Development and evaluation of an automated streaming potential measurement device

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Abstract

The streaming potential and other colloidal properties of aqueous suspensions of bleached kraft fibers were evaluated by a new laboratory instrument, the Streaming Potential Jar (SPJ). This device provides precise streaming potential data under moderately high electrical conductivity levels of 0.5 to 10 mS/cm. Features of the SPJ include automated operation, rapid acquisition and processing of data, continuous stirring, and applied pressures up to 276 kPa. The SPJ also provides data related to drainage rates and the turbidity of the filtrate. Test results showed a high degree of linearity of the streaming potential signals with applied pressure and little dependence of the results on the solids levels of the fiber slurries. These results, which are consistent with the Helmholtz-Smoluchowski equation, tend to justify the level of applied pressure used in this work. Changes in streaming potential with increasing pH were consistent with expected dissociation of surface-bound carboxyl groups on the fibers. The absolute magnitudes of the streaming potential values of bleached kraft pulps were strongly affected by increasing concentrations of Na₂SO₄. However, a high repeatability of measurements was obtained throughout the range of conductivities considered; relative standard deviations of streaming potentials were consistently below 3%. Titrations with
poly-(diallyldimethylammonium chloride) yielded curves that had shapes similar to those of parallel tests by micro-electrophoresis; however, the amount of titrant needed to reach the endpoints was about three times higher in the case of the streaming potential tests. The disagreement between the endpoints determined by the two types of test is attributed to a diffusion process of the titrant into the porous fibers.

*Keywords:* Streaming potential, zeta potential, cellulose fibers, titration, paper

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1. Introduction

The electrokinetic condition of paper furnish, or more popularly, the zeta potential, has been postulated in recent years to be an important determinant in the papermaking process. Electrical properties of fiber surfaces have been reported to be controlling factors in the retention of pigments, dyes, internal sizing agents, and wet strength resins, in drainage, and in flocculation [1-8]. Various experimental techniques have been developed, based on four types of electrokinetic phenomena, electrophoretic mobility [6-9], electro-osmosis [10], streaming potential [11-12], and sedimentation potential [13]. The common feature is that all these methods involve relative motion between charged surfaces and a fluid medium, usually water [5,13]. Related methods include the electrophoretic mass transport analyzer [14-15], electrokinetic sonic amplitude [16], and polyelectrolyte titration with a colorimetric [17-18] or streaming current [19-20] endpoint.

Compared to various other electrokinetic tests that have been used by papermakers, fiber-pad streaming potential tests offer some potential advantages. For one thing, the tests can be carried out with undiluted fiber slurry samples taken directly from a papermaking process [21-29]. Also, it is a potential advantage that the streaming potential signals originate at the surfaces of solids in the sample itself, not at a secondary surface as in the case of streaming current tests [19]. Because the measurements don’t involve light, there is no concern about focusing or the contamination of windows.

Briefly stated, streaming potential tests involve forming a pad or plug, pumping aqueous solution (e.g. water from the process) through the pad or plug. The flow causes movement of counter-ions adjacent to the charged surfaces. The flow of ions yields an
electrical potential that can be measured between electrodes on either side of the pad or plug. When reporting streaming potential data it is customary to also report the values of applied pressure and the conductivity of the solution.

In an effort to relate their results to a more fundamental quantity, many authors have reported use of the Helmholtz-Smoluchowski equation [30-31] to convert streaming potential data into zeta potentials [21-29]. This approach is sometimes justified by correlations between zeta potentials calculated from streaming potential data vs. results of micro-electrophoretic mobility methods [12,32]. At its best, a fiber-pad streaming potential approach can provide an easy-to-use, simple, and practical way to monitor and control the paper machine wet-end [22-29]. Several computer-based online measurement devices have been investigated [26-28,33-35].

