Invited Paper

Capillary Electrophoresis Separation Techniques and Mechanisms in Dilute Polymer Matrices

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Capillary electrophoresis in ultradilute derivatized solutions is used to separate nucleic acid fragment mixtures. Pulsed field techniques are used to improve resolution. Protocols in which pulse parameters are varied during the course of the separation are used to increase the range of fragment sizes separated. Fluorescence microscopy is used to study the separation mechanism in ultradilute solution.

INTRODUCTION

Interest in capillary electrophoresis has grown dramatically over the past few years. Sweeping advances have been made in many different areas of the technique such as instrumental and method development research including improved detection schemes, investigation into separation mechanisms, and use of capillary arrays to decrease analysis time. Progress on the experimental side of capillary electrophoresis is winning ever-widening roles in molecular biology, biochemistry and other fields.

Recently, our group has been exploring techniques for separations of long chain nucleic acid fragments by capillary electrophoresis in linear polymer matrices. To this end, we are investigating the use of ultradilute matrices and electric field modulation to increase fragment mobilities while improving resolution. Additionally, we are performing direct DNA imaging experiments, which allow the observation of DNA behavior under different separation conditions. Finally, intracapillary Raman thermometry has been used to measure temperature distributions of Joule heating caused inside a capillary during electrophoresis. The temperature studies have guided investigations into the effects of heating on fragment mobilities and resolution during electrophoresis, and have led to instrumentation for efficient dissipation of this energy.

Capillary Electrophoresis in Dilute Polymer Matrices

Agarose slab gel electrophoresis has been used for nucleic acid separations for many years. The cross-linked matrix is employed for its sieving properties, which are essential for nucleic acid and protein separations. The limiting time factor for slab gel electrophoresis is the very low electric field that can be applied. Joule heating that results from high electric fields will lead to sample degradation, gel failures, and band broadening. Capillary gel electrophoresis (CGE) was developed to take advantage of the superior heat dissipation characteristics of narrow bore capillaries and thus allow the use of high fields, which leads to faster and more efficient separations.

The pioneering capillary gel electrophoresis by Heiger et al. and our initial efforts in pulsed field CGE were performed in cross-linked polyacrylamide gel matrices. The use of cross-linked polyacrylamide followed naturally from the success of agarose slab gel matrices. The technique worked very well for protein and short chain nucleic acid separations. These capillary separations had high resolution and were completed in a fraction of the time of the corresponding slab gel separations.

Drawbacks of the cross-linked matrices are that they must be prepared inside the capillary and they are not very reproducible. In addition, during the course of a separation the stresses placed on the gel matrix cause damage in the matrix, reducing the lifetime of the capillary. Linear polymers were found to be equally useful separation matrices for nucleic acids and are much easier to use than the cross-linked polymers. Water soluble derivatized celluloses are commercially available and the sieving buffers are easy to prepare reproducibly. The solution viscosities are lower than cross-linked gels, so the separations are more rapid. We have shown that hydroxyethyl cellulose concentration about 0.05% (w/w) is near optimum for restriction enzyme fragment separations and Barron et al. also found that dilute linear polymers give good separations.

Surprisingly, baseline resolved separations are obtained even if the polymer concentration is reduced well below the entanglement limit. Barron et al. and Kim and Morris independently reported that hydroxyethyl cel-
ulose at a concentration of 0.002% (w/w) in 1X TBE gives good separations of λDNA/HindIll restriction enzyme fragments. Theories based on established polymer solution physics predict that DNA separations should not be possible in these dilute matrices. The theories require interaction between DNA and a fully entangled matrix.\textsuperscript{30-35} For the celluloses employed, the entanglement limit is about 0.1%. Sub-entanglement cellulose solutions have viscosities only slightly greater than aqueous solutions, yielding very rapid separations. For example, in Fig. 1 we show completely resolved λDNA/HindIll restriction enzyme fragments separated in less than 10 minutes in 0.015% methyl cellulose, using field modulation. This time is much shorter than required in a gel-filled capillary (20-40 min.), and a major improvement over the hour or more necessary in the conventional slab gel electrophoresis.

**Multi Frequency Pulsed Field Capillary Electrophoresis**

An instrumental technique which shows promise for rapid and efficient DNA separations is the implementation of field modulation in capillary electrophoresis. Pulsed field techniques have been used in slab gel electrophoresis for ten years. Theoretical studies and computer simulations have predicted that by disrupting the electric field the alignment of the DNA molecule is also interrupted, allowing for size based separations.\textsuperscript{36-42} In addition, the jostling brought about by the field modulation frees any trapped DNA molecules, eliminating the mobility minimum observed for very large nucleic acids. Rampino and Chrambach\textsuperscript{43,44} and Holzwarth et al.\textsuperscript{45-47} have verified these motions by fluorescence microscopy.

Early work by Heiger et al.\textsuperscript{15} demonstrated resolution enhancement by field inversion capillary gel electrophoresis. The effects of modulation depth and frequency were investigated by our group\textsuperscript{16} which introduced sinusoidal modulation. We showed that the use of single frequency, single amplitude sine waves to modulate the field resulted in separation enhancement for one specific size range of DNA fragments. Novotny et al. have recently extended field modulation techniques to the separation of polysaccharides\textsuperscript{48} and charged polystyrene spheres,\textsuperscript{49} illustrating the effects of pulsed fields on samples much smaller than kilo-base-sized nucleic acid fragments.

