<tr>
<td>Wet Elongation</td>
<td>%</td>
<td>4.8</td>
<td>2.7</td>
<td>2.1</td>
</tr>
<tr>
<td>Wet Toughness</td>
<td>cm/gm/sq.cm</td>
<td>29</td>
<td>13</td>
<td>8.8</td>
</tr>
<tr>
<td>Porosity</td>
<td>l/min</td>
<td>208</td>
<td>254</td>
<td>241</td>
</tr>
<tr>
<td>Burst</td>
<td>kg/sq.cm</td>
<td>1.69</td>
<td>1.23</td>
<td>1.34</td>
</tr>
<tr>
<td>MD Caustic Tensile</td>
<td>gm/25.4mm</td>
<td>270</td>
<td>610</td>
<td>495</td>
</tr>
<tr>
<td>25.4mm Water secs. Climb</td>
<td></td>
<td>10</td>
<td>8.5</td>
<td>6.0</td>
</tr>
<tr>
<td>25.4mm Water secs.</td>
<td></td>
<td>20</td>
<td>8.5</td>
<td>6.0</td>
</tr>
<tr>
<td>climb after 3 months</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

From the table of results, one can draw the following conclusions:

The chemical formulations consisting of polyamide-epichlorohydrin resin, polyethylene-imine and hydroxy ethyl cellulose had significantly increased the alkaline tensile compared with the standard sheet.

The same chemical formulation has also increased the water absorbency compared with the standard, after ageing.

Example 2

Handsheets of a typical teabag base furnish, of defibred abaca and softwood, were produced to give untreated sheets at
a basis weight (weight per unit area) of 34 gsm. (The value of the basis weight of the laboratory handsheets is not itself of significance; however, all the experimental results are relative to this basis weight in order to permit direct comparison).

Sheets of said paper were saturated with an aqueous liquor containing both Kymene 557 (the trade name of a polyamide-epichlorohydrin resin marketed by Hercules Powder Company) and CMC. This simulated, on a laboratory scale, one of the prior-art treatments discussed hereinbefore.

The treated webs were dried and then tested for their wet tensile strength and their alkaline wet tensile strength. The wet tensile strength was measured on strips of paper cut to a 15 mm width and saturated with water. The alkaline wet tensile strength was measured on strips of paper, 15 mm wide, saturated with 1% solution of sodium hydroxide. The results obtained were regarded as the standard and subsequent results from different treatments were compared to the standard.

Further untreated handsheets, as described above, were saturated with an aqueous liquor containing Kymene 557, HEC and Polymin SN (the trade name of a polyethylene imine marketed by BASF, Germany). The sheets were dried to give treated paper sheets in accordance with this invention. The treated paper was tested for wet tensile strength and alkaline wet tensile strength, in the manner described above.

The results (all corrected to a handsheet basis weight of 34 gsm) are given in Table 2 below, which also specifies the ratio (by weight) of the various treating agents (as solids) used in each test.

The teabag papers according to the invention were tested
to determine whether they would impart an undesirable taste to beverages. The papers were found to be as good in this respect as the prior-art papers containing just Kymene 557 and CMC.

**Example 2**

Untreated teabag paper was first treated with a 0.1% solids solution of Polyvinyl P and then treated with a mixture of Kymene 557 and HEC in the ratio of 1:2.9. The total pick up of the various solids was approximately the same as in Example 1. This gave a certain improvement in the alkaline wet strength, although this improvement was not quite as great as that obtained in Example 2, as may be seen from the results given in Table 2.

