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Study of the joint strength between regenerated cellulose fibres and its influence on the sheet strength

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SUMMARY: The joint strength between single fibres was investigated directly by measurements of perpendicular cross bonded fibres and compared with the corresponding paper properties. Regenerated cellulose fibres, rayon, were used. They were carboxymethylated in order to introduce charges to the fibres. Two different treatments were used to prepare fibres with charges located on the surface of the fibres and charges located throughout the bulk of the fibres. The charge characteristics, with respect to total charge and surface charge of the fibre were determined. The swelling caused by these charges was analysed with respect to pore volume distribution (NMR measurements) and water retention capacity (WRV). Charge characterisation showed that surface charged fibres could be produced by pre-drying fibres prior to carboxymethylation. Furthermore, WRV and NMR-relaxation measurements showed that an increase in the surface charge had no significant influence on the swelling whereas an increase in bulk charges had a large effect. The strength of the joints and the paper tensile properties are most affected by the surface charges, probably due to an increase in the molecular contact area with increasing charge. The results indicated that increased surface softness increased the critical strain of paper made from these modified fibres.

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The main constituents of the paper dry strength are the joint strength between the fibres and the strength of the individual fibres. Together with the paper formation, these entities will determine the final strength of the paper. Network models to describe the properties of the paper have been presented by several authors (Rögdahl et al. 1986; de Ruvo 1986), and network theories for the description of paper strength have recently been published (Heyden et al. 2001). In these latter theories, the strength of joints between fibres and the strength of fibres are included as separate parameters. However, since fibres are small, few exact measurements have been made of either fibre strength or joint strength between the fibres. There has, over the years, been a considerable debate as to which of these two components is the most important for paper strength (see for example Davison 1972).

Fibre strength is determined both by morphology and the strength of the interaction between the different chemical components in the fibre wall. Earlier work has addressed this. For example, the influence on the fibre strength by the orientation of the cellulose microfibrils was described several decades ago (Broughton, Wang 1955; Page et al., 1972). During fibre processing, both the chemical composition of the fibre wall and the properties of the constituents in the fibre wall are changed and hence the properties of the fibres as raw material for papermaking will change (Wågberg, Annergren 1997). On a fibre level, fibre damage also affects the fibre strength. Page (1985) demonstrated that "curl, crimps, kinks and microcompressions" on the fibres significantly reduced the ability to fully utilise the fibre properties in the paper.

The second and perhaps even more important component in the development of paper dry strength is the development of the fibre/fibre joint strength. The word "joint" is used in this work instead of "bond" in order to suggest adhesion between the fibres to be the primary factor affecting paper strength rather than chemical bonding between the fibres. The description of the joints between fibres is not unique to paper but generally valid for adhesive contact between materials in general (Caulfield 1973; Kinloch 1980). To develop strong joints between fibres, several different factors have to be taken into account. The fibres need to come into close proximity with each other, brought about by the capillary forces during pressing and drying. Van der Waals forces and electrostatic forces also affect the interaction between the fibres in the fibre/water/fibre contact zone (Wågberg, Annergren 1997). The properties of the molecular contact zone depend on the softness of the fibre surface. Increased swelling is an indication of a decrease in the transverse elastic modulus of the surface (Scallan, Tigerström 1992; Lindström 1986). This in turn leads to a larger molecular contact zone (Emerton 1957; Paavilainen 1993). Paper strength has also been shown to be highly dependent of the outermost layers of the fibre (Barzyk 1997; Laine et al. 1996). This may be probed by using for example Atomic Force Microscopy (AFM) as described by Nilsson et al. (2001).

