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ELECTROPHORESIS

High-resolution electrophoretic separation and integrated-waveguide excitation of fluorescent DNA molecules in a lab on a chip

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Keywords:	integrated optofluidics, lab on a chip, fluorescence, ultrasensitive



High-resolution electrophoretic separation and integrated-waveguide excitation of fluorescent DNA molecules in a lab on a chip

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List of abbreviations: Laser induced fluorescence (LIF), Microfluidic (MF), lab on a chip (LOC), Capillary electrophoresis (CE), Waveguide (WG), Limit of detection (LOD), epoxy-poly-(dimethylacrylamide) (EPDMA), Hydroxyethyl cellulose (HEC), Photomultiplier tube (PMT)

Keywords: fluorescence, integrated optofluidics, lab-on-a-chip, ultrasensitive

Abstract

By applying integrated waveguide laser excitation to an optofluidic chip, fluorescently labeled DNA molecules of 12 or 17 different sizes are separated by capillary electrophoresis with high operating speed and low sample consumption of ~ 600 picoliters. When detecting the fluorescence signals of migrating DNA molecules with a photomultiplier tube, the limit of detection is as low as 2.1 picomolar. In the diagnostically relevant size range (~150–1000 base-pairs) the molecules are separated with reproducibly high sizing accuracy (> 99%) and the plug broadening follows Poissonian statistics. Variation of the power dependence of migration time on base-pair size – probably with temperature and condition of the sieving gel matrix – indicates that the capillary migration cannot be described by a simple physical law. Integrated waveguide excitation of a 12- μm narrow microfluidic segment provides a spatio-temporal resolution that would, in principle, allow for a 20-fold better accuracy than is currently supported by state-of-the-art electrophoretic separation in microchips, thereby demonstrating the potential of this integrated optical approach to fulfill the resolution demands of future electrophoretic microchips.

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1 Introduction

Microfluidic (MF) lab-on-a-chip (LOC) devices [1, 2] have enabled integrated DNA sequencing [3] and genetic diagnostics [4, 5]. Capillary electrophoresis (CE) separation is a powerful method over a broad range of molecular sizes, thanks to flexible MF protocols, exploiting a variety of sieving gel matrices. The inherent advantages of MF CE, high-speed operation and low reagent volumes, can be combined with laser-induced fluorescence (LIF) detection [6-8], resulting in optofluidic integration [9, 10] toward compact, sensitive on-chip bio-analysis tools [11] which aim at solving real-life challenges in medicine [12], e.g. identification of genomic deletions or insertions associated with genetic illnesses.

Fluorescence excitation via an integrated optical waveguide (WG) provides advantages over focusing an external laser beam by lenses or through a microscope, such as stability and reproducibility of the optical excitation and avoidance of the requirement to realign the laser beam from time to time. Besides, it is one step toward full sensor integration in the MF LOC. We apply the technique of femtosecond-laser micromachining [13, 14] to post-process optical WGs into commercially mass-produced LOCs on a fast, flexible, chip-by-chip basis [15], thereby exploiting the existing mature MF fabrication infrastructure, while achieving integrated fluorescence excitation. Having pioneered the use of such integrated-WG laser excitation for electrophoretic DNA analysis [16], here we demonstrate excellent DNA-sizing accuracy and low background noise, resulting in a lower limit of detection (LOD) compared to earlier accounts [17]. Furthermore, the spatio-temporal resolution provided by integrated waveguide excitation

can allow for a 20-fold improvement in accuracy if better MF CE separation protocols become available.

2 Optofluidic chip fabrication and characterization

The lay-out of our optofluidic chip is presented in Fig. 1. Chips were fabricated in a two-step procedure. Firstly, the MF channel network and MF reservoirs were patterned photolithographically and wet-etched in fused silica glass and then sealed off by bonding another piece of fused silica glass on top. The chip has dimensions of 55 mm × 5.5 mm × 1 mm and the MF channels have a cross section of ~110 μm width and ~50 μm depth (LioniX BV) [<http://www.lionixbv.nl>].