The teabag papers according to the invention were tested to determine whether they would impart an undesirable taste to beverages. The papers were found to be as good in this respect as the prior-art papers containing just Kymene 557 and CMC.
<table>
<thead>
<tr>
<th></th>
<th>Base Handsheet + Kymene/CMC at a ratio of 1/8.8</th>
<th>Base Handsheet + Kymene/HEC/ Polymin SN at a ratio of 1/8.8/5</th>
<th>Base Handsheet + Kymene/HEC/ Polymin SN at a ratio of 1/2.9/1.67</th>
<th>Base Handsheet treated according to Example 3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Alkaline Wet</strong></td>
<td>55</td>
<td>350</td>
<td>350</td>
<td>150</td>
</tr>
<tr>
<td><strong>Tensiles</strong></td>
<td>70</td>
<td>215</td>
<td>275</td>
<td>135</td>
</tr>
<tr>
<td>gms/15 mm</td>
<td>80</td>
<td>350</td>
<td>625</td>
<td>155</td>
</tr>
<tr>
<td></td>
<td>100</td>
<td>245</td>
<td>400</td>
<td>160</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td></td>
<td></td>
<td>145</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td></td>
<td></td>
<td>180</td>
</tr>
<tr>
<td><strong>Average Value</strong></td>
<td>74</td>
<td>290</td>
<td>429</td>
<td>154</td>
</tr>
<tr>
<td><strong>Wet Tensiles</strong></td>
<td>500</td>
<td>300</td>
<td>475</td>
<td>250</td>
</tr>
<tr>
<td>gms/15 mm</td>
<td>600</td>
<td>320</td>
<td>450</td>
<td>300</td>
</tr>
<tr>
<td></td>
<td>450</td>
<td>480</td>
<td>600</td>
<td>245</td>
</tr>
<tr>
<td></td>
<td></td>
<td>450</td>
<td>600</td>
<td>250</td>
</tr>
<tr>
<td><strong>Average Value</strong></td>
<td>517</td>
<td>388</td>
<td>531</td>
<td>269</td>
</tr>
<tr>
<td></td>
<td>Base Handsheet + Kymene/CMC at a ratio of 1/8.8</td>
<td>Base Handsheet + Kymene/HEC/Polymin SN at a ratio of 1/8.8/5</td>
<td>Base Handsheet + Kymene/HEC/Polymin SN at a ratio of 1/2.9/1.67</td>
<td>Base Handsheet treated according to Example 3</td>
</tr>
<tr>
<td>-------------------------</td>
<td>-----------------------------------------------</td>
<td>---------------------------------------------------------------</td>
<td>---------------------------------------------------------------</td>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>Alkaline Wet</td>
<td>55</td>
<td>350</td>
<td>350</td>
<td>150</td>
</tr>
<tr>
<td></td>
<td>70</td>
<td>215</td>
<td>275</td>
<td>135</td>
</tr>
<tr>
<td>Tensiles</td>
<td>80</td>
<td>350</td>
<td>625</td>
<td>155</td>
</tr>
<tr>
<td>gms/15 mm</td>
<td>100</td>
<td>245</td>
<td>400</td>
<td>160</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td></td>
<td></td>
<td>145</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td></td>
<td></td>
<td>180</td>
</tr>
<tr>
<td>Average Value</td>
<td>74</td>
<td>290</td>
<td>429</td>
<td>154</td>
</tr>
<tr>
<td>Wet Tensiles</td>
<td>500</td>
<td>300</td>
<td>475</td>
<td>250</td>
</tr>
<tr>
<td>gms/15 mm</td>
<td>600</td>
<td>320</td>
<td>450</td>
<td>300</td>
</tr>
<tr>
<td></td>
<td>450</td>
<td>430</td>
<td>600</td>
<td>245</td>
</tr>
<tr>
<td></td>
<td>450</td>
<td>600</td>
<td></td>
<td>250</td>
</tr>
<tr>
<td>Average Value</td>
<td>517</td>
<td>388</td>
<td>531</td>
<td>269</td>
</tr>
</tbody>
</table>
Example 4

Untreated teabag paper having a basis weight of 12 gsm was obtained using a commercial paper-making machine (305 cm wide) operating at a constant speed of 200 metres per minute. The fibrous webs were treated with Kymene 557, HEC and Polymin SN, in the manner described in Example 2, and were tested for their alkaline wet tensile strength and wet tensile strength in both the machine direction (MD) and cross direction (CD).

