



EcoSEC High Temperature GPC System

Experts in Chromatography

EcoSEC® High Temperature GPC System
TOYOPEARL® & TSKgel® Bulk Resin
TSKgel HPLC Columns
EcoSEC GPC System



2015

Product Guide

TOSOH BIOSCIENCE

A Tradition of GPC Excellence

Tosoh Bioscience established itself as a world leader in the field of polymer analysis in 1971 with the introduction of TSKgel gel permeation chromatography (GPC) columns. The following year, Tosoh launched a dedicated instrument for GPC analysis. Since that first instrument there have been 7 generations of GPC systems with temperature control up to 50 °C, as well as 2 generations of high temperature GPC systems for analysis up to 220 °C. Today Tosoh Bioscience continues a tradition of GPC excellence with a 3rd generation high temperature GPC system: the EcoSEC High Temperature GPC System.

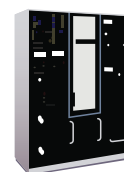
1972: HLC-801 GPC System

- First GPC instrument from Tosoh
- "All-in-one" concept incorporated



1977: HLC-811 HT GPC System

- First Tosoh high temperature GPC System



1982: HLC-802A GPC System

- Dual Flow refractive index (RI) detector
- "Stable RI Baseline" implemented



1986: HLC-8020 GPC System

- Temperature controlled pump system
- "High Reproducibility" obtained



1993: HLC-8120 GPC System

- First semi-micro GPC columns from Tosoh
- "Semi-micro" concept incorporated



1998: HLC-8121 HT GPC System

- 2nd generation Tosoh high temperature GPC System



2008: EcoSEC GPC System

- 7th generation Tosoh GPC System
- Released in overseas market



2013: EcoSEC High Temperature GPC System

- 3rd generation Tosoh high temperature GPC System
- Released in overseas market



History of *Performance*

With over 40 years in the GPC market Tosoh Bioscience is proud of our latest EcoSEC High Temperature GPC System. Designed by engineers, built by dedicated employees, tested by experienced polymer scientists, and supported by an entire organization, you can trust your analyses to the EcoSEC High Temperature GPC System.



Demanding high temperature analyses require a system that delivers results reliably, reproducibly, and produces these results in an easy to use, safe instrument. The EcoSEC High Temperature GPC System incorporates the proven design and technology used in our EcoSEC GPC System. The dual pump system, dual flow RI detector, spacious column oven, total system temperature control, autosampler, sample prep station, safety features, and all-in-one design controlled by intuitive software combine to deliver *Performance*.

Out of the box, the new EcoSEC High Temperature GPC System features the following:

- **Reliability**
- **Reproducibility**
- **Stability**
- **Safety**
- **All-in-One Design**
- **Ease of Use**

Engineered for *Performance*



Component

Description

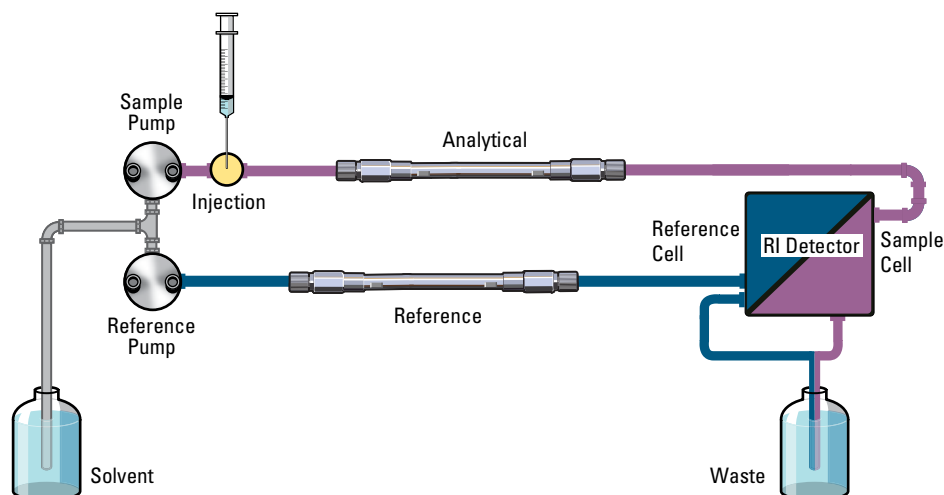
Benefit

Component	Description	Benefit
Solvent Holder	Maintains a constant temperature of 40 °C.	Prevent possible solvent freezing.
Control Panel	Allows the system to be controlled manually and at the discretion of the operator.	Saves time by controlling a series of operations without the use of the computer or software.
Temperature Controlled Pumps	Pump heads and solvent lines are maintained at a constant temperature.	Improves baseline stability by removing the effect of temperature fluctuations. This results in consistent and accurate flow rates and reproducible molar mass determinations.
Column Oven	Maintains 40 - 220 °C. Can accommodate up to 8, 30 cm length columns.	Constant column temperature ensures precise and reproducible molar mass determinations.
Autosampler	24 sample capacity. Temperature controlled by aluminum block from 40 - 220 °C.	Precise injection volume. Variety of loop sizes. Door is locked under sampling operation for safety.
RI Detector	Solvent flows through a separate reference cell. 10 µL volume flow cell.	Temperature controlled, stable baseline, quick response, low noise. Enhanced baseline stability from dual flow cell RI detector.
Purge unit and degasser	Variable degassing capacity. Temperature controlled degassing unit and auto purge function.	Saves time with rapid solvent changes via purge valve eliminating solvent replacement and other time-consuming manual operations.

Performance means Baseline Stability

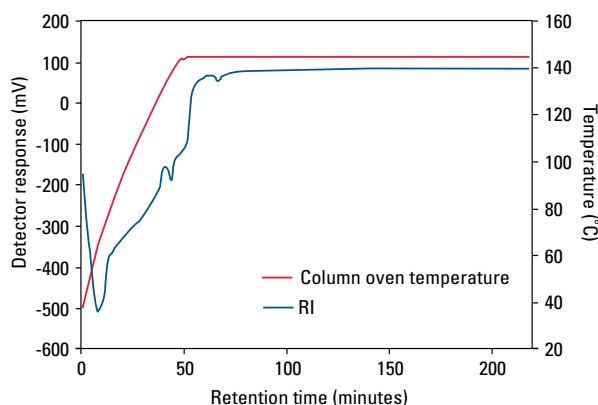
Incorporated into the design of our two pump delivery system is 40+ years experience in engineering. The EcoSEC High Temperature GPC System has a unique dual flow design which includes the use of two pumps. **Figure 1** demonstrates the flow paths of the sample and reference pumps. The sample pump flows solvent from the solvent reservoir through the following system components in sequence: autosampler, analytical column, sample side of RI detector cell, and waste container. The solvent flows via the reference pump from the solvent reservoir through a reference column, the reference side of the RI detector cell, and then the waste container. The entire flow system is temperature controlled to eliminate the effects of fluctuations in ambient temperature.

