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Size-characterization of natural and synthetic polyisoprenes by Taylor dispersion analysis

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ABSTRACT: Non-aqueous Taylor dispersion analysis (TDA) was used for the size-characterization of natural and synthetic polyisoprenes (4×10³ - 2×10⁶ g/mol molar mass). Not only the weight-average hydrodynamic radius ($R_h$), but also the probability distribution of the hydrodynamic radius, were both derived from the Taylorgrams by a simple integration of the elution profile and by a more sophisticated constrained regularized linear inversion of the Taylorgram, respectively. Results in terms of size characterization (hydrodynamic radii between 2 and 100 nm) were compared to size exclusion chromatography coupled to a refractive index-based mass detector. Multimodal size distributions were resolved by TDA for industrial and natural polyisoprenes, with the advantage over the chromatographic technique that, in TDA, there is no abnormal elution of microaggregates (hydrodynamic radii ~ 40-50 nm). Considering the importance and the difficulty of characterizing polyisoprene microaggregates, TDA appears as a promising and simple technique for the characterization of synthetic and natural rubber.

Key Words: polyisoprene; natural rubber; Taylor dispersion analysis; hydrodynamic radius; diffusion coefficient; size distribution.

Title Running Head. Taylor dispersion analysis of synthetic and natural rubbers
1 Introduction

Natural rubber (NR) is a biopolymer produced from latex of *Hevea brasiliensis* trees. NR is used to prepare many goods, the main one being tires that absorb nearly 70 % of the 12 million tons annually worldwide. As an agro-material produced by a tree, NR exhibits rather variable properties for many reasons (genotype of the tree [1-2], season [3], maturation of coagula [4-6], etc.). Thus, as for other polymeric materials used in industrial applications, it is important to characterize the NR macromolecular structure as a prelude to predicting NR end-use properties [7-8].

Natural rubber has a very specific associative structure. When dispersed in good solvents for polyisoprene, NR presents a soluble fraction and an insoluble fraction, often referred to as “macrogel” [9-10] or “gel phase” [11-12]. The composition of the soluble part is also specific as it contains two main entities: (i) polyisoprene linear macromolecules of different lengths, with a random coil structure, and (ii) very compact microaggregates or microgels (sphere-like structure) [13-15]. Links between non-isoprene compounds (lipids, proteins, and/or mineral elements) and the polyisoprene chains are thought to be the main reason for the macrogel formation in NR [12, 16-17]. Although an international standard (ISO 16564) exists to characterize the NR macromolecular structure by size exclusion chromatography (SEC) or gel permeation chromatography (GPC), the development of new analytical techniques is still required to improve the characterization of NR using orthogonal or complementary methods.

This paper describes a novel method to characterize the NR macromolecular structure: Taylor dispersion analysis in organic solvent. Taylor Dispersion Analysis (TDA) is an absolute sizing method requiring no calibration and allowing the determination of the diffusion coefficient (or, equivalently, the hydrodynamic radius) of solutes of any size between one angstrom and a few hundred nanometers. TDA is based on the dispersion of a sample band in an open tubular column under Poiseuille flow. The combination of the dispersive parabolic velocity profile with the molecular diffusion of the solute leads to a specific dispersion, called Taylor dispersion. Started with the pioneering work of Taylor on macroscopic tubing [18], TDA is now implemented in narrow capillaries (typically 50 µm internal diameter), leading to much faster analysis (a few min) and requiring only tiny sample amounts (a few nL) [19-22]. In its modern miniaturized version, TDA can be very easily implemented on commercial capillary electrophoresis apparatus. Alternatively, fast TDA can be performed on HPLC equipment at a high linear velocity (~30 cm s\(^{-1}\)) in a coiled open capillary tube of 250 µm i.d. × 30 m typical dimensions, as recently introduced by the Holyst group [23]. In the latter
configuration, the genuine conditions for the applicability of TDA are not fulfilled [24], however, an empirical correction can be used to get the correct diffusion coefficient [23].

TDA was applied to the characterization of proteins [25-27], macromolecules [28-30], drug delivery systems [31, 32], nanoparticles [21, 33, 34], polyelectrolyte complexes and polyplexes [35], pharmaceutical compounds [36, 37], and micelles or microemulsions [38, 39]. It can also be used for the study of molecular interactions [40, 41]. The application of TDA is not limited to water-soluble solutes, as demonstrated by the size characterization of polystyrenes in THF [28], asphaltene/bitumen in hexane or THF/acetonitrile mixtures [42, 43], and drugs in pharmaceutical solvents [36]. Basically, TDA yields the weight average hydrodynamic radius when using a mass sensitive detector, such as a UV detector, for a polymer absorbing via the repeating unit [44]. This information is obtained by simple integration of the elution temporal profile (also called Taylorgram). However, we recently demonstrated that more sophisticated data analyses allow one to determine the full hydrodynamic radius distribution, thus providing valuable information on the sample polydispersity [45, 46].

