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## アプリケーションノート

# Plasmid Topology and Digest Analysis Using a GTxResolve™ 250 Å Slalom Chromatography Column

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## **Abstract**

This application note illustrates the characterization of plasmid DNA topological isomers using a GTxResolve 250 Å Slalom Chromatography Column. Slalom chromatography discriminates large nucleic acid species by exploiting the dynamic differences in their rigidity and molecular stretching. Plasmid DNA provides the code for all recombinant expressions as well as a template for mRNA synthesis through *in vitro* transcription (IVT). Its quality is pivotal to ensuring the quality and efficacy of many different therapeutics, including mRNA drug substances. Purified plasmids can exist in supercoiled (S) as well as open-circular (OC) and linear (L) forms. Supercoiled species are desired for effective transfection when expressing recombinant proteins. Meanwhile, linearized plasmids are generally sought when performing IVT reactions to produce mRNA. In either case, impurities need to be kept to minimal levels. Here, the separation of plasmid isoforms in less than 5 minutes is shown by utilizing a slalom chromatography column manufactured from optimally packed high-purity, hydrophilic 2.5 µm particles and column hardware modified with MaxPeak™ High Performance Surfaces (HPS).

This column hardware and the packing material exhibit excellent inertness to produce high-resolution, high-recovery analyses. The identified chromatography conditions maximize the flow retardation effects of slalom chromatography to distinguish three different species of a large plasmid DNA of same size. Depending on the size of the plasmid, it is shown that slalom chromatography can potentially even separate closely related

supercoiled and open circle plasmid species. Thus, slalom chromatography provides an intriguing new alternative to agarose gel electrophoresis or capillary gel electrophoresis to analyze plasmid DNA species in <5 minutes by resolving them. The ability of slalom chromatography to analyze plasmid DNA impurities should enable quicker development of efficacious and more globally accessible cell and gene therapy medicines.

#### **Benefits**

- Excellent separation of plasmid isoforms, including S and OC forms depending on the length of the constructs
- Confirmation of gene insert in recombinant plasmid using restriction enzyme digests
- Excellent analyte recovery with minimal nonspecific adsorption
- High-throughput analyses with gel-like separation capabilities for large (≥ 3kbp) nucleic acids
- Robust and tunable chromatography to maximize the separation conditions

## Introduction

Genetic medicines use DNA and RNA to facilitate tailored medical interventions. Messenger RNA (mRNA) with protein-coding instructions can modify cellular gene expression through protein replacement or the introduction of novel proteins all together. Despite the promise and performance of mRNA drugs, the analytical strategies to maintain quality control are not fully developed or require tedious procedures. <sup>1</sup> The large size, complexity, heterogeneity, and high molecular weight of most mRNA sequences, and lack of robust and highthroughput analytical methods, are slowing down the deployment of mRNA therapeutics. Key steps in the mRNA vaccine production include (i) preparation of a plasmid DNA template of protein coding sequence for IVT, (ii) IVT reactions for mRNA synthesis and mRNA purification, (iii) encapsulation of mRNA into a lipid nanoparticle, and (iv) sterile filtration before use. The quality of the plasmid that provides the template for mRNA synthesis is pivotal for ensuring the integrity and efficacy of the resulting mRNA. The plasmid may exist in the S, OC, and L forms due to nicks (cuts) on any one or both strands. The presence of OC and incorrectly linearized forms of a plasmid affects mRNA integrity as the IVT reaction gets terminated at random nicks, leading to shorter and heterogeneous transcripts. Uniform and appropriate linearization of the IVT plasmid is essential for efficient RNA synthesis and generating correct 5' and 3' ends. Thus, the analysis and quality control of IVT plasmid isoforms has become a critical strategy for minimizing process-related impurities in mRNA drugs. Current techniques for investigating these impurities, such as agarose gel or capillary gel electrophoresis can be tedious or require large quantities of samples. As such, alternative methods are desired.

Slalom chromatography is a promising alternative because it provides separation selectivity for large nucleic acids using pressure versus electrophoretic-driven effects. This chromatographic approach provides high-throughput analyses as a result of its inherently fast flow rates.<sup>3–4</sup> In a slalom separation, the largest nucleic acid components and their topological isomers will experience flow retardation resulting in their separation.

Large nucleic acids will undergo entropic extension when exposed to the shear forces exerted during a run. Due to their rigidity, linear double-stranded DNA (dsDNA) will resist these forces and attempt to revert to its equilibrium configurations. Any single-stranded species that might be present will succumb to these shear forces and rapidly pass through the column. Differences in the conformational states of supercoiled and linear plasmid DNA are also significant enough for them to be separated by slalom chromatography. Interestingly, mobile phase shear forces can be optimized to achieve the desired levels of selectivity in a separation. Flow rate, solvent viscosity, and column temperature all contribute to the optimization of slalom chromatography separation.