When the fibres are in molecular contact the joint starts to develop and this is followed by a decrease of the specific surface area of the paper. This has been interpreted in terms of both inter- and intra-fibre bonding. The interaction most commonly suggested in the literature to explain the development of paper strength is probably hydrogen bonding. Results supporting this theory have been obtained by drying cellophane films together in close contact (Clark 1933) and the cohesion developed in a dried sheet can, according to McKenzie (1955), largely be attributed to hydrogen bonding between hydroxyl groups of carbohydrate chains. Considering the rough surfaces of wood fibres, the distance of 2-3 Å needed to
form a hydrogen bond has led to a debate regarding the validity of the hydrogen bond theory. New data on the surface softness of the fibres (Scallan, Tigerstrom 1992; Nilsson et al 2001) however show the fibre surface to be a highly swollen gel where the macromolecules have high mobility. With highly mobile molecules in the outer layer of the surface, the molecules (amorphous hemicellulose or added polymers for example) may be able to interdiffuse and create strong joints between the fibres (McKenzie 1984). The interaction between these molecules may be of a hydrogen bond type. Pelton and co-workers (Pelton et al, 2000) have presented results supporting McKenzie’s theory, but the focus in this investigation was more on polymer compatibility than on hydrogen bonding. By studying the compatibility of adsorbed polymers during sheet consolidation, they found that polymer compatibilities at the fibre surface was important for the development of paper strength (Pelton et al, 2000).

The strength of single fibre joints has been measured in only few studies. The joint strength is usually evaluated by application of the Page equation (Page 1969), where papers from the same type of fibres are pressed to different densities and a so-called related bonded area (RBA) is estimated by measuring the light scattering of these papers. Utilising the Page equation and the relative bonded area, the tensile strength of the paper and data concerning fibre geometry, a specific bond strength or a shear bond strength between the fibres can be calculated. Another more direct way of evaluating the joint strength between the fibres would be to measure directly the interaction between fibres in a sheet. Davison (1972) assessed the joint strength between fibres by pulling single fibres out from a paper and Stratton (1990) used a more direct approach and measured the joint strength directly in a perpendicular cross of two single fibres.

In order to clarify the importance of fibre interactions and molecular contact area on joint and paper strength in the present work, rayon fibres of different cross sectional area were chemically modified and the method of using single fibre crosses as described by Stratton (1990) was selected to determine the joint strength between fibres. To decrease the natural variations that occur in native fibres, regenerated cellulose fibres were investigated with CP/MAS(13)C-NMR (at 292 ± 1 K) on a Bruker AMX-300 instrument operating at 7.05 T (Hult 2001). It is known that the charge of cellulosic fibres is essential for their physical properties and hence also for their papermaking properties. The total charge of the fibres was determined by conductometric titration (Katz et. al. 1984) and the surface charge by polyelectrolyte adsorption (Wågberg et al. 1985; Winter et. al. 1986). In this method a high molecular polydimethylallylammonium chloride (polyDADMAC) with a molecular mass of 1,2*10^6 g/mol was used. The adsorption was carried out at pH 8.

To obtain a description of the fibre swelling the water retention capacity (Scallan, Carles 1972) and the pore structure of the fibres were analysed. To determine the pore structure the fibres were subjected to NMR relaxation measurements on a Bruker Avance spectrometer (5.87 T magnet. 1H resonance frequency 38.4 MHz) (Andreasson, Wågberg 2001). According to (Li et al, 1993) this method provides a good description of the pore size distribution of the pores within the fibre wall.

Handsheets were prepared in the Rapid-Kothen Blättbildner (ISO 5269-2). The thickness of the sheets was measured according to standard SS 843009. Force-displacement curves of the sheets were measured according to SCAN-P 67-93.

The fibre crosses were prepared according to the method initially described by Stratton and Colsson (1990).

**Materials and methods**

The fibres used in the experiments were rayon tow bright fibres from Svenska Rayon diameters of about 30 and 60 pm (5 and 20 dtex) cut by Bernhardt Stefferdt AB (Gothenburg, Sweden) to lengths of 4-mm. The fibres were thoroughly washed before use, first refluxed for 1 hour in deionised water at 10% pulp concentration then extracted in ethanol at 10% pulp concentration for 1 hour. Thereafter the fibres were reslushed in deionised water and dewatered. The washing continued until the first and last washing water had the same amount of total organic chloride (SS 028199 on a Shimadzu 5050A) and until Soxhlet extraction of the fibres at 10 g per litre in ethanol for 6 hours, showed a remaining amount of ethanol-extractable compounds being less than 0.01 mg/g.