Figure 1: Flow paths of sample and reference pumps in the EcoSEC High Temperature GPC System



On the EcoSEC High Temperature GPC System the RI baseline is considered stabilized when the drift in the signal is 3.0×10^{-7} RIU/h or less. When a new set of columns is manually placed on the EcoSEC High Temperature GPC System and the flow rate and temperature controls are started, the RI baseline stabilizes within 3 hours. **Figure 2** demonstrates the equilibration time from start-up of the EcoSEC High Temperature GPC System in orthodichlorobenzene (ODCB).

Figure 2: Refractive index detector signal during equilibration of the EcoSEC High Temperature GPC System

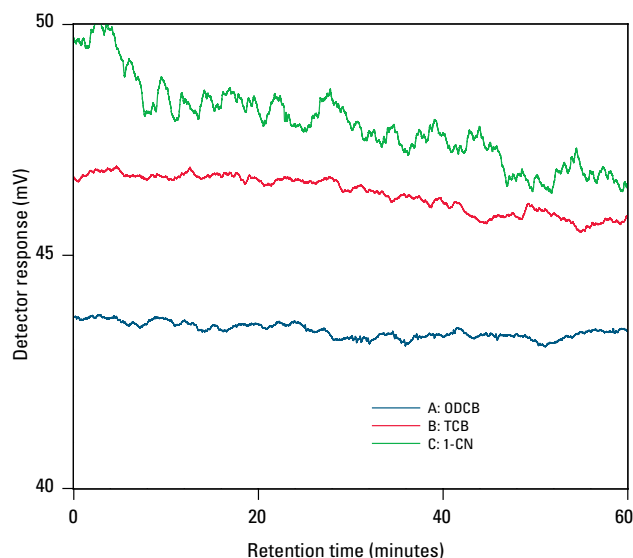


Column: TSKgel GMH_{HR}-H (S) HT2, 13 μm,
7.8 mm ID × 30 cm × 2
Mobile phase: ODCB with 0.05% BHT
Flow rate: 1.0 mL/min
Detector: RI (EcoSEC High Temperature GPC System)
Temperature: 145 °C

Performance means Baseline Stability

Advanced engineering, along with complete temperature control and a dual flow RI detector, means rock steady baselines in even the most challenging solvents and temperatures. The RI baselines as obtained for three commonly used high temperature GPC solvents: Trichlorobenzene (TCB) at 145 °C, orthodichlorobenzene (ODCB) at 145 °C and 1-chloronaphthalene (1-CN) at 210 °C are shown in Figure 3. The RI baseline drift for all three solvents is less than 1 mV/h.

Figure 3: Baseline drift of the dual flow refractive index detector of the EcoSEC High Temperature GPC System for TCB, ODCB, and 1-CN



Column: TSKgel GMH_{HR}-H (S) HT2, 13 µm,
7.8 mm ID × 30 cm × 2
Mobile phase: A: ODBC
B: TCB
C: 1-CN
Flow rate: 1.0 mL/min
Detector: RI (EcoSEC High Temperature GPC System)
Temperature: A and B: 145 °C
C: 210 °C

The unmatched baseline stability of the dual flow RI detector in the EcoSEC High Temperature GPC System is also shown in Table 1 through the drift, fluctuation, and noise obtained when ODCB at 145 °C, TCB at 145 °C, 1-CN at 210 °C, and THF at 40 °C are used as the mobile phase.

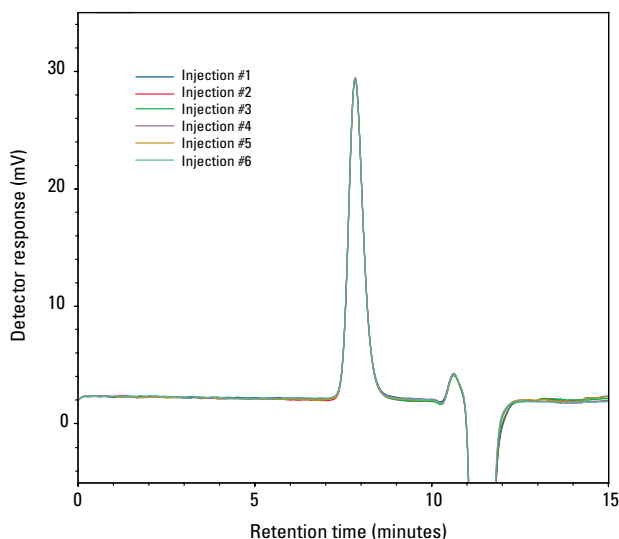
Table 1: Baseline drift, fluctuation and noise of the EcoSEC High Temperature GPC System in ODCB, TCB, 1-CN, and THF

Solvent (temperature)	Drift (mV/h)	Fluctuation (mV)	Noise (mV)
ODCB (145 °C)	-0.41	0.54	0.044
TCB (145 °C)	-1.30	0.69	0.046
1-CN (210 °C)	-0.91	1.61	0.098
THF (40 °C)	-0.35	0.23	0.022

Performance means Reproducibility

The dual flow design of the RI detector and the temperature controlled pumps of the EcoSEC High Temperature GPC System deliver precise flow rates at all temperatures, even when changes in environmental conditions occur, thus producing reproducible results sample after sample, day after day. The intraday and day-to-day reproducibility of the EcoSEC High Temperature GPC System are shown in Figure 4.