In this application note, the reproducible high-resolution separations of plasmid DNA isoform species is shown using GTxResolve 250 Å Slalom Chromatography Column. Large plasmid DNA isoforms do not access the intraparticle pore volume of this packing material with a 250 Å average pore diameter, but they are resolved across time and elution volume due to flow retardation differences. Optimization of column chromatography conditions, including column temperature, enabled the resolution of all three isoforms of an example plasmid DNA ( $\geqslant$  7kbp) in under 5 minutes, making it a very high-throughput analysis method.

## Experimental

1X TAE buffer: 40 mM Tris-acetate, 1 mM EDTA pH 8.3 made from

10X TAE (ThermoFisher Technologies p/n:

15558042)

Column storage: 10% acetonitrile, 90% aqueous 25 mM Sodium

Phosphate pH 7.0 + 100 mM Potassium Chloride

Seal and weak wash: 20% Methanol: 80% Water

Samples: Plasmids: pBR322 (New England Biolabs – N3033L), pCMV-

Cas9-2A-GFP (PP (CAS9GFPP-1EA) and pCMV-Cas9

(CAS9P-1EA) from Millipore Sigma. EcoRI (R0101S), Nhel-HF (R3131S), and XbaI (R0145S) were procured from NEB.

Plasmid digests: EcoRI digest of pBR322:

To 43  $\mu$ L of 18.2 M $\Omega^*$ cm water, 5  $\mu$ L of 10X NEBuffer (r2.1), 1  $\mu$ L of pBR322 (NEB p/n: N3033L) and 1  $\mu$ L of EcoRI (NEB p/n: R0101T) were added, vortexed, and briefly centrifuged to gather all the contents. After incubation at 37 °C for 1 hour, the enzyme was inactivated by incubating the digest at 70 °C for 20 minutes.

Double digest of pCMV-Cas9:

To 34  $\mu$ L of 18.2 M $\Omega^*$ cm water, add 10  $\mu$ L of pCMV-Cas9 plasmid (20 ng/ $\mu$ L) yielding a total of 200 ng of plasmid then add 5  $\mu$ L of 10X rCutSmart $^{\text{TM}}$  Buffer, 0.5  $\mu$ L of Xbal (NEB p/n: R0145S) and 0.5  $\mu$ L of Nhel (NEB p/n: R3131S) were added, vortexed and briefly centrifuged to gather all the contents. After incubation at 37 °C for 1 hour, the enzyme was inactivated by exposing the digest to 75 °C for 20 minutes to inactivate the enzymes.

Agarose gel electrophoresis:

0.6% agarose gel was prepared by using Agarose (A0539-500G) from Sigma Aldrich, LOT# 0000320705, Source SLCR2997. About 0.6 g of agarose was dissolved in 100 mL of 1X TAE buffer by boiling in a microwave, after initial cooling (~1-2 minutes), 10 μL of SYBR® Gold Nucleic Acid gel stain (Invitrogen p/n:S11494) was added, thoroughly mixed and poured into a gel tray with a comb. After solidification at room temperature, 1X TAE buffer was added, the comb was gently pulled to make wells for sample loading. Load 200 ng of sample into individual wells. The gel setup in

a horizontal Bio-Rad electrophoresis system was connected to the PowerPac HV High-Voltage
Power Supply. Electrophoresis was done at 50V for 2 hours and 10 minutes. After electrophoresis, the gel was imaged using the Gel Doc EZ Gel
Documentation System (Bio-Rad) to visualize the bands.

LC system:

ACQUITY™ UPLC™ I-Class Bio (or equivalent) consisting of QSM H-Class (Quaternary Solvent Manager with 50 μL mixer), TUV Detector - 1000 psi MAX Pressure 70 bars Max Pressure 500 nL, Sample Manager FTN (Ver. 2023 03 02), 15 μL Needle size (p/n: 700012820) Assy, Needle, FTN-15 μL, HPS, TXT w/Gd&st), 100 μL Sample syringe size, Injection Valve Flow Through (p/n: 700011791) External Pre-Heater, CH-30A, with Active Preheater (APH) MP35N, 0.004 ID, 18.5' (p/n: 430005558) and post column tubing to TUV: 0.004' ID 22.5' LG MP35N Welded Tube (p/n: 430002856), Assy, Pre-HRT to Column, Reusable MP35N (p/n: 700011809), ACQUITY UPLC 30 cm Column Heater/Cooler (p/n: 176015127)

Column:

Waters<sup>™</sup> GTxResolve 250 Å Slalom Column, MaxPeak Premier Technology, 2.5 μm, 4.6 x 300 mm and dsDNA 23K Ladder (p/n: 176006046)

Mobile phase A:

1X TAE buffer, pH 8.3

Column equilibration:

Column conditioning is done by ramping the flow rate by 0.1 mL increments to reach the flow rate of 1.3 mL/min over 20 minutes and continued for another 40 minutes.