The results are given in Table 3 below. Again the ratios by weight of the various treating agents (as solids) are specified. The table also shows the results obtained using standard prior-art teabag papers, one having a Kymene 557/CMC treatment and the other having a viscose treatment. The pickup of solids in these prior-art papers was comparable to that of the paper according to the invention.

The papers were also tested, using a panel of expert tasters, in order to determine the taste that they would impart to beverages. Each sample was rated on a 1-5 scale, 1 indicating an absence of imparted taste (the ideal and 5 indicating an intolerable taste. The paper treated according to this invention and the prior-art Kymene/CMC-treated paper both scored an average of 1.5, whereas the viscose-treated paper had the less satisfactory average score of 2.5.
<table>
<thead>
<tr>
<th>Untreated Web Off Machine + Kymene/HEC/ Polymin SN at a ratio of 1/8.8/5</th>
<th>Untreated Web + Kymene/HEC/ Polymin SN at a ratio of 1/2.9/1.67</th>
<th>Untreated Web + Kymene/HEC/ Polymin SN at a ratio of 1/1.8/1</th>
<th>Untreated Web + standard Kymene/CMC treatment</th>
<th>Untreated Web + standard viscose treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkaline Wet</td>
<td>290</td>
<td>230</td>
<td>375</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>265</td>
<td>232</td>
<td>400</td>
<td>290</td>
</tr>
<tr>
<td>Tensiles</td>
<td>290</td>
<td>220</td>
<td>385</td>
<td>265</td>
</tr>
<tr>
<td></td>
<td>270</td>
<td>240</td>
<td>385</td>
<td>288</td>
</tr>
<tr>
<td>gm/15 mm</td>
<td>300</td>
<td>215</td>
<td>460</td>
<td>312</td>
</tr>
<tr>
<td></td>
<td>285</td>
<td>218</td>
<td>315</td>
<td></td>
</tr>
<tr>
<td>Average Value</td>
<td>283</td>
<td>225</td>
<td>286</td>
<td>286</td>
</tr>
<tr>
<td>Wet Tensiles</td>
<td>375</td>
<td>275</td>
<td>510</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>375</td>
<td>280</td>
<td>450</td>
<td>375</td>
</tr>
<tr>
<td>gms/15 mm</td>
<td>320</td>
<td>275</td>
<td>425</td>
<td>335</td>
</tr>
<tr>
<td></td>
<td>275</td>
<td>240</td>
<td>540</td>
<td>375</td>
</tr>
<tr>
<td></td>
<td>360</td>
<td>245</td>
<td>450</td>
<td>380</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>215</td>
<td>470</td>
<td>280</td>
</tr>
<tr>
<td>Average Value</td>
<td>350</td>
<td>255</td>
<td>474</td>
<td>350</td>
</tr>
<tr>
<td>Untreated Web + Kymene/HEC/ Polymin SN at a ratio of 1/8.8/5</td>
<td>Untreated Web + Kymene/HEC/ Polymin SN at a ratio of 1/2.9/1.67</td>
<td>Untreated Web + standard Kymene/CMC treatment</td>
<td>Untreated Web + standard viscose treatment</td>
<td></td>
</tr>
<tr>
<td>-------------------------------------------------------------</td>
<td>-------------------------------------------------------------</td>
<td>------------------------------------------------</td>
<td>------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>M.D.</strong></td>
<td><strong>C.D.</strong></td>
<td><strong>M.D.</strong></td>
<td><strong>C.D.</strong></td>
<td><strong>M.D.</strong></td>
</tr>
<tr>
<td>Alkaline Wet</td>
<td>290</td>
<td>230</td>
<td>375</td>
<td>275</td>
</tr>
<tr>
<td>Tensiles</td>
<td>265</td>
<td>232</td>
<td>400</td>
<td>290</td>
</tr>
<tr>
<td>gm/15 mm</td>
<td>290</td>
<td>220</td>
<td>385</td>
<td>265</td>
</tr>
</tbody>
</table>