Figure 4: GPC elution profile of intraday reproducibility of the EcoSEC High Temperature GPC System



Reproducibility (intraday, n=6)

R.T.: CV 0.017%

Area: CV 0.42%

Reproducibility (day to day, n=5)

R.T.: CV 0.047%

Area: CV 0.71%

Column: TSKgel GMH_{HR}-H (S) HT2, 13 μ m,
7.8 mm ID \times 30 cm \times 2

Mobile phase: ODCB with 0.05% BHT

Flow rate: 1.0 mL/min

Detector: RI (EcoSEC High Temperature GPC System)

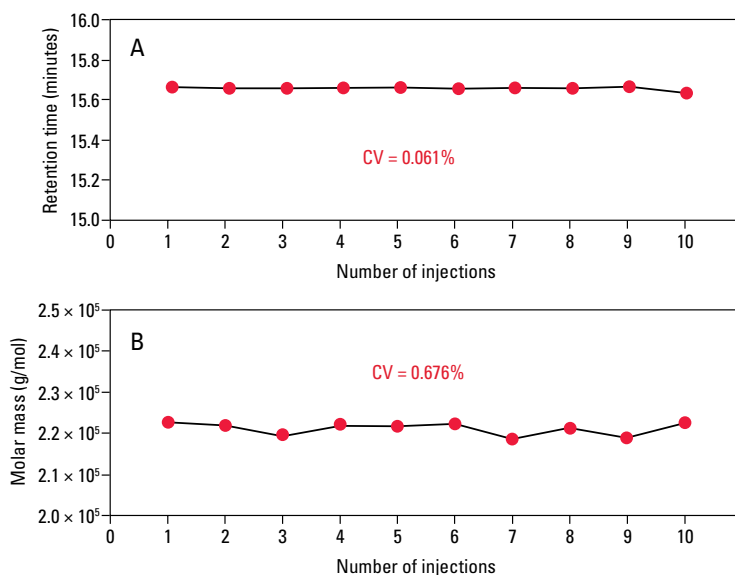
Temperature: 145 $^{\circ}$ C

Injection vol.: 300 μ L

Sample: polystyrene (F-20), 0.02%

The engineering design concepts of the EcoSEC High Temperature GPC System result in a high degree of reproducibility of retention times (Figure 5A) and molar mass determinations (Figure 5B). The coefficients of variation for retention time and weight-average molar mass, M_w , are well below 1% for successive injections.

Figure 5A and 5B: A: Intraday retention time reproducibility, B: Intraday weight-average molar mass reproducibility



Column: TSKgel GMH_{HR}-H (S) HT2, 13 μ m,
7.8 mm ID \times 30 cm \times 2

Mobile phase: ODCB with 0.05% BHT

Flow rate: 1.0 mL/min

Detector: RI (EcoSEC High Temperature GPC System)

Temperature: 145 $^{\circ}$ C

Injection vol.: 300 μ L

Sample: polypropylene

Performance Proof - Applications

Polyphenylene Sulfide

Polyphenylene Sulfide (PPS) has attracted a considerable amount of interest in the polymer industry due to its high tensile strength, good dimensional stability, flame resistance, and excellent stability in organic liquids. PPS is virtually insoluble in most organic solvents at ambient temperatures and thus can only be characterized in the solid state or by using elevated temperatures. The limited solubility of PPS makes it very difficult to determine macromolecular properties, such as molar mass and molar mass distribution, that play a vital role in the determination of mechanical, bulk and solution properties of the processing and end-use properties of a given material. Traditionally, PPS has been characterized by infrared spectrometry and thermal analysis methods. One method which can also be used to characterize PPS is high temperature GPC as PPS is soluble in 1-chloronaphthalene (1-CN) at extremely elevated temperatures (> 200 °C). 1-CN is a difficult solvent to use for analytical experiments as the solvent ambers over time and can cause havoc for detection methods such as RI. GPC analysis of PPS in 1-CN for the determination of molar mass averages and molar mass distributions is possible using the EcoSEC High Temperature GPC System due to the unique dual flow refractive index detector.

A new and a used PPS sample were compared for failure investigation through their GPC elution profiles, **Figure 6**, and their polystyrene relative molar mass averages, **Table 2**. As seen in **Figure 6**, the new PPS sample eluted prior to the used PPS sample. The shorter retention time of the new PPS sample indicated that the new PPS sample was larger in polymeric size than the used PPS sample, as the elution order in GPC is that of an "inverse-sieving" technique, larger analytes sample a smaller pore volume than smaller analytes resulting in the larger analytes eluting from the GPC column prior the smaller analytes. As seen in **Table 2**, the new PPS sample was determined to have a higher number-, weight-, and z-average molar mass and greater polydispersity index, *PDI*, than the used PPS sample. The approximately 20 to 50% decrease in the molar mass averages and 25% increase in *PDI* observed between the new PPS and the used PPS is potentially enough evidence to determine that after a predetermined amount of time the end-use product(s) made with this PPS sample will begin to fail or will no longer be able to perform up to standards. The use of GPC/RI for the failure investigation of PPS allows for immediate differentiation between the new and used PPS samples based on the GPC/RI elution profile, which was then confirmed through differences in the polystyrene relative molar mass averages and molar mass distributions between the new and used PPS samples.

Figure 6: GPC elution profile of new and used PPS samples as monitored by RI

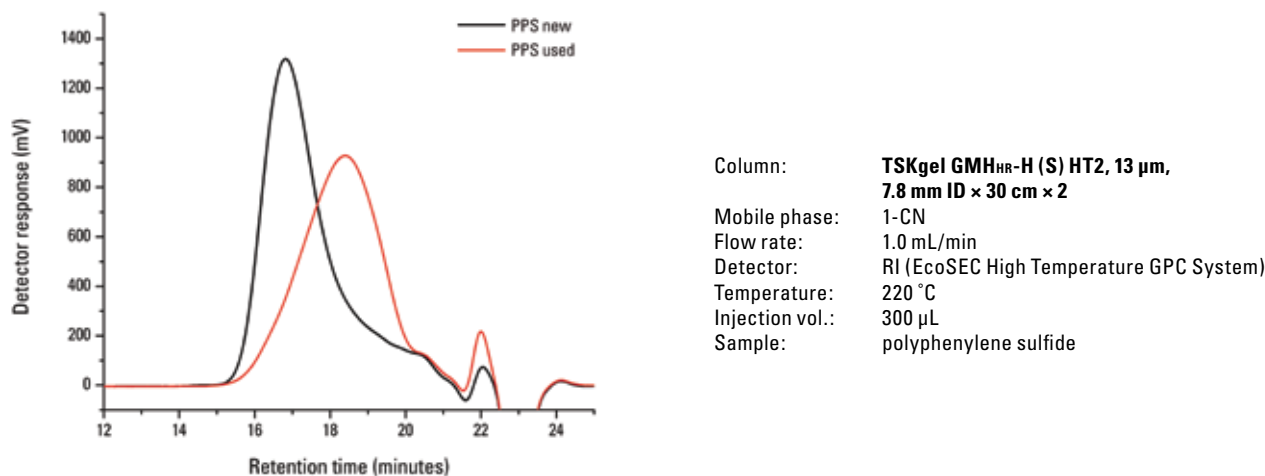


Table 2: Molar mass averages and polydispersity index of new and used PPS samples via GPC/RI