To obtain two different charge distributions of the fibres, bulk-charged and surface-charged fibres, a modified carboxymethylation was used (Fors 2000). To achieve bulk-charged fibres, the washed pulp was solvent exchanged to ethanol (spectrophotometric grade) prior to the carboxymethylation. The solvent exchange from water to ethanol was carried out in six steps with a four-hour equilibrium time for each step. To achieve a surface-carboxymethylated pulp the fibres were hornified (dried from water at 105°C) before carboxymethylation. The carboxymethylation was then performed according to Wålecka (1956). The surface-carboxymethylated fibres were dried in the sodium form at 50 % R.H whereas the bulk-carboxymethylated fibres were not dried at all prior to use.

To study the compatibility of adsorbed polymers during sheet consolidation, they found that polymer compatibilities at the fibre surface was important for the development of paper strength (Pelton et al. 2000). The strength of single fibre joints has been measured in only few studies. The joint strength is usually evaluated by application of the Page equation (Page 1969), where papers from the same type of fibres are pressed to different densities and a so-called related bonded area (RBA) is estimated by measuring the light scattering of these papers. Utilising the Page equation and the relative bonded area, the tensile strength of the paper and data concerning fibre geometry, a specific bond strength or a shear bond strength between the fibres can be calculated. Another more direct way of evaluating the joint strength between the fibres would be to measure directly the interaction between fibres in a sheet. Davison (1972) assessed the joint strength between fibres by pulling single fibres out from a paper and Stratton (1990) used a more direct approach and measured the joint strength directly in a perpendicular cross of two single fibres.

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except that the fibres were not dyed. After preparation, the fibre crosses were stored at room temperature until they were tested. An Environmental Scanning Electron Microscope (ESEM 2020) from Electro Scan, USA was used to study the behavior of rayon fibers and the joints between the fibres during loading. A tensile testing stage was designed at the Swedish Pulp and Paper Research Institute (STFI) by modifying the Electroscan standard tensile stage, see Fig. 1. During tension, both stage jaws moved outwards simultaneously while the test piece remained centered below the detector to permit imaging. A water-cooled unit was attached to the stage in order to maintain a stable test piece temperature and to remove the local heat caused by the electron beam in long-term experiments. The load cell from Sensotec had a working range of 0-150 gms.

The sample was glued to the table of the load bench with Loctite 401" glue. The glue was allowed to dry for five minutes before the bench was placed in the ESEM. The conditions inside the microscope were 27°C and 4 torr, approximately 20% RH. The sample was conditioned for 15 minutes before measurement. The microscope was operating at 15 kV accelerating voltage with a 1 mm working distance of the stage, the displacement rate was 1μm/s.

**Results and discussion**

The influence of the carboxymethylation reaction on the cellulose crystallinity was examined using CP/MAS 13C-NMR. There was no difference in the spectra between a treated and an untreated fibre, which indicates that the cellulose was not affected on the supermolecular level by the carboxymethylation reaction and was mainly of the cellulose II crystalline form.

According to the STFI fibremaster measurements the weighted fibre length was 4.00-4.08 mm for both the 5dtx and 20dtx. The fibrewidths were 36.3-36.6 pm for the 5dtx fibres and 65-67.8 μm for the 20dtx fibres, and the aspect ratio of the fibre cross section was 92.6-96.8% for both the 5dtx and the 20dtx fibres respectively. These values were not affected by the carboxymethylation reactions.