In this work, we demonstrate that TDA can be used to get, within a few minutes, the hydrodynamic radius distributions of commercial natural and synthetic polyisoprenes. The data obtained by TDA are compared to those derived from size exclusion chromatography (SEC) coupled to multi-angle static light scattering (MALS).

2 Experimental

2.1 Materials

The two NR samples used for this study were all TSR (Technically Specified Rubber). Sample NR1 was industrial TSR10 grade prepared by natural coagulation of latex followed by several days of coagulum maturation before processing. This sample was treated with neutral hydroxylamine sulfate (NHS) to stabilize its properties during storage, yielding a TSR10CV grade (CV stands for constant viscosity), a special grade not prone to storage hardening. Sample NR2 was prepared by acid coagulation (formic acid) of fresh field latex. Neutral hydroxylamine sulfate (NHS) was added to the latex prior to coagulation to inhibit storage hardening of rubber [47]. Sample NR2 was TSR5CV grade. Three synthetic industrial cis-1,4-polyisoprenes (PI) were used directly as received: IR2200 (NatSyn 2200, Goodyear chemical), IR307 and IR309 (Kraton polymers). Standard linear polyisoprene (PI) of 4.5, 22, 111, 307, 361, 608 and 766 × 10^3 g/mol weight-average molar masses were obtained from Polymer Standard Service (PSS, Germany) and were used as received.
2.2 Determination of the number- and weight-average molar mass by size exclusion chromatography coupled to multi-angle light scattering (SEC-MALS)

For all samples, the macromolecular structure was characterized by SEC-MALS, except for standard polyisoprene, for which the number average, $M_n$, and mass-average, $M_w$, molar mass were given by the supplier. The samples (30 ± 5 mg), from NR pellets, were dissolved in tetrahydrofuran (THF, 30 mL, HPLC grade, VWR France) stabilized with 2,6-di-tert-butyl-4-methylphenol (BHT). The solutions were stored for 7 days in the dark at 30°C (during storage, the samples were stirred for 1 h/day), they were then filtered (Acrodisc 1 µm, glass fiber, Pall France) and finally injected into the SEC-MALS apparatus. For each sample, three solutions were independently prepared and measured. The SEC equipment comprises an online degasser (EliteTM, Alltech), a Waters 515 pump, a refractive index detector (Waters 2410) and a multi-angle light scattering detector (Dawn DSP, Wyatt Technology). Two Waters HMW6E columns (Styragerel HMW columns packed 20 µm particles, 300 mm x 7.8 mm I.D.), maintained at 45°C, and a guard column (same stationary phase as the two separation columns) were used. The mobile phase was stabilized THF injected at a flow rate of 0.65 mL.min$^{-1}$; the injected volume was 150 µL. The MALS detectors at all 18 angles were calibrated using a THF solution of a polystyrene standard with low polydispersity ($M_w = 30.3$ kg.mol$^{-1}$, Wyatt technology). The same solution was used to determine the interconnection volume between the two detectors (0.235 mL). The basic theory of determining the weight-average molar mass and radius of gyration for a dilute solution of a macromolecule is well known and described in numerous papers in the literature [48, 49]. The weight average molar masses and radius of gyration at each slice of the chromatogram were calculated using the Berry method for extrapolation, as implemented in ASTRA software (version 5.3.1, Wyatt technology). The order of polynomial fit for the Berry analysis was two. Twelve angles, from 38.8° to 138.8°, were used for the analysis. The differential refractive index increment at $\lambda = 633$ nm was $dn/dc = 0.130$ mL.g$^{-1}$ [50].