Sample	M_n (g/mol)	M_w (g/mol)	M_z (g/mol)	<i>PDI</i> ^a
PPS new	5,790	3.91×10^4	7.19×10^4	6.74
PPS used	3,176	1.62×10^4	5.54×10^4	5.10

^a $PDI = M_w/M_n$

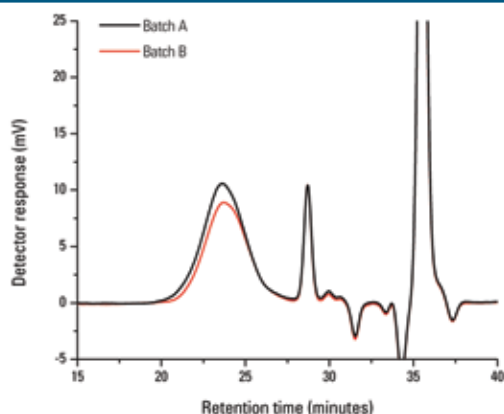
Performance Proof - Applications

Polyethylene

One of the most common plastics and commercially available polymers on the market is polyethylene. Polyethylene in general describes a huge family of resins obtained by the polymerization of ethylene gas. Polyethylene is available in a range of flexibilities and properties depending on the production process. Properties of polyethylene such as toughness, hardness, and clarity can be regulated by altering the molar mass averages, comonomer type, and comonomer content. Most polyethylene resins for commercial products are fabricated by controlling the molar mass average, molar mass distribution and branching characteristics. The molar mass averages and molar mass distributions of polyethylene can be determined using the EcoSEC High Temperature GPC System.

High temperature GPC experiments provide two forms of comparison between the two difference batches of polyethylene samples: GPC chromatograms and polystyrene relative molar mass averages and molar mass distributions. **Figure 7** shows the GPC elution profiles as monitored by the RI detector in the EcoSEC High Temperature GPC system for the difference batches of polyethylene. Batch B extends further in the larger polymeric size, shorter retention time direction of the GPC elution profile than Batch A, an indication that the two batches differ slightly in polymeric size, as elution order in GPC is that of an “inverse-sieving” technique, as smaller analytes elute after larger analytes.

Figure 7: GPC elution profile of two batches of polyethylene as monitored by RI



Column: **TSKgel GMH_{HR}-H (S) HT2, 13 μm, 7.8 mm ID × 30 cm × 2**
 Mobile phase: ODCB with 0.05% BHT
 Flow rate: 1.0 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 135 °C
 Injection vol.: 300 μL
 Sample: polyethylene

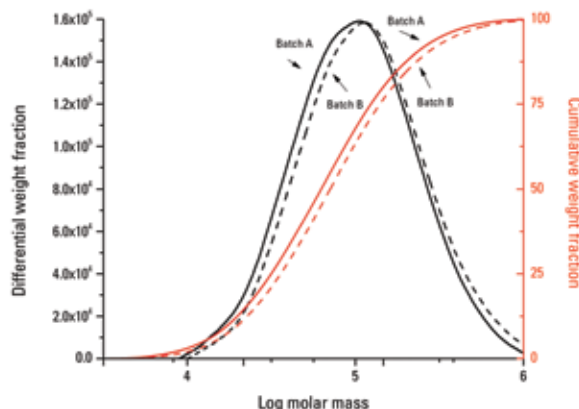
The molar mass averages and polydispersity index, *PDI*, as determined by the polystyrene RI calibration curve are given in **Table 3**. A comparison of the molar mass averages and molar mass distribution, **Figure 8**, of the two different batches of polyethylene reveals an approximately 10 to 15% difference in the polystyrene molar mass averages and distributions between the two batches. The molar mass averages and distributions of the two different batches of polyethylene obtained by high temperature GPC are different enough to distinguish the two batches from one another but may be similar enough to both create a successful commercial plastic with the same end-use properties.

Table 3: Molar mass averages and polydispersity index of two batches of polyethylene via GPC/RI

Sample	M_n (g/mol)	M_w (g/mol)	M_z (g/mol)	<i>PDI</i> ^a
Batch A	4.48×10^4 ± 364 ^b	1.18×10^5 ± 790	2.95×10^5 ± 1,821	2.64 ± 0.06
Batch B	3.66×10^4 ± 135	1.03×10^5 ± 124	2.64×10^5 ± 2,806	2.80 ± 0.01

^a $PDI = M_w/M_n$

Figure 8: Overlay of cumulative and differential molar mass distribution of two batches of polyethylene



Performance Proof - Applications

Polythiophene

Conducting polymers, such as polythiophenes, have been widely investigated over the past several decades due to their potential industrial applications based on their conductivity and organic light-emitting capability. To date polythiophenes have been used in the development of electronics, energy storage batteries, photochromic devices and nonlinear optical devices. The heavy focus on synthesis of conducting polymers facilitates the need for characterization methods. Among the methods employed for the characterization of the intermediates and final conducting polymers are FT-IR, NMR, GPC, and microscopy. Some conducting polymers have limited solubility thus require the use of high temperature GPC for determination of the molar mass averages and molar mass distributions. Similar to other polymers, the molar mass averages and molar mass distributions of conducting polymers play a role in determining the end-use properties of the applications for which the polymer is used.

The molar mass averages and molar mass distributions of two conducting polymers similar to polythiophene were determined using the EcoSEC High Temperature GPC System. The polystyrene relative molar mass averages, M_n , M_w , and M_z , are given in Table 4. The variation between the molar mass averages of the two conducting polymers may be enough to change the conductivity of the polymers, thus their end-use applications. In addition to the molar mass averages, the molar mass distribution can also influence various properties of conducting polymers. The molar mass distributions of the two conducting polymers are compared in Figure 9. The molar mass distribution of polymer A is significantly larger than that of polymer B.

Information regarding the difference between the two conducting polymers can be seen by comparing their GPC elution profiles, Figure 10. The shift in GPC retention time amongst the two conducting polymers indicates a variation in polymeric size between the two conducting polymers, as elution order in GPC is that of an "inverse-sieving" technique, large analytes sample a smaller pore volume than smaller analytes resulting in larger analytes eluting from the GPC column prior to the smaller analytes. Based on the GPC elution profile, polymer A is significantly larger in polymeric size than polymer B.