In a second step, the optical WG was inscribed into the bulk of such a fused silica chip by femtosecond-laser writing at a translation speed of 20 μm/s, using a Ti:Sapphire laser operating at 800 nm wavelength with 150 fs, 4 μJ pulses at a repetition rate of 1 kHz. Employing astigmatic beam shaping, an elliptical cross section of the written WG was obtained, with a major diameter of ~50 μm in the vertical direction, in order to excite the maximum possible volume of the MF channel, while the minor diameter in the horizontal direction is ~12 μm in order to retain a high spatial resolution along the direction of DNA flow and separation. The WG displays a graded refractive index profile and a maximum refractive index increase of 2×10^{-3} . Fiber-to-chip coupling of laser light with coupling losses of < 2 dB was achieved and the WG propagation losses were measured to be in the range of 0.5–0.9 dB/cm at the wavelength of 543 nm.

The excitation WG intersects the CE separation channel orthogonally at a distance of ~3.6 cm from the MF crossing junction at which the separation commences, toward the end of the MF CE separation channel, close to reservoir 4, where the plug separation is highest. The fluorescence signal was detected through a small detection window in order to reduce the background noise. Constant, inherent, mutual alignment of the excitation WG with its detection window has the potential to render our system more compact, faster to operate, and more reproducible compared to conventional LIF approaches [6-8].

3 Results and discussion

3.1 Electrophoretic separation protocol

Prior to the experimental runs the inner walls of the MF channel network were coated with an epoxy-poly-(dimethylacrylamide) (EPDMA)-based polymer in order to suppress the electro-osmotic flow and to minimize adsorption of DNA molecules on the MF channel wall. Subsequently, the channels were filled with a sieving gel matrix consisting of hydroxyethyl-cellulose (HEC) (2% wt./vol.), dissolved in 20 mM MES / 20 mM His buffer (pH 6.2) in order to maximize the resolution of the DNA CE separation. SYBR Green I (Molecular Probes Inc.) was added according to the manufacturer's protocol. The reagents were sterilized, filtered (0.22 µm pore size) and stored at 269 K. The CE sample loading and separation protocol was based on actuation voltages of up to 1.5 kV, delivered by Pt electrodes integrated into the MF reservoirs, with the help of a Labview

(National Instruments, Inc.) program to steer the MF control system (CapiliX BV) [<http://www.capilix.com>]. Application of a high voltage forced the negatively charged DNA molecules to migrate into the CE injection channel from sample reservoir 1 to waste reservoir 2. By switching the voltages at all four reservoirs simultaneously a well-confined plug of DNA molecules – with a volume of ~605 picoliters at the crossing junction of the two MF channels – was injected into the CE separation channel, from the MF crossing junction toward waste reservoir 4, and the DNA molecules contained in the plug volume were separated according to their size.

3.2 Fluorescence excitation and detection

Double-stranded DNA molecules were injected as a single mixture plug into the MF CE separation channel. For the movie of Fig. 2 Hg arc lamp illumination was applied in order to visualize migration of the plugs, and the fluorescence was recorded by a CCD camera. Excitation through the optical WG was provided by the 488-nm line from a continuous-wave Ar-ion laser. For ultrasensitive fluorescence detection, a photon-counting photomultiplier tube (PMT) (H7421-40, Hamamatsu Photonics K.K.) cooled down to 203 K was built onto the output port of an inverted microscope (DM5000, Leica Microsystems GmbH) and aligned to collect the fluorescence signal through the detection window. Combination of excitation/emission band filters (K3, Leica Microsystems GmbH, and XF57, Omega Optical, Inc.) in the fluorescence collection path ensured that only the fluorescence signal emitted by the fluorescently labeled DNA molecules reached

the PMT, while all other sources of background signal, including the scattered laser excitation signals, were rejected.

A commercial DNA ladder consisting of molecules with 17 different sizes (in the range of 50-3000 bp) was injected, separated, and the individual plugs detected. The movie (Fig. 2, online) depicts the flow of the separated DNA plugs across the excitation WG under background Hg arc lamp illumination. The four snapshots in Fig. 2 correspond to the flow of the first well-separated DNA plug across the excitation WG. At this point in time, the remaining well-separated 16 plugs were not yet in the field of view of the CCD camera. Upon laser excitation of such a plug through the WG a sharp fluorescent segment only $\sim 12\text{ }\mu\text{m}$ in width (matching well with the minor diameter of the WG cross-section) can be observed along the WG-MF-channel intersection (Fig. 2). Separation of these 17 DNA molecules was repeated in the same microchip, providing reproducible results, as shown in Fig. 3A-B.