Vials: Max Recovery Sample Vials (Waters p/n:186009186

(vial) 186009186 (caps))

Column temperature: 5 °C, 25 °C, and 40 ⊠°C

Sample temperature:  $5 \boxtimes \pm \boxtimes 3 \boxtimes ^{\circ} C$ 

Sample amount:  $1 \mu L$ 

Flow rate: 0.5 or 0.7 mL/min

Gradient: Isocratic

Sampling rate: 40 Hz

## Results and Discussion

The capability of a GTxResolve 250 Å Slalom Column to produce fast yet reproducible separations of large DNA molecules was demonstrated in a previous application note. In the current studies, applying slalom chromatography to the characterization of plasmid samples was taken into consideration. Plasmid DNA, when purified from cells, exists mostly as a S configuration with varying amounts of OC and L forms as impurities. When transfecting cells, it is desired to have high-purity supercoiled DNA. When performing *in vitro* transcription reactions to produce mRNA, it is important to work with high-purity linearized DNA. It is critical that supercoiled DNA be cleaved at the designated restriction site for linearization to ensure appropriate 5' and 3' termini are obtained for synthesized mRNA drug substance.

Figure 1 shows the ability of slalom chromatography to evaluate the extent of linearization of plasmid DNA in <5 minutes. In this example, *E. coli* plasmid pBR322 (4,361 bp) was examined. It harbors a single cleavage site for *Eco*RI at 4,359 bp; therefore, upon cleavage the supercoiled form will be converted into a linear form. Due to its rigid supercoiled structure, the untreated pBR322 is expected to differ in its stretching ability compared to the linearized version, thus differing in their flow retardation effects. As predicted, the supercoiled plasmid eluted early at approximately 1.142 minutes, while the linearized form was retained longer, leading to their elution at ~1.277 minutes. This pattern was reproduced again when a mixture of supercoiled and linearized forms of pBR322 was subjected to slalom chromatography (Fig 1; bottom panel). Thus, if plasmid preparations

involve linearization, analysts can easily apply slalom chromatography to measure percent linearization.

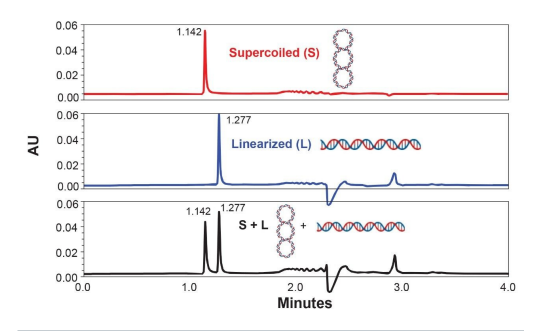


Figure 1. Plasmid linearization efficiency analysis by slalom chromatography. 100 ng of either undigested/supercoiled form of pBR322 (top) or EcoRI - treated (middle) or a mixture of supercoiled and linear forms of pBR322 plasmid (bottom) were subjected to slalom chromatography using 1X TAE mobile phase at 1.25 mL/min flow rate (10,850 psi) and 40 °C to understand their elution behavior. Note that the retention times of the two peaks in the mixture correspond to the supercoiled and linearized forms of the plasmid DNA.

To investigate whether an open circular form of a plasmid can be detected by slalom chromatography, the refrigerated column temperatures and a set of plasmids of varying sizes was examined. Separations were performed at 5 °C column temperatures using an auxiliary column heater/cooler oven. Although pBR322 did not show any significant effects at lower temperature (data not shown), larger size plasmids pCMV-Cas9-GFP (7906 bp) and pCMV-Cas9 (7,043 bp) exhibited multiple features, including pre-peaks before the main peak as well as a late eluting peak (Figure 2). Interestingly, the main peaks (RTs of 2.399 and 2.343 minutes for Cas9-GFP and Cas9, respectively) in both plasmids exhibit a retention time difference of 0.056 minute, reflecting the difference in their size. Further, *Nhel*-treated pCMV-Cas9 exhibited a peak that corresponded to the late eluting peak of the undigested plasmid. The linearized sample of pCMV-Cas9 (Figure 2c) also exhibited additional features, namely small peaks at a retention time window corresponding to supercoiled plasmids. These observations provide further evidence that the extent of plasmid digestion and generation of any potential

artifacts can be monitored by slalom chromatography as long as the plasmid sample is greater than or equal to 7 kbp in size. Smaller (<7 kbp) plasmid samples will require the use of anion exchange chromatography to effectively resolve supercoiled and open circular variants. Simultaneous use of anion exchange and slalom shearing forces is also promising, as recently reported by Finny *et al.*<sup>9</sup>