Table 4: Molar mass averages and polydispersity index of two conducting polymer samples via GPC/RI

Sample	M_n (g/mol)	M_w (g/mol)	M_z (g/mol)
Polymer A	2.58×10^4 $\pm 0.01 \times 10^4$	6.51×10^4 $\pm 0.02 \times 10^4$	1.34×10^5 $\pm 0.03 \times 10^5$
Polymer B	9.39×10^3 $\pm 0.01^a \times 10^3$	1.26×10^4 $\pm 0.04 \times 10^4$	1.60×10^4 $\pm 0.01 \times 10^4$

^a Standard deviation from two injections

Figure 9: Overlay of cumulative and differential molar mass distribution of two conducting polymer samples

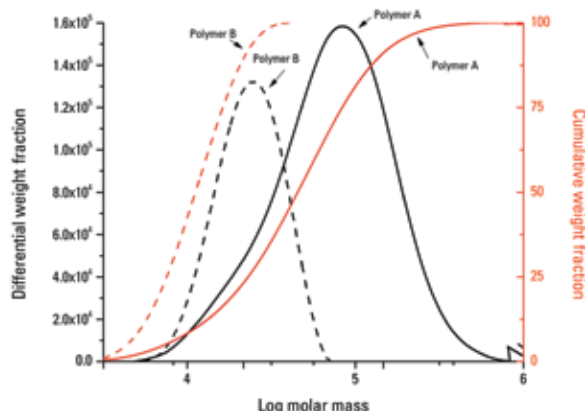
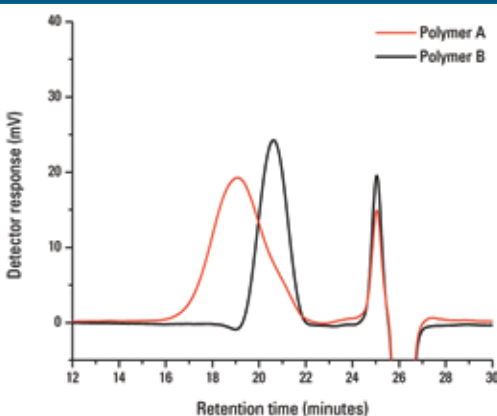


Figure 10: GPC elution profile of two conducting polymer samples as monitored by RI



Column: **TSKgel GMH_{HR}-H (S) HT2, 13 μ m, 7.8 mm ID \times 30 cm \times 2**
 Mobile phase: TCB
 Flow rate: 1.0 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 135 $^{\circ}$ C
 Injection vol.: 300 μ L
 Sample: polythiophene

Performance Proof - Applications

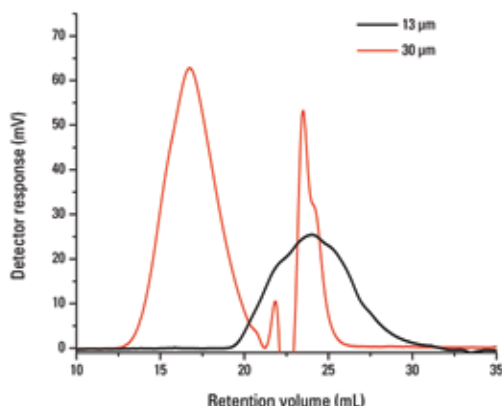
High Molar Mass Polymers

High temperature GPC is a common and important technique used for the characterization of polyolefins. GPC analysis of polyolefins can be difficult as those containing over 10% ethylene and polypropylene monomers have limited solubility due to their characteristically high strength and toughness that results from their high crystallinity. In addition to the limited solubility of most polyolefins, high molar mass polyolefins, such as ultra-high molar mass polyethylene (UHMPE) present their own subset of issues when being analyzed by GPC. High molar mass polyethylenes are extremely long polymer chains with a molar mass greater than 2×10^6 g/mol. Polymers greater than a million in molar mass have been shown to experience on-column flow induced degradation when analyzed by GPC. To decrease the amount of degradation that occurs when UHMPE samples are analyzed by high temperature GPC and thus obtain the most accurate molar mass averages and molar mass distributions, GPC columns packed with larger size particles with large pores are ideal.

An EcoSEC High Temperature GPC System with a dual flow refractive index detector was used in conjunction with 13 μm and 30 μm TSKgel high temperature GPC columns to determine the molar mass averages and distributions of a UHMPE. Figure 11 show the GPC elution profiles obtained on both column sets. The shape of the GPC elution profile varies between the two column sets. The elution profile obtained using the 13 μm high temperature GPC column has a shoulder in the high molar mass region (the molar mass region most likely affected by on-column flow induced degradation) while the elution profile obtained using the 30 μm high temperature GPC column does not.

The polystyrene RI relative molar mass averages of the UHMPE obtained using two different high temperature GPC column sets are given in Table 5. The molar mass averages obtained using the 13 μm high temperature GPC columns are significantly smaller than those obtained using the 30 μm high temperature GPC columns. The sample degradation is more prevalent in the high molar mass region of the sample as the z-average molar mass is two orders of magnitude greater when analysis is performed on the 30 μm high temperature GPC columns. The molar mass distribution of the UHMPE obtained by both high temperature GPC column sets indicate an extremely polydisperse polymer. The use of 30 μm high temperature GPC columns provides a better representation of the polystyrene relative molar mass averages as the larger size particles and pores decrease the amount of degradation experienced by UHMPE.

Figure 11: GPC elution profile of UHMPE samples as monitored by RI with 13 μm and 30 μm TSKgel high temperature GPC columns



Column: TSKgel GMH_{HR}-H (S) HT2, 13 μm , 7.8 mm ID \times 30 cm \times 2 + TSKgel G2000H_{HR}(20) HT2, 20 μm , 7.8 mm ID \times 30 cm
 Mobile phase: TCB
 Flow rate: 1.0 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 135 $^{\circ}\text{C}$
 Injection vol.: 300 μL
 Sample: polyethylene

Table 5: Molar mass averages and polydispersity index of UHMPE samples via RI with 13 μm and 30 μm TSKgel high temperature GPC columns

Column (particle size)	M_n (g/mol)	M_w (g/mol)	M_z (g/mol)	PDI^a
13 μm	2.23×10^4	5.76×10^5	4.41×10^6	25.75
30 μm	9.21×10^4	7.74×10^6	2.55×10^8	84.07