2.3 Determination of the weight-average hydrodynamic radius $R_h$ and of the $R_h$ distribution by Taylor dispersion analysis

TDA experiments were performed on a PACE MDQ Beckman Coulter (Fullerton, CA) apparatus. Capillaries were prepared from bare silica tubing purchased from Composite Metal Services (Worcester, United Kingdom). The capillary dimensions were 40 cm (30 cm to the detector) × 50 µm I.D. New capillaries were conditioned with the following flushes: 1 M
NaOH for 30 min and water for 10 min. Before sample injection, the capillary was filled with cyclohexane (8.62 × 10^{-4} Pa.s viscosity). Between two TDA analyses, the capillary was flushed with cyclohexane (30 psi for 2 min). A mobilization pressure of 1 psi (~71 mbar) was applied with cyclohexane vials at both ends of the capillary, resulting in a flow rate of 184 nL.min^{-1}. Samples were dissolved in cyclohexane at a concentration of 1 g/L. Sample injection (2.8 nL) was performed hydrodynamically on the inlet side of the capillary (0.3 psi for 3 s; ~1% of the capillary volume). Solutes were monitored by UV absorbance at a wavelength of 200 nm. The elution time was systematically corrected for the delay in the application of the pressure (pressure ramp time, 15 s) by subtracting 7.5 s (half-time of the pressure ramp) to the recorded elution time. The temperature of the capillary cartridge was set at 25 °C.

Two different methods were used to determine the weight-average hydrodynamic radius by TDA. The first method (method 1) is based on the integration of the Taylorgram [22], and the second method (method 2) is based on a deconvolution of the Taylorgram using a recently published Constrained Regularized Linear Inversion (CRLI) method [45].

In method 1, the average diffusion coefficient $D$ is obtained by integration of the Taylorgram (or temporal elution profile) in order to calculate the temporal variance of the Taylorgram, $\sigma^2$. The integration of the elution profile was performed using the discrete form of the following equation [22]:

$$\sigma^2 = \frac{\sum_{i=m}^{n} S(0)(t - t_0)^2 dt}{\sum_{i=m}^{n} S(t) dt} = \frac{\sum_{i=m}^{n} S_i(0)(t_{i+1} - t_0)^2(t_{i+1} - t_i)}{\sum_{i=m}^{n} S_i(t_{i+1} - t_i)}$$

(1)

where $S(t)$ is the detector response, $t_i$ is the elution time for a given point $i$ of the Taylorgram and $t_0$ is the average elution time. $n$ and $m$ are the starting and ending points that are considered for the integration of the Taylorgram.

The integration of the Taylorgram was only performed on the left part of the elution profile (the end point was $t_0$), before and after subtracting the Gaussian contribution due to the small molecules (see results and discussion section). Corrections of the temporal variance and average detection time due to the injected volume were also taken into account, as described elsewhere [29], but these corrections are negligible when the injected volume is ≤ 1% of the capillary volume to the detection point [22], as is the case here. Finally, the so-called Taylor average diffusion coefficient $\langle D \rangle_{TDA}$ is obtained from:

$$\langle D \rangle_{TDA} = \frac{R_c^2 t_0}{24 \sigma^2}$$

(2)

where $R_c$ is the capillary radius.
Equation (2) is valid as far as \( \frac{\langle D \rangle_{TDA} R_c^3}{R_c^2} \geq 1.25 \) \(^{[18, 24]}\) to ensure that the analysis time is larger than the characteristic diffusion time of the solutes over the capillary cross section. Equation (2) is also only valid if the axial diffusion can be neglected compared to the Taylor dispersion contribution. This latter condition is fulfilled if \( \frac{R_c u}{\langle D \rangle_{TDA}} \geq 40 \), where \( u \) is the linear velocity of the mobile phase \(^{[18, 24]}\). It is worth noting that the Taylor average diffusion coefficient obtained by integration of the Taylorgram (method 1) is a weight harmonic average diffusion coefficient defined by:

\[
\langle D \rangle_{TDA} = \left[ D^{-1} \right]^{-1} = \left[ \int_0^\infty P_D(D) D^{-1} dD \right]^{-1}
\]

(3)

where \( P_D(D) \) is the (mass-weighted) probability distribution function (PDF) of the diffusion coefficient. The TDA weight-average hydrodynamic radius \(<R_h>_{TDA}\) is finally determined using the Stokes-Einstein relationship:

\[
<R_h>_{TDA} = \frac{k_B T}{6\pi \eta \langle D \rangle_{TDA}}
\]

(4)

where \( k_B \) is the Boltzmann constant, \( T \) the temperature (K) and \( \eta \) the viscosity of the carrier liquid (Pa.s).

Method 2 is based on a recently developed Constrained Regularized Linear Inversion (CRLI) approach \(^{[45]}\). This method allows the PDF of the hydrodynamic radius to be extracted from the experimental Taylorgrams. In brief, this method consists of introducing additional constraints in the usual least-squares fitting of the Taylorgram (“regularization”), in order to reject non-physical solutions, finally converging to the desired PDF of the hydrodynamic radius, \( P_R(R_h) \). Typically, the constraints penalize those solutions for which \( \partial^2 P_R / \partial R_h^2 \) is, on average, too large, i.e. those PDFs that exhibit too detailed a structure, see Ref. \(^{[45]}\) for more details.