Based on these findings and building on the work of Noolandi in slab gels,\textsuperscript{41} we have recently developed variable frequency sinusoidal modulation protocols for field inversion CGE.\textsuperscript{11} Because there is generally a wide range of fragment sizes in one separation mixture, a variable frequency protocol provides optimum separation conditions for the entire fragment mixture during one separation. We found that the net mobilities are primarily a function of the overall applied field, almost independent of the modulation frequencies. However, the resolution is directly dependent on the frequency range of the pulse protocol.

We have been developing more sophisticated pulse protocols in which both the field amplitude and the frequency are varied. By starting the modulation after the smaller fragments in a mixture have eluted, we are able to observe the small fragment resolution found under DC conditions, as well as the large fragment resolution which requires modulation.\textsuperscript{10} We vary both the field strengths and modulation frequencies to now achieve rapid separations of fragments ranging from 70-23,000 base pairs in length. For example, Fig. 2 shows the separation achieved by increasing the AC field strength from 100% of DC to 140% during the course of the separation after a short DC only period achieves complete separation of a fragment mixture ranging from 75 to 12216 base pairs. By using of dynamic pulse protocols during an electrophoretic separation, the separated fragment size range can be extended far beyond the present limits of DC electrophoresis.

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**Fig. 1.** Electrophoretic separation of λDNA/HindIll digest in 0.015% (w/w) methyl cellulose in 1X TBE. Capillary dimensions: 75 μm i.d., 365 μm o.d.; 33.0 cm long, with 28.5 cm from inlet to detection window. **Sample:** 12 ng/μl in 1X TBE; electrokinetic injection, 0.7 kV for 4 s. **Detection:** laser induced fluorescence of DNA/ethidium bromide (3 ng/μl) complex. Separation fields: (a) -180 V/cm DC, (b) -180 V/cm DC, 252 V/cm AC with 1 KHz sine wave modulation. Peak legend: 1 = 554, 2 = 2028, 3 = 2322, 4 = 4371, 5 = 6657, 6 = 9419, 7 = 23130 base pairs.
DNA Fluorescence Imaging

We are supplementing our electrophoretic studies with fluorescence microscopic investigations of DNA migration. The technique was introduced by Smith et al. and extended by Rampino and Chrambach for various agarose slab gel electrophoresis conditions. Perkins et al. have recently looked at the DNA motions under field-free conditions by immobilizing a bead to one end of a nucleic acid molecule and dragging it through a matrix with optical tweezers. This work provides insight into the DNA migration characteristics and reorientation mechanics in gels and fully entangled polymers, but not under the conditions in dilute polymer solutions near and below entanglement concentrations.

Our images show that indeed the basics of entanglement theories are valid. The DNA appears to become entangled at one point along its length forming a U- or V-shaped conformation (Fig. 3a). This is in contrast to entanglement at many discrete points along the backbone of the molecule, which has been observed in agarose and postulated by Barron et al. in the dilute matrices. Additionally, this point of entanglement remains almost stationary in matrices whose concentrations are well above the entanglement limit of the polymer. The entanglement point migrates in the direction of the electric field in matrices near or more dilute than the cellulose entanglement concentration (Fig. 3b). As most theories of gel electrophoresis predict, separation occurs under conditions where an irregular DNA conformation can be generated. The surprise is that an entangled matrix is not needed to produce this conformation.

This dynamic entanglement between the DNA and the matrix allows for such techniques as field modulation to be useful for ultradilute polymer solutions in capillary electrophoresis, as we have demonstrated. We are now beginning to use imaging to study migration under pulsed fields. We expect to observe the perturbations in DNA motions caused by field reversal and to establish what field strengths and frequencies would be optimum for a given size range.

Intracapillary Temperature Studies

Our interest in intracapillary temperatures derives from our field modulation work. The imposed modulation increases power dissipation in the capillary, thus increasing temperature and mobility. As we have reported, in field inversion electrophoresis the heat generated by higher electric fields results in faster fragment mobilities. But resolution is reduced in isorheic DC electrophoresis.

Theoretical calculations by Bello and Righetti and Gobie and Ivory predict an average temperature increase inside the capillary as a consequence of the current flow. We have used Raman thermometry to measure intracapillary temperatures and compare them to predictions and existing data.

The images are yeast chromosomal DNA. The migration direction is indicated by the arrows, and the scale bars represent 10 µm. Electrophoretic conditions: (a) 0.5% hydroxyethyl cellulose in 50% sucrose and 0.5X TBE, 25 V/cm DC field. The images are at 16.0 s intervals. (b) 0.16% hydroxyethyl cellulose in 55% sucrose and 0.5X TBE, 25 V/cm DC field. The images are at 8.0 s intervals.
ing theories. We have found that the theories are fundamentally correct, but there are discrepancies between predictions and actual measurements. The discrepancies are caused by the use of unrealistic boundary conditions in the calculations. The most important of these is the neglect of the heat sinks created in fastening the capillary to external supports. The heat sinks cause substantial thermal gradients along the length of the capillary, which affect the average temperature, and more importantly, are a source of band broadening.

Presently there are two practical ways of dissipating the heat generated by the electric fields. One is by active convective cooling (i.e. blowing air over the capillary) and the other is by flowing a liquid around the capillary housing. We prefer to the first method because if it is by far the simpler to implement. We have developed a simple and effective solid state system to remove the heat from the capillary and have been able to handle fields up to 1300 V/cm without an appreciable temperature increase. The capillary is enclosed in a metallic heat sink with a thin ceramic layer in contact with the capillary itself. The large area of the heat sink allows very efficient convective cooling and the ceramic functions as an electrical insulator. This should allow for DNA separations at higher electric fields without the thermal degradation of the sample and matrix during the electrophoresis and without deleterious resolution effects.

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Capillary electrophoresis; Nucleic acid; DNA; Mechanisms; Gel; Sieving buffer.

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