After fitting the fluorescence intensity peaks in Fig. 3A-B with Gaussian profiles (the Gaussian fit of the peaks in Fig. 3B is shown in Fig. 3C) the obtained peak widths were plotted vs. the square root of the DNA molecule sizes, see Fig. 3D. The square-root dependence, confirmed by the linear fit, indicates that the migration of DNA molecules follows Poissonian statistics, the dispersion being governed by the number of collisions a molecule typically undergoes as it traverses through the sieving gel matrix [18]. The Gaussian fit is nevertheless justified, as for high expectation values the Poissonian approaches a Gaussian shape.

The LOD of DNA analysis in the integrated optofluidic chip were explored under integrated-WG laser excitation. Replacing the CCD camera by an ultrasensitive PMT

allowed us to record the fluorescence intensity vs. migration time, as depicted in the electropherograms of Fig. 3A-B. We estimate the LOD by considering that an injected DNA concentration of 10 nM diluted by, firstly, the spatial separation of 17 different molecule sizes and, secondly, plug broadening from 110 μm to at least 248 μm , provided a fluorescence signal which is 125 times the four-fold standard deviation of the detected background noise. Based on this method [19], our conservative estimate of the LOD is ~2.1 pM, which surpasses previously reported LOD values in laboratory setups involving on-chip-integrated fluorescence monitoring [15, 17].

3.3 Base-pair resolution

In an independent experiment carried out in a different microchip, another set of double-stranded DNA molecules with 12 different sizes, in a relatively smaller size range (150-500 bp), was separated and the individual plugs detected. The resulting electropherogram is shown in Fig. 4.

Optimization of the coating and sieving gel matrix led to excellent CE separation of the DNA molecules under integrated-WG laser excitation in both experiments, see Figs. 3A-B and 4. When plotting the temporal occurrence of the fluorescence peaks of the 17 molecules in Fig. 3A-B vs. their *a priori* known base-pair sizes, the two smallest (insufficient interaction with the sieving matrix) as well as the two largest DNA molecules (tendency to fold) exhibit clear deviations, while the peaks of all DNA molecules with sizes of 150-1000 base-pairs, in the diagnostically relevant region, are resolved with a sizing accuracy higher than 99% (Fig. 5A). This high sizing accuracy is

verified when analysing the result of Fig. 4 for the 12 DNA molecules in a smaller size range. The high sizing accuracy (> 99%) is comparable to state-of-the-art performance in microchip CE-based DNA sequencing without integrated-WG laser excitation [20, 21].

Each measured fluorescence peak in the electropherogram is a convolution – along the flow axis – of the Gaussian WG mode-field cross-section Δx_{WG} and the diffusion-induced physical width Δx_{MF} of the MF plug. The resulting spatial plug widths Δx are related to the temporal electropherogram peak widths Δt (provided in Fig. 3D as full width at half maximum) as

$$\Delta x = \frac{L}{t} \Delta t = \sqrt{\Delta x_{WG}^2 + \Delta x_{MF}^2} ,$$

where $L = 3.6$ cm and t (data on the abscissa of Fig. 3A-C) are the migration length and time, respectively. In our experiment, the minimum electropherogram peak width was found to be $\Delta x = 248$ μm (the very first peak in Fig. 3B-C). With $\Delta x_{WG} = 12$ μm , we obtain $\Delta x_{MF} = 247.7$ $\mu\text{m} \gg \Delta x_{WG}$, i.e., the current spatial resolution is limited by the MF plug width, while the obtained integrated-WG mode profile would allow for a 20-fold better resolution. MF technological advances, e.g. by improving the MF channel-wall coatings and sieving gel matrices, will lead to smaller MF plug widths, thereby fully exploiting the monitoring resolution achieved with integrated optical WGs. Clearly, integrated optical instrumentation is well suited to handle the resolution demands of future MF-chip generations.

From the measured migration time t vs. DNA size (Fig. 5A) we derive the electrophoretic mobility μ according to the equation

$$\mu = \frac{L}{E t},$$

where L is the distance between the MF crossing junction and the detection window and E is the electric field strength in the MF CE separation channel. The observed linear decrease (on double logarithmic scale) of the electrophoretic mobility with increasing DNA size (Fig. 5B) matches well with observations in conventional bulk capillary-based DNA gel electrophoresis setups [22].