Some authors have objected that such use of the Helmholtz-Smoluchowski equation is unjustified [36-37]. The equation was derived for the case of cylindrical capillaries, a physical situation that is grossly different from the case of fiber pads. Not only is the flow pattern much more complex, but it is subject to change as the structure and density of a fiber pad respond to the applied pressure. If the flow is controlled by application of vacuum, then there is a further possibility that bubbles will form on the electrode probes and affect the signal [37]. Questions have been raised also about the influence of surface conductance and its effect on zeta potential determinations [38]. The effects of surface conductance on streaming potential data might be expected to change unpredictably with changes in fiber pad structure or density. Assuming that the density of the pad increases as a result of the pressure application, one also can expect a corresponding decrease in the average or characteristic pore size within the pad structure.
It has shown that such effects can affect streaming potential measurements due to the differing degree of overlap of ionic double layers on the facing surfaces [39]. As a further complication, plots of measured streaming potential versus applied pressure usually fail to pass through the origin [40]. This tendency has been attributed to an asymmetry potential or “rest potential” between the sensing electrodes. To minimize the errors associated with asymmetry potentials it has been recommended to calculate zeta potentials from the slopes of a linear regression between streaming potential and differential pressure [41]. In light of these considerations it is perhaps not surprising that results from streaming potential tests sometimes correlate poorly with micro-electrophoresis, regardless of how things are calculated [9,42]. A classic study by Bull [43] showed that zeta potentials calculated from streaming potential data agreed quantitatively with the results of microelectrophoresis only when (a) both tests were conducted in the same capillary cell, and (b) the surfaces of the suspended particles and the capillary walls were both well covered with a uniform layer of protein, eliminating the likelihood of chemical differences between the surfaces.

The purpose of the present work was to develop and evaluate a new laboratory approach to the monitoring and control of electrokinetic properties of paper furnish. The result of this work is called the Streaming Potential Jar, or SPJ. The working hypothesis is that an especially stable and reliable measure of the colloidal charge state of aqueous slurries at increased electrical conductivity can be achieved with a system comprising the following elements:
1. A system based on streaming potential measurements of the type in which events at the surface of pulp fibers in the aqueous sample themselves give rise to the detected signals;

2. Operation under pressure, rather than vacuum, to avoid generation of bubbles that tend to interfere with electrokinetic measurements;

3. Operation under conditions of pressure that amplify the original electrical signal so that the device can be used at higher electrical conductivity levels where such signals tends to be weak;

4. Automated subtraction of the potential measured at zero pressure in order to minimize effects of asymmetry between the electrodes or their environments;

5. Reliance on titration endpoints, rather than non-zero streaming potential values to minimize problems associated with variable fiber pad structure and density, and chemical interferences;

6. Semi-continuous mixing of a titrant with the slurry sample, together with selection of titrant properties, to favor a uniform distribution of titrant onto the surfaces of the suspended solids.

Additional goals included automated measurement of a drainage rate, filtrate turbidity, and conductivity, all within the context of a user-friendly computer system for operations and data acquisition.

The microelectrophoresis method was used for reference data. Microelectrophoresis has a more extensive and proven theoretical basis, compared to streaming potential tests [13,36-37]. Micro-electrophoresis also has been used by previous authors for
determination of titration endpoints [6,44-45]. Despite its use here as a reference method, microelectrophoresis is not necessarily an ideal method for charge measurement of papermaking furnish. Factors tending to make microelectrophoresis less suitable for routine use in papermaking operations include its restrictions of particle size, a need for high dilution, and a high requirements for user skill and maintenance time.

2. Experimental

2.1 Assembly and procedures for automated streaming potential measurement

Figure 1 shows a schematic diagram of electronic components of the experimental system. The four main parts of the SPJ system are: (1) a computer equipped with a plug-in data acquisition (DAQ) board, a general purpose interface bus (GPIB), and an AT serial card; (2) such commercial devices as a digital millivolt meter, a conductivity meter, a turbidity meter, and a pH meter, each with a probe or transducer to connect it to the wetted parts of the equipment; (3) the wet-end process itself, i.e. the streaming potential jar; and (4) signal conditioning. The system was controlled by LabVIEW (Laboratory Virtual Instrument Engineering Workbench). Like C or Basic, this is a general-purpose programming system with extensive libraries of functions. Measurements included streaming potential, filtrate mass (for drainage), turbidity (related to retention), conductivity, and pH. Data were collected and then output to the computer display.

As shown in Fig. 2, 500 ml of the fiber slurry to be evaluated were placed in an 800 ml cylindrical jar. The contents were stirred continuously with a magnetic stirbar
during each test. A 70 mesh stainless steel screen, having a circular opening with a
diameter of 40 mm was attached to an inverted Büchner funnel within the jar. The jar
was sealed except for the screen opening (connected through the lid), a pressure relief
valve, and the air pressure control system. Experiments were set up so as to allow
addition of titrants and other additives during times when the jar interior was at ambient
pressure.