^a $PDI = M_w/M_n$

Performance Software

EcoSEC High Temperature GPC System Workstation Software

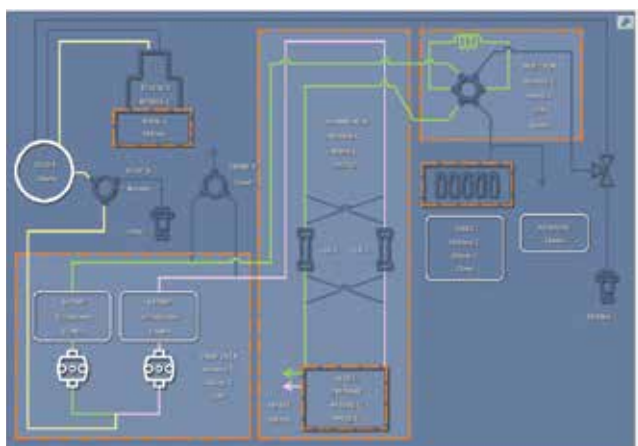
- GPC-specific EcoSEC High Temperature GPC System software to simplify system control and data handling
- Controls up to 2 EcoSEC High Temperature GPC Systems
- Excellent data handling and report generation
- Fully featured data handling system; analyze data from two detectors
- Start and stop system automatically
- One license for multiple locations

Features include:

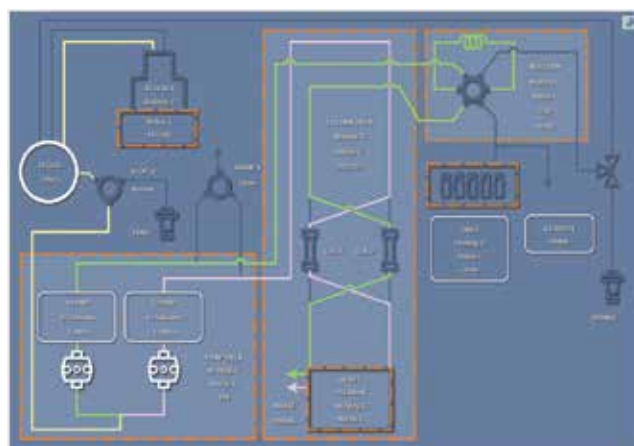
Flow Diagram

- Unique screen allows you to easily modify running conditions of an individual component

Typical flow

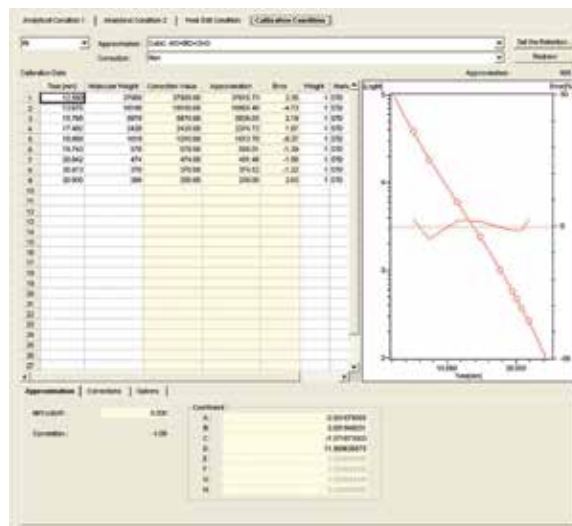


With use of column switching valve



Method

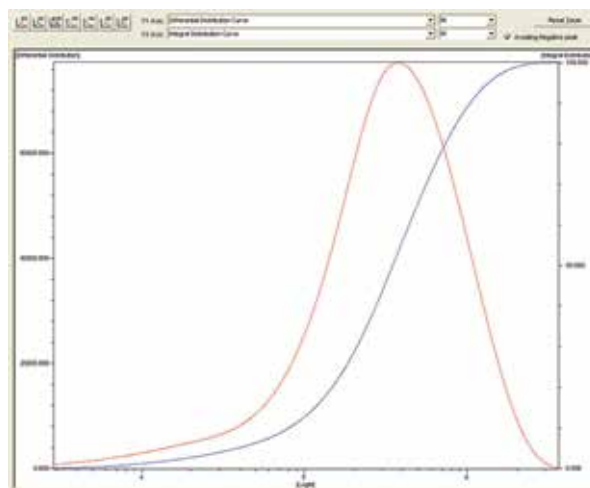
- All parameters for data acquisition and peak integration, including baseline operations, are saved in the template method
- One click switching between calibration curves



Performance Software

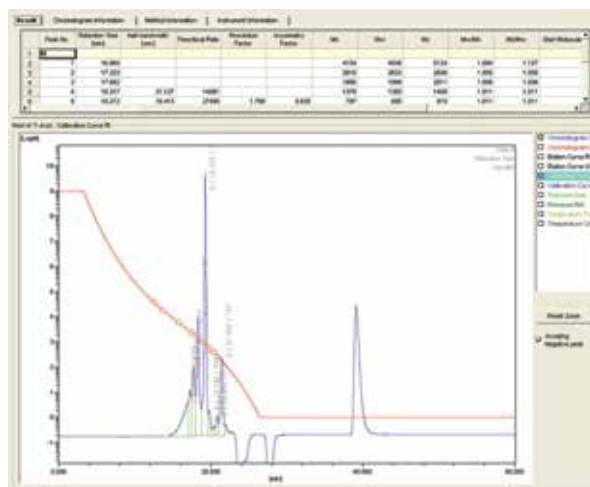
GPC Specific Quantitative Calculations

- M_n , M_w , and M_z molar mass averages
- Cumulative and differential molar mass plotting
- Polydispersity index (*PDI*) values



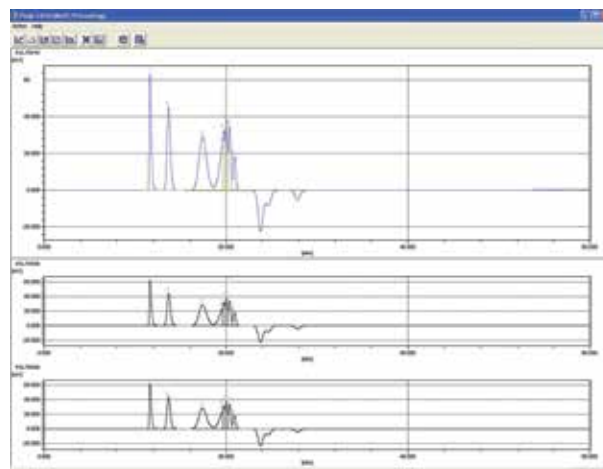
Data Management and Report Generation

- Allows viewing of chromatograms, elution, curve, flow rate, pressure, and temperature.
- Large number of built in reports
- Fully customizable reports
- Easily export data into text or pdf files.