The CRLI algorithm inverts the following relationship relating the Taylorgram to \( P_R(R_h) \) \(^{[45]}\):

\[
S(t) = \frac{k_R m_{tot}}{Q} \sqrt{\frac{\lambda_R}{\pi}} \int_0^\infty \frac{P_R(R_h)}{\sqrt{R_h}} \exp \left[ -\frac{\lambda_R (t-t_0)^2}{R_h} \right] dR_h
\]

(5)

where \( k_R \) is the detector response factor, \( m_{tot} \) the total sample mass injected, \( Q \) the carrier liquid flow-rate and with

\[
\lambda_R = \frac{2k_B T}{\pi R_c^2 t_0 \eta}
\]

(6)
After normalization, eq. (5) reduces to:

$$s(t) = \frac{S(t)}{S(t_0)} = c_R \int_0^\infty \frac{P_R(R_h)}{\sqrt{R_h}} \exp \left( -\frac{2 \lambda R (t-t_0)^2}{R_h} \right) dR_h$$  \hspace{1cm} (7)$$

with

$$c_R = \left[ \int_0^\infty \frac{P_R(R_h)}{\sqrt{R_h}} dR_h \right]^{-1} = \left[ R_h^{-1/2} \right]^{-1}$$  \hspace{1cm} (8)$$

The PDFs of $R_h$ reported in the following are obtained by averaging individual distributions measured in 2 to 5 independent repetitions. For each sample, a mass-average $R_h$ value was calculated from $P_R(R_h)$, via:

$$\langle R_h \rangle_{CRL} = \frac{\int_0^\infty R_h \times P_R(R_h) dR_h}{\int_0^\infty P_R(R_h) dR_h}$$  \hspace{1cm} (9)$$

3 Results and discussion

Our main objective is to compare the average radii and the $R_h$ PDFs obtained by SEC and by TDA on different PI standards, industrial PI and natural rubbers (NR).

3.1 Determination of the average hydrodynamic radius by TDA

Examples of raw Taylorgrams obtained for the polyisoprene (PI) standards are displayed in Figure 1A. Each Taylorgram has been normalized by the maximum absorbance at $t=t_0$. A quite good baseline return and peak symmetry is seen in Figure 1A. It is clearly observed that the higher the molar mass, the broader the Taylorgram, due to the higher Taylor dispersion for the species with lower diffusion coefficient. Nevertheless, some PI standards contain a tiny amount of small molecules (possibly monomers, residual solvent) absorbing at the detection wavelength (200 nm). This small molecule contribution appears as a thin peak, most notable for the 361 and 766 × 10^3 g/mol PI standards (blue and green lines in Figure 1A). The contribution due to the small molecules should not be taken into account in the data analysis. Accordingly, we systematically corrected for it by subtracting the corresponding Gaussian peak from the raw data, leading to the Taylorgrams presented in...
The corresponding weight-average hydrodynamic radii obtained by integration of the Taylorgrams presented in Figure 1B (method 1, \( \langle R_h \rangle_{TDA} \)) are gathered in Table 1. For the sake of comparison, the \( R_h \) values obtained without subtraction of the small molecule contribution are also given in Table 1. The impact of the small molecule contribution on the average \( R_h \) value is less than 3%.

The same data processing was applied to the industrial PI samples (IR2200, IR307 and IR309) and the natural rubber samples (NR1 and NR2). The raw Taylorgrams and the corrected ones are displayed in Figures 2A and 2B for industrial PI, and Figures 3A and 3B for NR, respectively. It should be noticed that the symmetry of the Taylorgrams was not perfect, especially in the case of NR. This may be due to some polymer adsorption onto the capillary surface. This is why the Taylorgrams were analyzed on the left part, where adsorption has a much lower impact. The contribution of the small molecules to the average \( R_h \) value is less than 6% (see Table 1). The correlation between the hydrodynamic radius (method 1, \( \langle R_h \rangle_{TDA} \)) and the PI weight average molar mass \( (M_w, \text{SEC}) \) is given in Figure 4.

For a random coil in a good solvent, the theoretical value of the exponent \( b \) of the power law \( R_h = aM_w^b \) is predicted to be 0.588 [52]. This value was confirmed experimentally using light scattering, for PI in cyclohexane (Fetters et al., \( b=0.592 \) [53]) and in tetrahydrofuran (Kim et al., \( b = 0.606 \) [14]). In our experiments, we find an exponent \( b=0.54+/0.06 \) (confidence interval at 95%) for the seven PI standards, slightly lower than the expected theoretical value. Industrial PI grades and NR samples are close to the scaling law obtained for the PI standards (see Figure 4).