Pressure and vacuum sequences were actuated by the DAQ system, using the
valve arrangement illustrated in Fig. 2. A “high” level of pressure P1 was achieved with
a conventional air pressure regulator. Rapid adjustment to a steady “low” pressure P2
was achieved with the combination of pressure regulator, air bleed, and bellows. The
continued application of low pressure could be used when needed to maintain the
integrity of the fiber pad during a critical part of the experiment. A value of P2=0 was
selected for the work reported here. The function of vacuum system shown in Fig. 2 is to
return filtrate to the test vessel for the next measurement.

For a measurement cycle, a high pressure P1 was applied at first, then a low
pressure P2, and then the vacuum valve was opened. The duration of pressure and
vacuum applications were pre-set up to suit the drainage properties of various pulp
samples. Changes in the relative electrical potential on the two sides of the screen were
detected with 1.58 mm (1/16 “) diameter silver alloy wires (45% Ag, 30% Cu, 25% Zn).
Voltage information was obtained with a GDM-8055 digital multi-meter, which had an
electrical impedance rating of 1 GΩ over the range of use. In comparison, radio-
frequency conductance values of the sample cell, with fibers included, were in the range
of 50 to 2000 Ω over the course of the work. Output from the digital meter was transferred to data acquisition system and computer display.

As shown at the right of Fig. 2, the amount of filtrate flowing through the mat was evaluated with an analytical balance. The collection system was set up so that filtrate properties such as turbidity, conductivity, and pH could be measured as soon as a relatively small amount of filtrate had passed through the fiber pad collected on the screen. These data, together with the electrical signals for evaluation of streaming potential, were evaluated with the data acquisition system. The Helmholtz-Smoluchowski equation for zeta potential [30-31] was incorporated into the instrument’s software system, for optional use in future research. Electrical conductivity, pH, and turbidity tests were carried out during the period of time form T1 to T2 when low pressure P2 was applied.

2.2 Materials

Pulps used in the study were southern U.S. bleached hardwood kraft pulp (HKP) and bleached softwood kraft pulp (SKP). The pulp was refined in deionized water at 1.56% consistency using a Valley type beater to a freeness value 270 ml in the case of HKP and 260 ml in the case of SKP. Each pulp was disintegrated in deionized water at 0.3% consistency just before the start of an experiment.

For studies on the effects of salts and solution conductivity, sodium sulfate or sodium chloride solutions were added.

Poly-(diallyldimethylammonium chloride) with an average molecular mass of 250,000 Daltons (poly-DADMAC, Aldrich Chemical Co., U.S.A) was used as the titrant.
2.3 Test procedures

Streaming potential tests were carried out with 500 ml samples of 0.3% solids fiber slurry. The applied pressure was 207 kPa (30 psi.) except where indicated otherwise. The durations of high pressure P1, low pressure P2, and vacuum applied periods were set at 30, 15, and 10 seconds, respectively. The rest time between successive measurements was set at various values, depending on the type of experiment. Data were collected for 10 seconds and averaged before and after the pressure was changed from its high to zero setting. The streaming potential value was taken as the millivolt reading at 207 kPa minus the reading at zero applied pressure. In the case of titration with poly-DADMAC, the polymer was added from the pressure relief valve port during the rest time, and the streaming potential before and after polymer addition was measured and recorded respectively. The pH of the furnish for measurements was in a range of 5.5 to 5.8, except where indicated otherwise.

Zeta potential data were obtained with a Charge Analyzer 2 microelectrophoresis meter from SKS Associates. Tests were carried out with fresh filtrate obtained during the course of the corresponding SPJ tests.

3. Results

3.1 Output and reproducibility of signals
Figure 3 illustrates typical data from measurements of streaming potential and drainage. As shown, when high pressure P1 was applied, the potential signal E1 was obtained and recorded, and the balance shown in Fig. 2 started to record the collected filtrate mass. When the pressure was lowered to P2 (set to zero), the reference potential signal E2 was obtained, and the filtrate mass remained at its maximum value.

Drainage was defined as the difference of filtrate mass at T1 minus the corresponding value at T0, divided by the time from T0 to T1 (g/min). As shown in Fig. 3, the drainage rate decreased during each period of steady application of pressure. This is consistent with the build-up and densification of fiber pads. It is to be understood that drainage measurements are affected by the pressure applied and the filtering area of the screen, so the results reported here are unlikely to agree quantitatively with data from other methods. However, the simultaneous measurement of drainage, together with the other data, may be useful for judging the relative effects of chemical treatments.