Peak Editing and Multiprocessing Function

- Full editing functionality including baseline setting and peak splitting using the mouse
- Automatic peak editing
- Automatic application of peak detection and integration parameters to multiple chromatograms of the same sample using the multiprocessing function; resulting in identical processing for similar chromatograms for enhanced reproducibility.



Accessories to Maximize *Performance*

Sample Prep System

- Sample shaker 10 - 100 RPM
- 24 vial capacity
- Aluminum heated block
- 40 - 220 °C



Column Switching Valve

- Easily change between 2 column sets
- Equipped above column oven
- Manual switching
- Position is recognized by software



External Light Scattering Detector

- DAWN® HELEOS™ II and DAWN 8+ from Wyatt Technology
- Integrated heated transfer lines
- Absolute molar mass averages and radius of gyration
- 40 - 210 °C

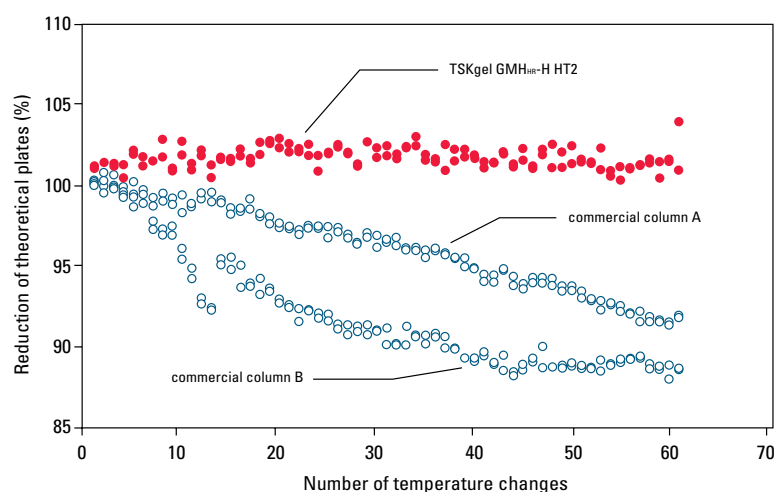


The Heart of Performance – Columns

TSKgel H series columns are recommended for the analysis of organic-soluble polymers and are packed with spherical particles composed of polystyrene cross-linked with divinylbenzene (PS-DVB). The “GM” prefix denotes a column packed with particles of different pore sizes blended to provide an extended linear calibration curve. The TSKgel HT columns are for high temperature applications ($\leq 140\text{ }^{\circ}\text{C}$) while the TSKgel HT2 columns are used in ultra-high temperature (up to $220\text{ }^{\circ}\text{C}$) applications.

Figure 12 demonstrates the performance stability of the TSKgel GMH_{HR}-H HT columns compared to other commercially available high temperature GPC columns during repetitive temperature changes. The TSKgel H_{HR} HT columns and two commercially available high temperature GPC columns were subjected to drastic changes in temperature by raising the temperature for 2 hours followed by lowering the temperature for two hours for a total of 60 cycles. The number of theoretical plates was shown to remain constant for the TSKgel H_{HR} HT columns and to decrease by 15% for the two commercially available high temperature GPC columns; thus revealing the superior performance stability of the TSKgel H_{HR} HT columns.

Figure 12: Durability of TSKgel H_{HR} HT columns compared to two commercially available high temperature GPC columns



Column: **TSKgel GMH_{HR}-H HT2, 5 μm , 7.8 mm ID \times 30 cm \times 2**
 Mobile phase: ODCB with 0.05% BHT
 Flow rate: 1 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 40 to 145 $^{\circ}\text{C}$



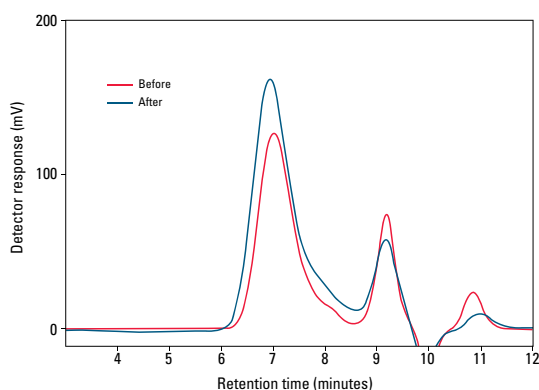
The Heart of Performance – Columns

Column Durability at 220 °C

Column durability in high temperature GPC polymer analysis is essential as these columns are continuously exposed to harsh organic solvents, extremely elevated temperatures and temperature cycling as GPC systems are turned on and off. The durability of a high temperature GPC column directly influences the quality, applicability and selectivity, or resolution, of the GPC column, thus the accuracy of the molar mass averages obtained. As a high temperature GPC column begins to fail or lose resolution due to the extreme experimental conditions required for high temperature GPC polymer analysis, the number- and z-average molar mass values obtained become inflated and the GPC elution profile begins to shift due to a decrease in multiple factors that affect the ability of the columns to separate species varying in hydrodynamic volume.

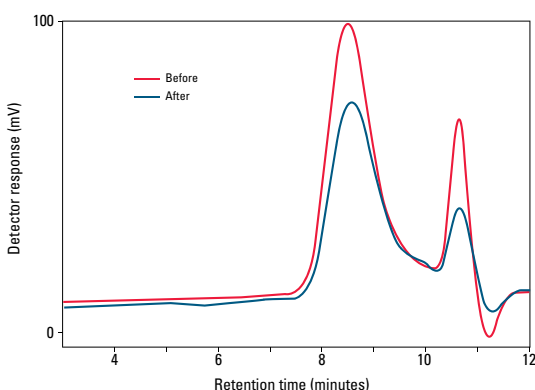
A durability and stability study of a TSKgel GMH_{HR}-H (S) HT high temperature GPC column was performed and the results were compared to another commercially available column for polymer analysis at 220 °C. The deterioration of the commercially available high temperature GPC column is observed in the GPC elution profiles, [Figure 13](#), as the resolution between the sample and solvent peaks decreases after the column is exposed to temperature cycling. The GPC elution profiles obtained for the TSKgel GMH_{HR}-H (S) HT column before and after temperature cycling remain superimposable, [Figure 14](#).