| **Average Value** | **283** | **225** | **286** | **286** | **455** | **365** | **43.5** | **12.6** | **151** | **33** |

| Wet Tensiles   | 375  | 275  | 510  | 355  | 450  | 460  |
| gms/15 mm      | 375  | 280  | 450  | 375  | 470  | 325  |

| **Average Value** | **350** | **255** | **474** | **350** | **453** | **396** |
From the results of the foregoing Examples 2 - 4, the following conclusions could be drawn.

(i) The alkali resistance of teabag paper can be substantially increased by treating the web with a hydroxy ethyl cellulose/polyamide-epichlorohydrin/polyethylene imine mix when compared to a carboxy methyl cellulose/polyamide-epichlorohydrin mix.

(ii) It was also found that the treatment with Kymene 557/Polymin SN/HEC gave the same taste neutrality that is a feature of the prior-art Kymene 557/CMC-treated paper.

(iii) The alkali resistance (measured as the alkaline wet tensile strength) of teabag paper increased as the proportion of Kymene 557 approached that of polyethylene imine and hydroxy ethyl cellulose.

Note: Polyethylene imine is also marketed in other forms, such as Polymin P. Polymin SN and Polymin P give similar results when used in the present invention.
Example 5

Samples of paper were treated with various treating agents and then soaked for 20 minutes in a 6% by weight, aqueous sodium hydroxide solution. The tensile strength of each sample was then measured.

The paper used was a pure abaca fibre base web obtained from a commercial production run used for the manufacture of casing paper. Such a paper was characterised by being virtually unmodified by treatments such as beating. Thus, the bonded areas of such a sheet prior to saturation with the treating agent was very small, with the result that the initial web was very weak.

The treating agents were as follows:

A : HEC/Xymene 557/Polymin P (ratio 1:1:1 by weight)
B : Polymin SN
C : Polymin P
D : Kynene 557
E : HEC

The results are shown below in Table 4.
### TABLE 4

<table>
<thead>
<tr>
<th>Treatment</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pick-up (dry-on-dry)</td>
<td>3%</td>
<td>3%</td>
<td>3%</td>
<td>3%</td>
<td>3%</td>
</tr>
<tr>
<td>Alkaline Wet</td>
<td>90</td>
<td>60</td>
<td>35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Tensiles (gm/15mm)</td>
<td>75</td>
<td>40</td>
<td>35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>after 20 minutes in 6% caustic solution</td>
<td>70</td>
<td>35</td>
<td>60</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>Average value</td>
<td>78.75</td>
<td>46.25</td>
<td>45</td>
<td>1.25</td>
<td>0</td>
</tr>
</tbody>
</table>

Note: the level of the solids pick-up at 3%, is the same as that specified in connection with Table III in U.S. Patent No. 3,484,256.
### TABLE 4

<table>
<thead>
<tr>
<th>Treatment:</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pick-up</strong> (dry-on-dry)</td>
<td>3%</td>
<td>3%</td>
<td>3%</td>
<td>3%</td>
<td>3%</td>
</tr>
<tr>
<td>Alkaline Wet</td>
<td>90</td>
<td>60</td>
<td>35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Tensile (gm/15mm)</td>
<td>75</td>
<td>40</td>
<td>35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>after 20 minutes in</td>
<td>70</td>
<td>35</td>
<td>60</td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td>8% caustic solution</td>
<td>80</td>
<td>50</td>
<td>50</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><strong>Average value</strong></td>
<td>78.75</td>
<td>46.25</td>
<td>45</td>
<td>1.25</td>
<td>0</td>
</tr>
</tbody>
</table>

**Note**: the level of the solids pick-up at 3%, is the same as that specified in connection with Table III in U.S. Patent No. 3,434,256.
From the results shown in Table 4, the following conclusions can be drawn.

