

# **Application** News

# No. L561

#### **High Performance Liquid Chromatography**

### High Speed GPC Analysis by Nexera™ GPC System: **Calculation of Molecular Weight and Quantitative Analysis of Additives**

#### Introduction

Measurement of the molecular weight distribution of polymers is one field of HPLC utilizing the size exclusion mode and has been called GPC (gel permeation chromatography) for a long time. Recent requirements for increased throughput by high speed analysis are also considered as a priority in GPC, which is an established analytical technique. In this article, a high speed GPC analysis was carried out using a GPC system consisting of a high speed GPC column packed with smaller particles and a Nexera XR UHPLC system that absolutely minimizes external column dispersion to verify its performance. In addition, this article also introduces a simultaneous determination of low molecular weight polymer additives within a high speed GPC analysis using an additional PDA (photodiode array) detector and a deconvolution function provided in the LabSolutions™ workstation to show an improved efficiency of GPC analysis, whereas ordinary quantitative analysis of these additives is carried out by HPLC separation mode other than GPC such as reversed phase mode after fractionation by preparative LC that requires a long time and tedious procedures.

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### High Speed GPC Analysis

A refractive index detector (RID), which responds to the sample weight, is commonly used in GPC analysis for calculation of the average molecular weight and the polydispersity of polymer compounds, but RID is known to be sensitive to temperature changes. Moreover, because GPC calculation is affected not simply by the peak area, but by the shape of the peak over the entire elution interval, the evaluation of the repeatability of GPC calculation is severer in comparison with that of ordinary HPLC determination. In the analysis described here, a high speed GPC analysis was carried out using a GPC column for rapid analysis having a particle size of 3.5 µm and a Shodex™ RI-504, which is a RID that supports semi-microscale analysis. The average molecular weight and the polydispersity were measured, and the repeatability of five consecutive analyses was verified. Table 1 shows the details of the analytical conditions. The table also shows the settings of the PDA (photodiode array) detector, which is discussed on the following page. Fig. 1 shows the calibration curve created for GPC calculation using polystyrene standard markers.

Table 1 Analytical Conditions			
System	: Nexera XR GPC		
Column	: Shodex GPC HK-404L (150 mm $\times$ 4.6 mm I.D., 3.5 $\mu$ m)		
Mobile phase	: THF		
Flow rate	: 0.8 mL/min		
Column temp.	: 40 °C		
Injection vol.	: 5 μL		
Detection	: RI-504 (RID) Cell temp.: 40 °C SPD-M40 (PDA, UHPLC cell equipped) Wave length range: 190-400 nm Cell temp.: 40 °C Response: 0.025 s Sampling: 0.025 s		
Sample	: 0.2% (w/v)		
Analysis time	: 3.5 min		

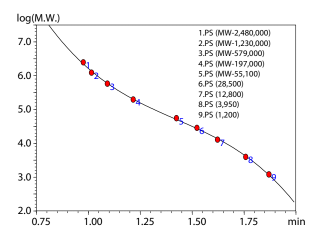


Fig. 1 Calibration Curve Created Using Polystyrene Standard Markers

#### Results of GPC Analysis and Discussion

Fig. 2 shows the high speed GPC chromatogram obtained with the RID. The elution volume around the actual permeation limit appears to be approximately 1.6 to 2 mL. However, considering the stability of the baseline, the time per one analysis in continuous analysis, or so-called cycle time, was set to approximately 3.5 min. In addition, the molecular weight corresponding to the peak front was also about 600,000, suggesting that an appropriate GPC calculation was executed using the calibration curve shown in Fig. 1.

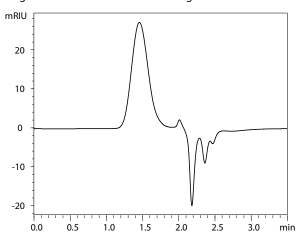


Fig. 2 Chromatogram of Polystyrene Sample by High Speed

Fig. 3 shows a comparison of the chromatograms obtained by five consecutive analyses by high speed GPC.