Charge and charge distribution of the fibres are important properties for both joint and paper properties since they affect the swelling of the fibre surface and hence the transverse elastic modulus of the surface. This in turn will affect the molecular contact area and may also affect the molecular mobility at the surface. A large difference in total charge was found for fibres that had been hornified and ethanol-exchanged before carboxymethylation. The hornified fibres reached a plateau at 100 μeq/g whereas the solvent-exchanged fibres reached a level of 550 μeq/g, see Fig. 2a. Fig. 2h on the other hand shows that the amount of adsorbed high molecular polyDADMAC reaches the same level for the two types of fibres, despite the five times higher total charge of the solvent-exchanged fibre. These results indicate that the surface of these fibres becomes saturated with charges at a level of about 100 μeq/g.

![Fig. 1. Rayon fibres joined together in a fibre cross. Joint strengths analyzed in tensile stage.](image1.png)

![Fig. 2a. and h. Effect of carboxymethylation on hornified and solvent-exchanged fibres measured as (a) total charge and (b) polyelectrolyte adsorption.](image2.png)

The total amount of charges normally has a large influence on the swelling or water-holding capacity (WRV) of fibres (Grignon, Scallan, A.M., 1980; Lindstrom, Carlsson 1978) as well as the pore size distribution inside the fibre wall (Li 1993). As shown in Fig. 3, bulk-charged fibres gave a significantly higher WRV and a higher average pore size and a wider pore volume distribution compared to the surface charged fibres. This is most likely due to the internal swelling of the bulk-charged fibre. As indicated in Fig. 3, the hornified fibres did not show any significant swelling since there was no change in WRV or in pore volume distribution. This most probably depends on the location of the charges on the surface of the fibres, only creating a softer surface of the fibres in water.
break of the fibres will not affect the strain properties of the sheets made from these fibres.

As discussed in the introduction, the joint strength between fibres is determined by several factors, such as molecular contact area and mixing of materials in the external parts of the adjacent fibres etc. The final network strength is naturally also dependent on fibre strength, fibre orientation and chemical interactions to mention only a few factors.

In order to establish whether or not there is a link between fibre properties, joint strength and paper strength, both single fibre crosses and paper sheets were prepared from the same kind of fibres. The results for fibre crosses and paper sheets are shown in Fig. 5. The total charge of the fibres has been recalculated into total charge per square meter, and used as a basis for comparison of fibres with different dimensions. Dimensions of the fibres and the mass per 10 000 meters of the fibres were used for this calculation where the fibres were assumed to have a cylindrical shape. As clearly shown in Fig. 5 the same trend in strength can be found for both the joints and the paper. In turn this shows that surface charges are by far most important for the development of joint and paper strength than the bulk charges. These results are in accordance with earlier measurements of papers with differences in charge distribution of the fibres (Barzyk 1997; Laine 1996).

As shown in Fig. 6, the strain at break of the paper
increases with the charge of the fibres. This indicates that the critical strain of each individual joint did increase since the critical strain of the single fibres was reduced.

At the present time we have no direct explanation of this phenomenon. It may be speculated that the higher charge and higher surface softness may permit a greater mixing of the external surface layers of adjacent fibres creating a large contact zone in the z-direction of the fibres. During loading of the paper this in turn means that a larger part of the fibre surface will be affected by the stress in the fibre/fibre joint resulting in a larger overall strain of the paper. This hypothesis is being tested in current investigations.

It should also be added that the chemical treatment of the surface did not affect the density of the sheet and that this straightforward comparison therefore can be made between the sheets.

Conclusion

The joint strengths of single fibre crosses of regenerated cellulose fibres were measured in order to link the properties of the joints formed during web consolidation to the strength properties of the paper. It was found that surface charges of fibres are most important for the development of both joint strength and paper strength. It is suggested that this may be due to surface softening of the fibres, increasing the molecular contact area and promoting a better molecular mixing of the external material on adjacent fibres. Such a mixing of polymers on the fibre surface also seems to increase the critical strain of the network.

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Literature

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Rapid Kotten Blattbildner (delivered by PTE AB) standard procedure ISO 5269-2.

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