By plotting the results of Fig. 5A on a double-logarithmic scale, see Fig. 5C, slopes of 0.24 and 0.25 (indicated by the straight lines) are derived for the two consecutive experiments, respectively, investigating the flow of the 17 DNA molecules. These results are significantly different from the slope of 0.31 derived for the flow of the 12 DNA molecules, despite the fact that in all cases a high sizing accuracy was obtained. This variation of power dependence points toward different flow mechanisms in these experiments, potentially depending on ambient temperature and/or condition of the sieving gel matrix, a result that questions whether a general, simple model of electrophoretic flow in micro-capillaries can be established and universally applied.

4 Concluding remarks

Simple and straight-forward integration of optical waveguides into commercial microfluidic chips by femtosecond-laser post-processing enables separation of fluorescently labeled DNA molecules with low limit of detection of 2.1 pM and high

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sizing accuracy of > 99%, leaving plenty of room for further improvement of accuracy via better electrophoretic separation protocols. This fabrication technique also allows for three-dimensional waveguide structures [23], thereby further enhancing the potential of the optofluidic chip technology for monitoring on-chip CE.

For Peer Review

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Figure Captions

Figure 1. Schematic of the optofluidic chip showing reservoirs 1-4, sample injection channel (reservoir 1 → reservoir 2) and CE separation channel (reservoir 3 → reservoir 4), as well as the integrated optical WG and detection window.

Figure 2. Movie recorded with a CCD camera showing transient fluorescence from several molecule plugs formed by CE separation of the DNA ladder as these plugs pass by the point of integrated-WG laser excitation at 488 nm.

Figure 3. CE separation in an optofluidic chip of a DNA ladder consisting of double-stranded molecules with 17 different sizes (50-3000 base-pairs), fluorescence-labeled with an intercalating dye. (A) Electropherogram of fluorescence intensity vs. migration time recorded by a PMT after integrated-WG laser excitation. The *a priori* known base-pair sizes are indicated; (B) repetition of the same experiment in the same microchip; (C) Gaussian fits to the fluorescence peaks of Fig. 3B, The peak widths are indicated; (D) peak widths obtained from Gaussian fits to the fluorescence peaks of Fig. 3A and 3B vs. square root of the base-pair size.

Figure 4. Fluorescence intensity vs. migration time for 12 double-stranded DNA molecules in the size range of 150-500 bp, flown and separated independently in a different microchip.

Figure 5. For the two flow experiments on the same chip with 17 double-stranded DNA molecules in the size range of 50-3000 bp, and the flow experiment with the 12 double-stranded DNA molecules in the size range of 150-500 bp, in a different chip, the plots of (A) Migration time (on a linear scale) vs. DNA size (on a logarithmic scale), indicating a reproducibly high sizing accuracy; (B) Electrophoretic mobility vs. base-pair size; on a double logarithmic scale; (C) Migration time vs. base-pair size on a double logarithmic scale; depicting the different flow mechanisms.