(i) The combination of HEC, a polyamide-epichlorohydrin resin (i.e. Kymene 557) and a polyethylene imine (i.e. polymin P) produced a far greater alkaline wet strength than the individual agents used alone, in equivalent quantities. This demonstrates the surprising synergistic effect of the ternary treatment in accordance with the present invention.

(ii) HEC is completely unsuitable on its own as an alkaline wet-strength agent.

(iii) The polyamide-epichlorohydrin resin is also unsuitable on its own as an alkaline wet-strength agent.

The last conclusion seems, at first, to be contradictory to the results shown in Table III of U.S. Patent 3,484,356, to which reference was made earlier. However, it is probable that the paper web used in the aforesaid U.S. Patent was itself intrinsically stronger than the paper used in the present Example, which as stated above was very weak, since this showed more clearly the effect due to the various treating agents.

More importantly, there is no indication in the U.S. Patent that the test webs were soaked in the aqueous caustic solution for any longer than was necessary completely to wet the webs before testing them. Thus, very little time had elapsed in which the alkali could disrupt the binder. It is our experience that the alkaline wet strength of a fibrous web, as is currently used for casing paper, diminishes with time, presumably as the disruption of the binder by the alkali gradually proceeds.
For this reason, the alkaline wet tensiles were measured on samples that had been soaked in the alkaline solution for 20 minutes; after this period little further change in the alkaline wet tensile occurs. The results clearly show that after this period had elapsed the alkaline wet strength of the web treated with Kymene 557 diminished virtually to zero.

Our findings are corroborated by literature issued by Hercules Powder Company, who are leading manufacturers of polyamine-epichlorohydrin resins for the wet-strengthening of papers. They suggest that to reclaim such paper for re-use, one should add an alkali such as caustic soda in order to destroy the wet-strength.

Example 6

Samples of paper were treated with various agents in the manner described in Example 5, except that the total pick-up of solids during the treatment was 6% on a dry-on-dry basis. This is the level that might be expected in the commercial treatment of paper webs with wet-strength agents or the like. The dry tensile and alkaline tensile were measured (the latter being measured only after soaking the sample for 20 minutes in 6% aqueous NaOH).

The treating agents were as follows:

A : HEC/Polymin P (weight ratio 1:1)
B : Kymene 557/Polymin P (weight ratio 1:1)
C : HEC/Kymene 557/Polymin P (weight ratio 1:1:1)
D : HEC/Kymene 557/Polymin P (weight ratio 5:2:1)

The results are shown below in Table 5.
As can be seen from the above table, the ternary mixture according to the invention imparted a significantly higher alkaline wet strength to the web than the tested binary mixtures. It will be noted that, assuming uniform pick-up, composition B with 3% pick-up of each of the known wet-strength agents Kymene 557 and Polymin P still gives a lower alkaline wet-strength than composition C, with the equivalent of only a 2% pick-up of each of these two agents. (As shown in Example 5, HEC by itself is unsuitable as an alkaline wet-strength agent.) The superiority of the invention is even more marked in the case of composition D, which has even lower amounts of Kymene 557 and Polymin P.

It should also be noted that the high levels of Kymene 557 and Polymin P in the web treated with composition B may, in some circumstances, exceed the permitted levels. Thus, the ternary system used in the present invention is especially advantageous.
The claims defining the invention are as follows:

What we claim is:

1. A process for the production of paper, which process comprises forming a fibrous paper web and treating the fibres with a water-soluble cationic, thermosetting, epihalohydrin-containing resin, a non-viscose film-forming material and a polyalkylene imine.

2. A process according to claim 1, wherein the water-soluble cationic, thermosetting epihalohydrin-containing resin, the non-viscose film-forming material and the polyalkylene imine are applied to the fibres after the formation of the web.