Figure 4 illustrates the reproducibility of the streaming potential signals. These data were obtained from six cycles of measurements with the same HKP fiber sample in a 0.01 M sodium sulfate solution. As shown, successive values of E1 and E2 gave highly repeatable results. Table 1 shows an analysis of these data. Within an individual test the standard deviation (SD) was no larger than 0.02 mV for the potential signal at high pressure (E1) and 0.35 mV at low pressure (E2), respectively. However, when comparing results of successive tests, the standard deviation of streaming potential was 0.10 mV and the average was –6.72 mV. The calculated relative deviation of streaming potential in the case considered was 1.5%, a typical value for tests reported in this article.
3.2 Effect of varying the applied pressure

Figure 5 shows an approximately linear relationship between streaming potential values and the applied pressure $P_1 - P_2$. The two sets of results shown correspond to experiments in 0.1N NaCl or Na$_2$SO$_4$ solutions, respectively. The pressure differential was adjusted within the range 65 to 275 kPa.

Although the Helmholtz-Smoluchowski equation strictly applies to the case of simple capillaries [13,30,31], it is reasonable to expect that the same physical phenomena also govern streaming potential observations in packed beds or pads of fibers. The excellent linear regression fits are consistent with this supposition. However, the fact that the regression lines failed to pass through the origin suggests that the analogy between cylindrical capillaries and the present experiments is only approximate. As noted by Jayaweera and Hettiarachchi [41], a non-zero intercept is likely to result from asymmetry between the pair of electrode surfaces. In the present work one such asymmetry may result from the fact that only one of the metal probes is in contact with fibers. A further reason to expect some deviations from ideality is the fact that cellulosic fibers are compressing, causing a change in effective sizes of pores as a mat of fibers is subjected to varying levels of pressure. As noted by Ricq et al. [39] deviations from ideality increase with decreasing pore size due to increased overlap of ionic double layers on the facing surfaces and an increasing relative importance of surface conductivity.

The practical implication of the relatively linear results in Fig. 5 is that it supports the use of data obtained at moderately high values of applied pressure, e.g. 207 kPa used in most of the work reported here. As will be shown, streaming potential values tend to
decrease with increasing electrical conductivity of solutions, making the measurements more difficult. Fortunately, the random deviations of streaming potential signals also decreased with increasing electrical conductivity. Since the streaming potential signal is proportional to the pressure applied, a higher applied pressure tends to amplify the signal. It follows that the SPJ method may be advantageous for papermaking processes with relatively “closed” white water systems with correspondingly high electrical conductivities [46].

3.3 Effect of varying the slurry solids and fiber type

A change in pulp consistency can be expected to have some effect on the density of fiber pads formed during a streaming potential test. Consequently, one might expect pulp consistency to affect the results of fiber-pad streaming potential tests [21,39,47]. To evaluate the suitability of the SPJ in various concentration of fiber slurry, both bleached hardwood kraft pulp (HKP) and softwood kraft pulp (SKP) with consistencies from 0.1% to 1.0% were compared. The effect of pulp consistency on streaming potential is shown in Fig. 6.

It is interesting that the streaming potential of HKP was almost unchanged, while that of SKP was changed by less than 10% with changing pulp consistency in the given range. These results suggest that streaming potential was only weakly dependent on the structure of the fiber pads under the conditions of testing. Based on the results of Chang and Robinson [21] the lower absolute values of streaming potential measured at the lowest consistency conditions could be attributed to a lower density of the resulting fiber
pads. In summary, the SPJ appears to be well suited for typical headbox pulp consistencies or more dilute samples.

3.4 Effects of cationic polymer at various pH levels

Figure 7 shows the changes in streaming potentials and titration endpoints as a function of pH. From Fig. 7 it is clear that the streaming potential titration endpoints changed greatly when pH increased from 3 to 4.6 but changed only slightly with increasing pH above 4.6. This phenomenon is only in rough agreement with the dissociation of carboxyl groups of hemicellulose in the fiber [48]. At pH values lower than 3, most of the carboxyl groups take on a proton and the charge on the fibers and fines becomes essentially zero. When the pH reaches 5, more than 65% of these carboxyl groups will become dissociated. Almost all the carboxyl groups will be ionized and contribute to the charge of fibers when pH is 7 or higher. Further work may be required to find out why the apparent charge of the fiber surfaces did not increase significantly in the intermediate range of pH. One possibility is that the fiber surfaces were already sufficiently anionic at pH 4.6 to be able to interact with all of the cationic titrant capable of fitting into the available space.