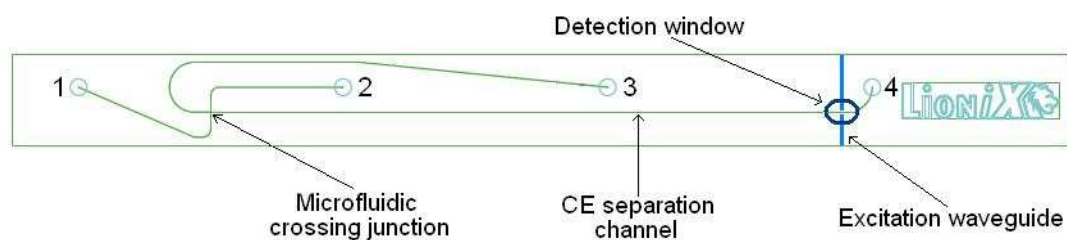
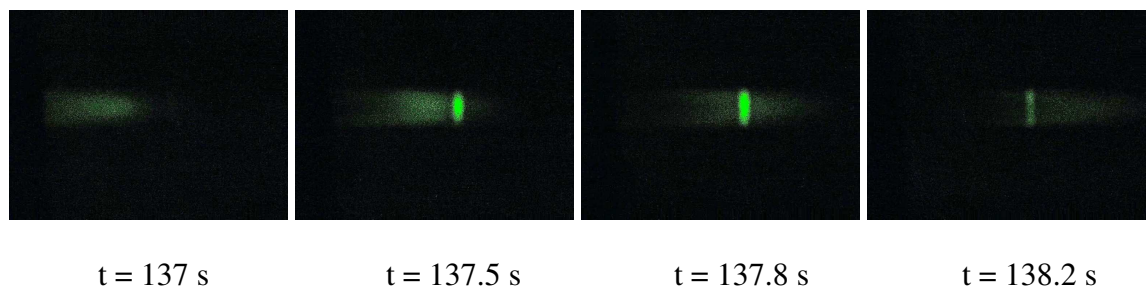
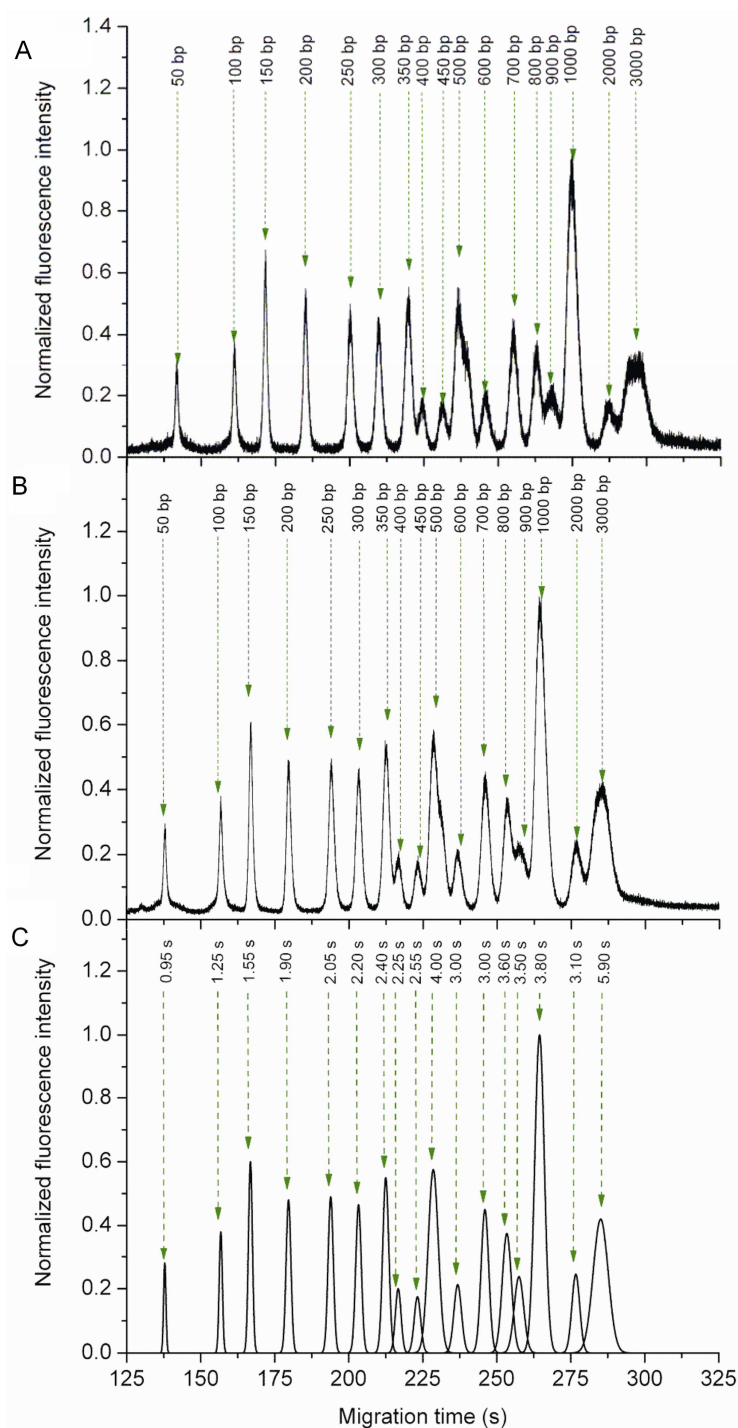
Figure 1