Table 2 summarizes the number average molecular weight, weight average molecular weight, and polydispersity, together with their repeatabilities. Stable consecutive analyses were able to be done even with a short cycle time of 3 to 4 min, and satisfactory GPC calculation repeatability was obtained.

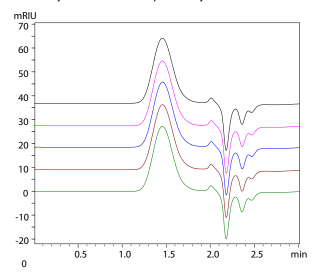


Fig. 3 Comparison of Chromatograms in Continuous High Speed GPC Analysis

Table 2 GPC Calculation Results of Polystyrene Sample (n=5)

	Mn	Mw	Mw/Mn
Polystyrene	$2.73 \times 10^{4}$	$4.67 \times 10^{4}$	1.71
%RSD	0.89	0.68	0.23

#### Deconvolution of Unseparated Additives in Polymer Sample by i-PDeA II

In quantitative analysis of polymer additives, complete separation of two or more low molecular weight additives is generally difficult using only GPC columns. Consequently these additives are collected from the polymer by preparative LC or the like followed by for example reversed phase chromatography, which can be expected to provide higher separation performance.

However, it is possible to obtain deconvoluted chromatographic approximate solutions for the respective compounds by simulation using the *i*-PDeA II (Intelligent Peak Deconvolution Analysis II) function of the LabSolutions workstation, which deconvolutes unseparated peaks based on the spectral information obtained with a PDA detector, and the chromatographic information obtained by deconvolution can be used as is in quantitative determinations. Moreover, deconvolution is not limited to chromatograms, but can also be applied to spectral analysis, also contributing to enhancement of the qualitative analysis performance. Details concerning *i*-PDeA II can be found in Shimadzu Technical Report No. C191-E042.

Among the additives contained in the polystyrene sample analyzed in this article, elution times were confirmed for Irganox® 1010 and Tinuvin® 144, for which standard compounds were able to be prepared, and these compounds showed almost same elution position at UV254 nm in the PDA chromatogram in Fig. 4. As shown by the data processing correction line in Fig. 4, these two compounds formed an unseparated peak. Fig. 5 shows the overlaid deconvoluted chromatograms of the respective compounds obtained by *i*-PDeA II processing during the elution interval of polystyrene and low molecular weight additives. The peaks originating from Irganox 1010 and Tinuvin 144 were deconvoluted to traces #2 and #4, and their content ratios in the polystyrene were quantitively determined as 26.9 mg/g and 3.9 mg/g, respectively, using calibration curves created separately based on their peak areas.

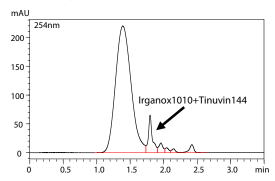


Fig. 4 PDA Chromatogram of Polystyrene Containing Additives

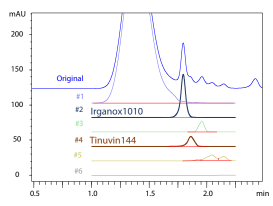


Fig. 5 Result of Deconvolution of Polystyrene and Additive During Elution Interval

#### Conclusion

The results of this experiment confirmed that the Shimadzu Nexera XR GPC system was able to be applied to high speed GPC analysis with equivalent reliability to that of conventional GPC analysis in a shorter time. Although complete separation and quantitative determination of low molecular weight additives were difficult when using only a conventional GPC column, this experiment demonstrated that the separation and quantitative analysis of these additives were possible simultaneously within a high speed GPC analysis by the *i*-PDeA II deconvolution function of the LabSolutions workstation.

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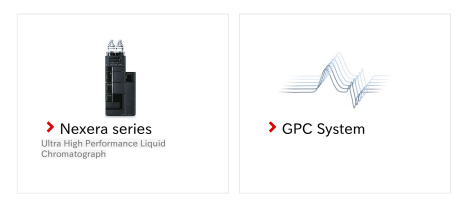
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