3.5 Effects of varying the electrical conductivity

Figure 8 compares streaming potential data with zeta potentials obtained by the reference method, micro-electrophoresis. It is worth noting that the values obtained by
the two methods are expected to have a linear relationship only when certain other
variables are held constant [13]. One such variable is conductivity. As shown in Fig. 8,
the streaming potential values obtained with the SPJ were affected to a greater relative
extent by increasing electrical conductivity, compared to the zeta potential data. This
trend is again consistent with common practices of evaluating streaming potential data
[22-29,32-34]. The standard deviation among replicate determinations of migration time
of individual particles was typically about 0.5 seconds, which corresponds to relative
standard deviations of streaming potential in the range 0.5 to 20%. This source of
deviation of the micro-electrophoresis data becomes more important at high absolute
values of zeta potential and low conductivity, whereas convection currents are expected
to affect measurements at the high end of the conductivity range [4].

Figures 9 and 10 show results of titrations of bleached hardwood kraft pulp
measured with streaming potential and microelectrophoresis methods in different
concentrations of sodium sulfate solution. First let us consider a relatively low salt
concentration of 0.001N Na₂SO₄. As shown in Fig. 9, approximately three times as much
titrant was required to achieve a zero signal in the case of the streaming potential tests,
compared to parallel tests with micro-electrophoresis. This type of result also has been
reported by Penniman [32], who attributed it to more rapid mutual diffusion of titrant to
the fine materials, compared to the case of fibers having slower Brownian motion [see
49]. However, in the present case the pulp samples were well washed and relatively free
of fine colloidal materials. Therefore, the rate of adsorption of polyelectrolytes is
expected to be dominated by (a) flow phenomena, and (b) diffusion rates of just the
titrant molecules [49]. By comparison to those rates it is assumed that Brownian
diffusion of fibers and relatively large fiber fines can be neglected. It follows that transport of cationic titrant will occur at a nearly equal rate toward all of the solid surfaces, and the results in Fig. 9 must have some other explanation.

Another way to account for the greater amount of cationic titrant to reach a zero value of streaming potential, compared to the microelectrophoresis test, involves the known surface areas [50] and porous nature of kraft fibers and fines [51-52]. Fiber fines have larger specific surface area than fibers [50]. Assuming a uniform rate of diffusion of titrant to all of the surfaces, more of the positively charged materials will be adsorbed onto the fines, per unit mass, when a titrant is added to a mixture of fines and fibers. The expected result is that the fines will most likely become saturated with titrant and neutralized much earlier than the fibers, after polymer addition. By contrast, the fiber surfaces offer a deeper, porous sink into which the titrant molecules may continue to diffuse [51-53]. Since the streaming potential method measures the whole pulp fibers, but the microelectrophoresis method measures only the fines that passed through the screen, the results in Fig. 9 are consistent with this explanation.

As shown in Fig. 10, a ten-fold increase in salt concentration resulted in titration curves with a different characteristic shape. Results of both types of electrokinetic tests showed “plateau” features, as described earlier by Penniman [32] and Onabe [54]. In other words, the measured electrical potential values changed rapidly with the first small aliquots of titrant, but the output became much less sensitive to titrant addition in an intermediate range. Beyond the plateau, the electrokinetic potential values again responded to titrant addition more rapidly and passed through zero. Tentatively, these
results are attributed to a process of penetration of titrant into pores of the cellulosic materials, after the sites of highest anionic charge on the surface have been satisfied.

Comparing Figs. 9 and 10, the titration endpoint of hardwood pulp with poly-DADMAC were significantly affected by the concentration of sodium sulfate. To neutralize the charge of the whole pulp and to reach at endpoint, the amount of poly-DADMAC titrant required increased with increasing salt concentration. The dosage of poly-DADMAC required for neutralization was 30, 70, and 96 µeq/g pulp in 0.001, 0.005, and 0.01 M Na₂SO₄, respectively. This greater demand for titrant at the higher salt concentration is consistent with the expected lower radius of gyration of the polyelectrolytes, making them able to penetrate into a much higher fraction of the available pores [51-52]. Alternatively, the effect also is consistent with an expected three-dimensional conformation of adsorbed polyelectrolytes at intermediate ranges of electrical conductivity [55-57]. The present results do not yet provide a way to discriminate between these two explanations.