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For Peer Review

Figure 3

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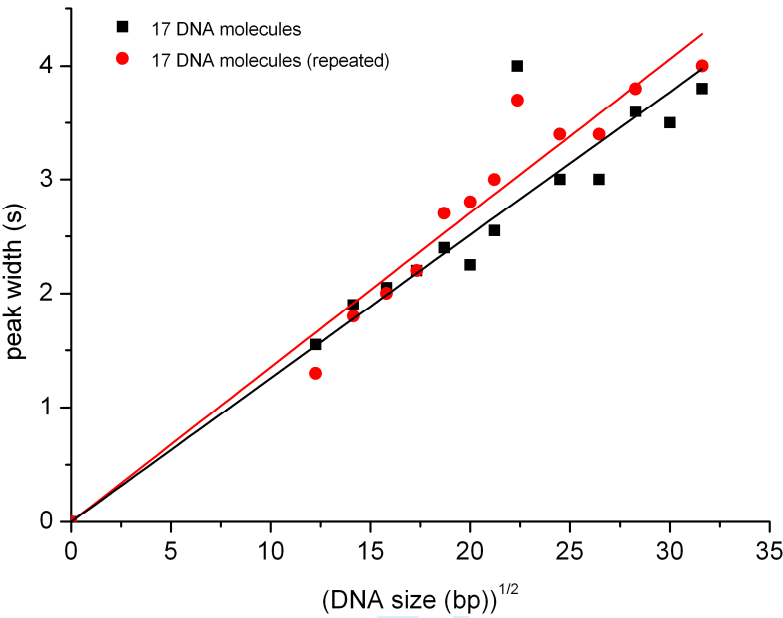


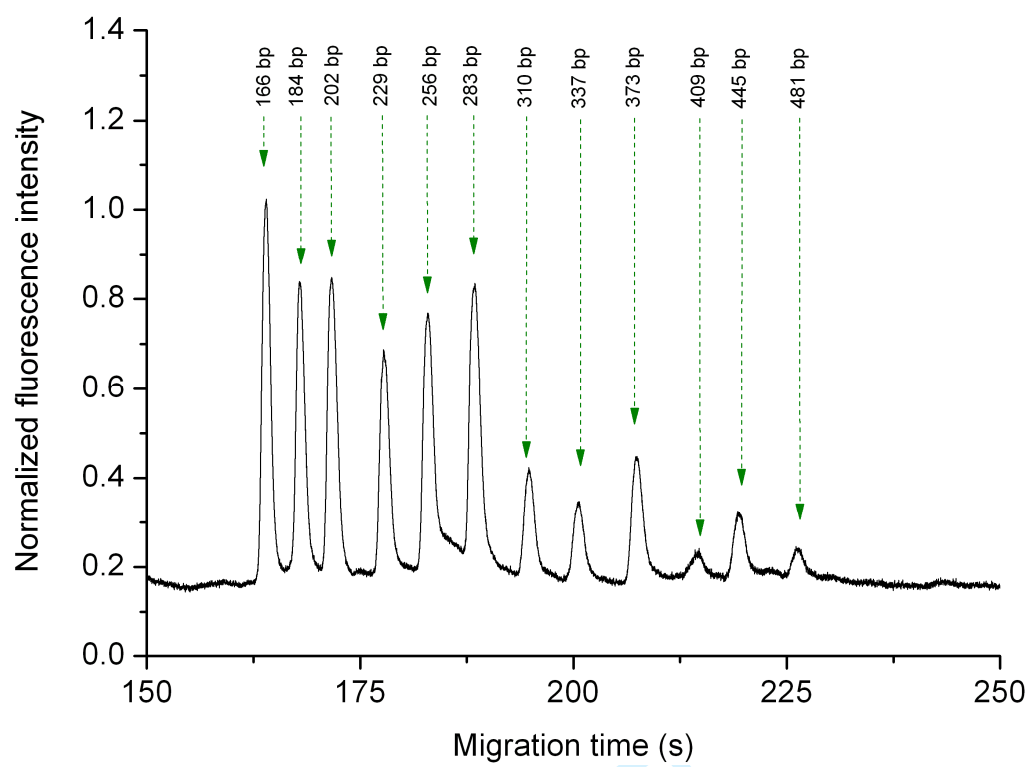
Figure 4

Figure 5