Figure 13: GPC elution profile for a polymer before and after temperature cycling obtained using a commercially available high temperature GPC column



Column: Commercially available high temperature GPC column, 13 μ m, 7.8 mm ID \times 30 cm
 Mobile phase: 1-CN
 Flow rate: 1.0 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 220 °C
 Injection vol.: 200 μ L
 Sample: synthetic polymer

Figure 14: GPC elution profile for a polymer before and after temperature cycling obtained using a TSKgel GMH_{HR}-H (S) HT column



Column: **TSKgel GMH_{HR}-H (S) HT, 13 μ m, 7.8 mm ID \times 30 cm**
 Mobile phase: 1-CN
 Flow rate: 1.0 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 220 °C
 Injection vol.: 200 μ L
 Sample: synthetic polymer

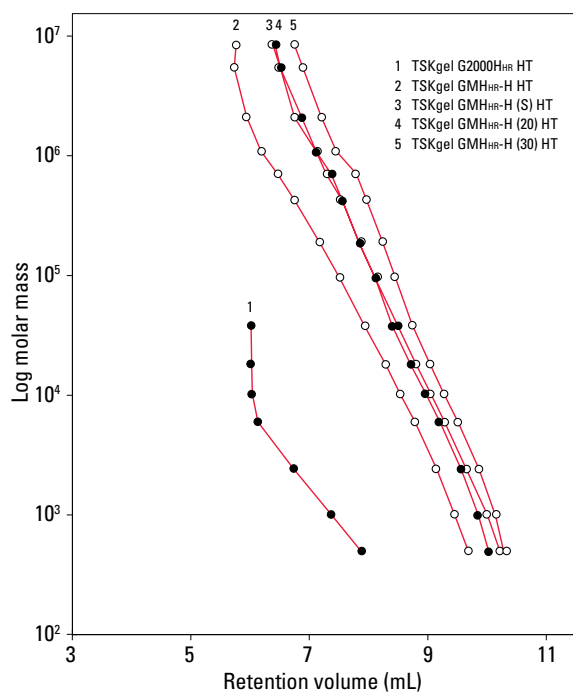
The Heart of Performance – Columns

Table 6 lists the attributes of the TSKgel HT columns which are for high temperature applications up to 140 °C.

Table 6: Properties and separation ranges for TSKgel HT columns

TSKgel column	Particle size	Pore size	Exclusion limit	Max. temp.
GMH _{HR} -H HT	5 µm	mixed pore sizes	4.0 × 10 ⁸ Da	140 °C
GMH _{HR} -H (S) HT	13 µm	mixed pore sizes	4.0 × 10 ⁸ Da	140 °C
GMH _{HR} -H (20) HT	20 µm	mixed pore sizes	4.0 × 10 ⁸ Da	140 °C
GMH _{HR} -H (30) HT	30 µm	mixed pore sizes	4.0 × 10 ⁸ Da	140 °C
G2000H _{HR} (20) HT	20 µm	2 nm	1.0 × 10 ⁴ Da	140 °C

Figure 15: Polystyrene calibration curves for TSKgel HT columns



Columns:
 TSKgel G2000H_{HR} (20) HT, 20 µm, 7.8 mm ID × 30 cm
 TSKgel GMH_{HR}-H HT, 5 µm, 7.8 mm ID × 30 cm
 TSKgel GMH_{HR}-H (S) HT, 13 µm, 7.8 mm ID × 30 cm
 TSKgel GMH_{HR}-H (20) HT, 20 µm, 7.8 mm ID × 30 cm
 TSKgel GMH_{HR}-H (30) HT, 30 µm, 7.8 mm ID × 30 cm

Mobile phase: ODCB with 0.05% BHT
Flow rate: 1.0 mL/min
Detector: RI (EcoSEC High Temperature GPC System)
Temperature: 135 °C
Injection vol.: 300 µL
Sample: polystyrene standards

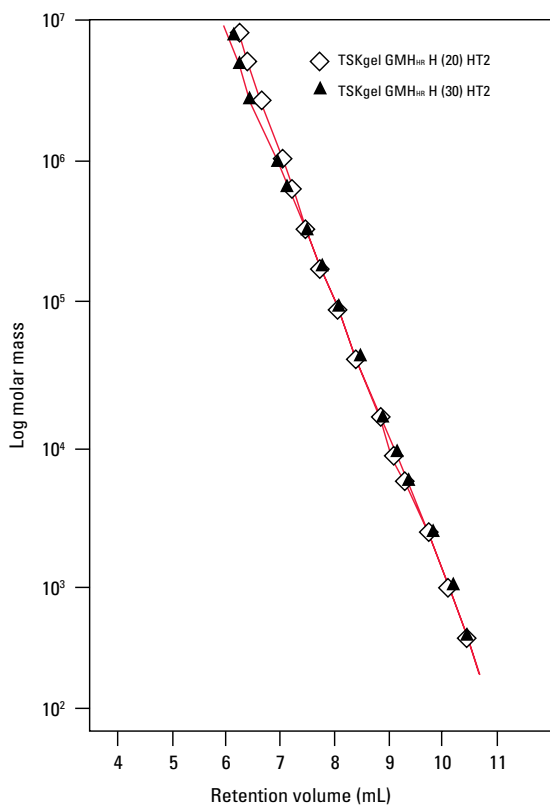
The Heart of Performance – Columns

The TSKgel column series also includes four columns for the analysis of polymers at ultra-high temperatures (up to 220 °C). The TSKgel HT2 columns are specifically designed for the analysis of organic-soluble polymers at extremely elevated temperatures. The attributes of the TSKgel HT2 column series are listed in [Table 7](#).

Table 7: Properties and separation ranges for TSKgel HT2 columns

TSKgel column	Particle size	Pore size	Exclusion limit	Max. temp.
GMH _{HR} -H (20) HT2	20 µm	mixed pore sizes	4.0 × 10 ⁸ Da	220 °C
GMH _{HR} -H (30) HT2	30 µm	mixed pore sizes	4.0 × 10 ⁸ Da	220 °C
GMH _{HR} -H (S) HT2	13 µm	mixed pore sizes	4.0 × 10 ⁸ Da	220 °C
G2000H _{HR} (20) HT2	20 µm	2 nm	1.0 × 10 ⁴ Da	220 °C

Figure 16: Polystyrene calibration curves for TSKgel HT2 columns



Columns: **TSKgel GMH_{HR}-H (20) HT2, 20 µm, 7.8 mm ID × 30 cm**
TSKgel GMH_{HR}-H (30) HT2, 30 µm, 7.8 mm ID × 30 cm
 Mobile phase: ODCB with 0.05% BHT
 Flow rate: 1.0 mL/min
 Detector: RI (EcoSEC High Temperature GPC System)
 Temperature: 135 °C
 Sample: polystyrene standards