It is interesting to compare the amounts of adsorbed polyDADMAC in the present work with previous estimates of cellulose fiber charge. For example, Öhman et al. [58] estimated a charge density of about 4 µeq/g of charge at the surface of bleached softwood kraft fibers, based on poly-DADMAC adsorption tests in distilled water. This relatively low number, compared to the present work, is consistent with a highly extended conformation at very low ionic strength and a tendency to adsorb with 1:1 stoichiometry on the outer surface of fibers under those conditions [59]. The higher numbers obtained in the present case are also consistent with (a) a lower mass of poly-DADMAC, (b) a somewhat higher carboxyl content in bleached hardwood kraft pulp, compared to
softwood, and (c) the fact that the fibers were refined [60]. By contrast Herrington et al. [61] measured between 40 and 100 \( \mu \text{eq/g} \) of carboxyl groups associated with bleached hardwood kraft samples. These relatively higher numbers are consistent with the potentiometric titration procedures used and the ability of small ions to penetrate into pores that are not easily accessible by polyelectrolytes.

4. Conclusions

A pressurized and automated streaming potential measurement device was developed and tested for laboratory research, and the device showed good reproducibility for electrokinetic measurements. Within a range of solution conductivities between 0.5 and 10 mS/cm, the relative standard deviation among repeated measurements with the SPJ was consistently below 3%.

The magnitude of streaming potential values varied linearly with applied pressure, and it was a weak function of the solids content of the fiber slurries. These facts are consistent with some practical uses of the Helmholtz-Smoluchowski equation. Further work is needed to determine the extent to which values calculated from the Helmholtz-Smoluchowski equation are proportional to zeta potentials determined by microelectrophoresis over a range of conditions, including changes in electrical conductivity. Though the Helmholtz-Smoluchowski equation was derived for the case of cylindrical capillaries, the same equations have become widely used in practical fiber-pad streaming potential devices. The results also help to justify the use of the SPJ within the range of applied pressures used in this work.
Streaming potential results responded in qualitatively reasonable ways to changes in pH, electrical conductivity, and the addition of a cationic titrant. The fact that the apparent surface charge increased greatly in the pH range between 3 and 4.6, but only slightly at higher pH values suggests that surface charge density was not the only factor governing adsorption of the oppositely charged polymer. Rather, the degree of dissociation of carboxyl groups at the fiber surfaces probably was sufficient at pH 4.6 and above to provide anchoring sites for the maximum amount of poly-DADMAC able to fit within the available space. This phenomenon deserves further study.

The titration endpoints determined by the SPJ streaming potential method differed by a factor of three or four, compared to endpoints determined by micro-electrophoresis of filtrate obtained during the same SPJ tests. These results are in qualitative agreement with earlier studies [9,32]. The phenomenon is attributed to a much higher surface area per unit mass of fine material, allowing the fines to more quickly become saturated with titrant that is diffusing at a given rate from the bulk solution to all of the surfaces. It is proposed that the longer fibers, which mainly are sensed by the fiber-pad streaming potential tests, maintain a net negative electrokinetic charge at higher titrant addition levels to mixed furnish because of continual diffusion of the polyelectrolytes into the fine pores in the cell walls.

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References


TABLES

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* Each individual measurement involved averaging of signals over ten seconds; this averaging also made it possible to obtain standard deviation data for each individual E1 and E2 data point.
FIGURE CAPTIONS

Fig. 1. Illustration of electronic components of the Streaming Potential Jar (SPJ) and their arrangement.

Fig. 2. Illustration of SPJ sample jar, pressure control system, and filtrate evaluation system.

Fig. 3. Filtrate mass (relative) and electrical potential data collected during a single cycle of high pressure (P1), low pressure (P2) and low vacuum.

Fig. 4. Six replicate tests of streaming potential of bleached hardwood kraft pulp (BHKP).

Fig. 5. Streaming potential (E2 – E1) values versus applied pressure difference for BHKP suspended in 0.01 M salt solutions.

Fig. 6. Influence of pulp slurry percent solids on the streaming potential of bleached hardwood and softwood pulps.

Fig. 7. Effect of pH on the titration of BHKP slurries with Poly-DADMAC at a conductivity of 1050 µS/cm.

Fig. 8. Comparison of streaming potential data with results of micro-electrophoresis (ME) for BHKP in the presence of different amounts of Na2SO4.

Fig. 9. Comparison of poly-DADMAC titration data by ME vs. the SPJ test in the presence of 0.001 M Na2SO4.

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