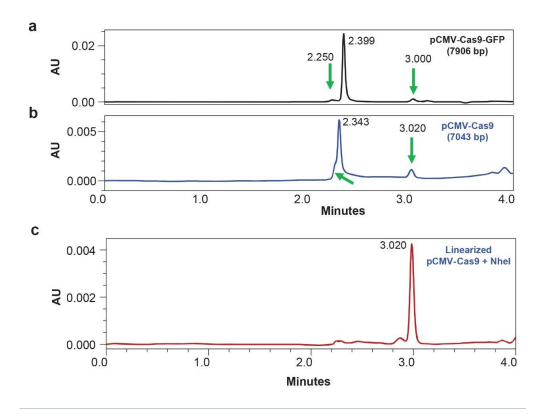


Figure 2. Plasmid topology studies by slalom chromatography. Two different plasmids pCMV-Cas9-GFP (panel a) and pCMV-Cas9 (panel b) from Sigma were analyzed at 5 °C with 1X TAE pH 8.3 buffer as mobile phase under slalom conditions (10,500 psi back pressure). These plasmids are predominantly present in supercoiled form. Open circular and linear forms are expected to be present as low-abundance impurities (top and middle panel chromatograms). Early and late eluting impurity peaks are indicated by arrows. It is hypothesized that these additional peaks can be attributed to open circular and linear impurities, respectively.

Slalom chromatography can also be applied for another type of investigation: confirming the insertion of a gene of interest. Successful cloning of a gene sequence into the designated sites of a plasmid backbone is generally monitored by agarose gel electrophoresis followed by double digestion with two restriction endonucleases. Figure 3 shows that slalom chromatography can be used to monitor the presence of a gene

insert in less than 5 minutes. Double digestion of pCMV-Cas9 was performed by using *Nhe*I and *Xba*I (panel c) and compared against undigested (panel a) as well as linearized (*Nhe*I treated) plasmid (panel b). Successful cleavage of pCMV-Cas9 at *Nhe*I and *Xba*I sites was confirmed to release the 4147 bp Cas9 gene sequence. Although the linear Cas9 insert (4147 bp) and the residual plasmid backbone (~2896 bp) exhibit a relatively small difference in length (1259 bp), they can still be resolved (a difference of 0.077 min) by slalom chromatography indicating its sensitive detection capability to evaluate digestions as compared to agarose gel electrophoresis (Figure 3d). About 200 ng of digest were needed to see clear bands for gene insert and plasmid backbone on an agarose gel. In contrast, 40 ng of digest were sufficient to detect both the peaks, indicating 5X improvement in sensitive detection by slalom chromatography.

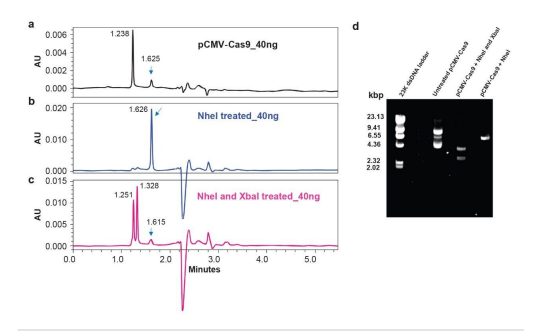


Figure 3. Confirmation of gene insert in a recombinant plasmid by slalom chromatography at 25 °C. Digestion of pCMV-Cas9 by Nhel and Xbal releases Cas 9 insert (panel c) and both insert, and plasmid backbone are resolved by slalom chromatography despite small differences in their lengths. Undigested plasmid (panel a) and linearized (Nhel-treated) plasmid (panel b) are shown for comparison. (d) Agarose gel electrophoresis of the pCMV-Cas9 samples (200 ng) subjected to no enzyme or restriction endonuclease treatments. The negative peaks on digested samples are suspected to be coming from storage solvent of restriction endonucleases.

## Conclusion

Slalom chromatography by GTxResolve 250 Å Slalom Column is highly suitable for plasmid analysis at various steps in both cell culture processing and cell-free syntheses.

- a. Extent of plasmid linearization before synthesis of mRNA drug substance by in vitro transcription.
- b. Plasmid quality evaluation to detect the levels of open circular and linear forms of impurities on constructs ≥7 kb.
- c. Confirmation of gene inserts in recombinant plasmids by digestion with appropriate restriction endonucleases.
- d. Sensitive detection of plasmid digests with 5x improvement compared to agarose gel electrophoresis.

As such, slalom chromatography with a GTxResolve 250 Å Slalom Column provides a viable high-throughput alternative to tedious agarose gel electrophoresis methods. Its quick readout on plasmid DNA samples in <5 minutes will be helpful to a number of different molecular biology and process development laboratories.

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