3. A process according to claim 1, in which the water-soluble cationic, thermosetting epihalohydrin-containing resin, the non-viscose film-forming material and the polyalkylene imine are added to the fibre slurry prior to the formation of the web.

4. A process according to claim 1, 2 or 3 wherein the water-soluble cationic, thermosetting epihalohydrin-containing resin, the non-viscose film-forming material and the polyalkylene imine are applied to the fibres simultaneously.

5. A process according to any of claims 1 to 4 in which the film-forming material is hydroxy ethyl cellulose.

6. A process according to any of claims 1 to 5, in which the cationic, thermosetting resin is an epichlorohydrin-containing resin.

7. A process according to claim 5, wherein the cationic, thermosetting, epihalohydrin-containing resin is a polyoxido-epichlorohydrin resin.

8. A process according to any of claims 1 to 7, in which the polyalkylene imine is a polyethylene imine.

9. A process according to any of claims 1 to 3, wherein the cationic, thermosetting epihalohydrin-containing resin is
applied in an amount to give 0.05% to 4.0% by weight of the fibrous web, on a dry weight basis.

10. A process according to any one of claims 1 to 9, wherein the film-forming material is applied in an amount to give 0.5% to 8% by weight of the fibrous web, on a dry weight basis.

11. A process according to any of claims 1 to 10, wherein the polyalkylene imine is applied in an amount to give 0.05% to 2.0% by weight of the fibrous web, on a dry weight basis.

12. A process according to any of claims 1 to 8, for the production of casing paper, in which the treating agents are applied in amounts to give, by weight of the fibrous web on a dry weight basis, 0.3% to 4.0% of the epihalohydrin-containing resin, 0.6% to 8% of the film-forming material and 0.05% to 2.0% of the polyalkylene imine.

13. A process according to claim 12, in which the treating agents are applied in amounts to give, by weight of the fibrous web on a dry weight basis, 1.2% to 2.4% of the epihalohydrin-containing resin, 2.4% to 4.8% of the film-forming material and 0.4% to 0.8% of the polyalkylene imine.

14. A process according to any of claims 1 to 8, for the production of teabag paper, in which the treating agents are applied in amounts to give, by weight of the fibrous web on a dry weight basis, 0.05% to 1.0% of the epihalohydrin-containing resin, 0.5% to 5% of the film-forming material and 0.1% to 1.0% of the polyalkylene imine.

15. A process according to claim 14, in which the treating agents are applied in amounts to give, by weight of the fibrous web on a dry weight basis, 0.25% to 0.5% of the epihalohydrin-containing resin, 1.0% to 3.0% of the film-forming material and 0.2% to 0.3% of the polyalkylene imine.
16. A process according to claim 1, substantially as described in Example 1.

17. A process according to claim 1, substantially as described in Example 2, 3 or 4.

18. Paper whenever produced by a process according to any of claims 1 to 11.

19. Casing paper whenever produced by a process according to claim 12, 13 or 16.

20. Teabag paper whenever produced by a process according to claim 14, 15 or 17.


22. Paper according to claim 21, in which the film-forming material is hydroxy ethyl cellulose.

23. Paper according to claim 21 or 22 wherein the cationic, thermosetting, epihalohydrin-containing resin is a polyaide-epichlorohydrin resin.

24. Paper according to claims 21, 22 or 23 in which the polyalkylene imine is a polyethylene imine.

25. Paper according to any of claims 21 to 24, being teabag paper.

26. Infusion pouches whenever prepared from a paper according to claim 20 or 25.

27. An infusion pouch according to claim 26, containing a quantity of herbal tea.

28. Paper according to any of claims 21 to 24, being a casing paper.

29. Casing material whenever prepared from a casing paper.
according to claims 19 or 28 by impregnating the casing paper with a caustic viscose solution and subsequently regenerating the cellulose in the viscose by means of acid.

30. A processed meat product whenever encased in casing material according to claim 29.

DATED this 14th day of June 1977.

C.H. DEXTER LIMITED.