3.2 Hydrodynamic radius distributions obtained by TDA and comparison with SEC

The elution profile in SEC gives access to the size distribution of the polymers, since the separation parameter in SEC is the hydrodynamic volume of the solute. For the SEC
measurements, THF was used as the eluent. THF gives a higher signal for the LS and refractive index analysis, thanks to a higher dn/dc compared to cyclohexane. For the TDA analysis, by contrast, cyclohexane is the best choice, since THF is incompatible with UV detection and with the TDA setup (THF dissolves the MDQ Beckman Coulter plastic interface in contact with the sample and eluent vials). In spite of the need to use different solvents, the comparison between the TDA and SEC-MALS results for the same PI and rubber samples sheds light on the strengths and limitations of each method.

In SEC, the logarithm of the hydrodynamic radius is a decreasing linear function of the elution volume. In the elution profile, the refractive index (RI) detector yields a signal proportional to the polymer mass concentration. The MALS detector combined with the RI detector allows the polymer molar mass to be measured. All the 3 industrial PI analyzed by SEC-MALS exhibited a main peak eluting at about 16-17 min, followed by one or two other modes eluting between 18-22 min, which were more or less separated depending on the sample. IR309 appeared as a tri-modal elution profile (see Figure 5A), while the two other samples (IR307 and IR2200) presented unresolved shoulders. Figure 5B shows hydrodynamic radius ($R_h$) mass-distributions obtained by SEC assuming that $R_g/R_h = 1.78$, as expected for monodisperse random coils in good solvent [52]. Indeed, SEC coupled to a MALS detector allows determining, for each slice of the chromatogram, both the molar mass ($M_i$) and the radius of gyration ($R_g$). The $R_h$ were calculated from the $R_g$ for each slice of the chromatogram using $R_g/R_h = 1.78$ relationship. In the SEC analysis, all samples display rather broad distributions with an unresolved shoulder for IR2200 and IR307, and three partially resolved modes for IR309. The $R_h$ corresponding to the peak maximum for the mass-distribution of $R_h$ determined from the SEC data (Figure 5B) were 32 nm for IR2200, 45 nm for IR309 and 47 nm for IR307.

Figure 5C displays the hydrodynamic radius distributions obtained by TDA after data processing of the Taylorgrams from Figure 2A according to method 2 (CRLI, see section 2.3). Size distributions obtained by TDA display two well-defined main modes, and a less important (minor) third mode (see inset of Figure 5C), corresponding to low molar mass polymers (oligomers) or monomers with $R_h$ below 3 nm. These oligomers/monomers appear as a sharp peak in the Taylorgrams presented in Figure 2A. All IR samples display a main mode at $R_h$ ranking between 30 and 50 nm (peak apex at 34.8 nm for IR 2200, 41.4 nm for IR 309 and 42.3 for IR 307). These $R_h$ values are in reasonably good agreement with SEC analysis, given that the solvents are different and taking into account the uncertainty in the actual value of $R_g/R_h$ required to calculate the $R_h$ PDF from the SEC data. A second mode in the $R_h$ distribution was observed by TDA, corresponding to sizes between 5 and 11 nm (peak apex at 10.6 nm for IR 2200, 9.6 for IR 309 and at 5.6 nm for IR 307). This peak is not
resolved in the PDF calculated from the SEC data, except perhaps for the IR 309 sample, which exhibits a pronounced shoulder in the 10-20 nm range. Overall, TDA showed was better resolution for the size characterization of the IR samples compared to SEC.

<Figure 5>

For natural rubber (NR) samples, the TDA analysis again gave $R_h$ profiles with better size resolution than those retrieved from the SEC elution profiles (Figure 6). The SEC data exhibited elution profiles rather unimodal, with a more or less pronounced shoulder. Unlike the PI industrial samples, for the NR samples, it is not possible to assume that each slice of the chromatogram is only composed of monodisperse random coils, because of the co-elution of microaggregates and random coil macromolecules [13]. For this reason, it was not possible to calculate the $R_h$ PDF. By contrast, in TDA, $R_h$ is directly determined, with no need of making any assumptions on the nature of the sample (e.g. on the presence of aggregates). As displayed in Figure 6B, the NR samples had trimodal $R_h$ profiles with small oligomers ($R_h$ below 2-3 nm) and two larger polymer modes at 7–10 nm and 20–60 nm, respectively. Remarkably, the TDA analysis reveals unambiguous differences in the position and the width of the peaks for the two samples, not captured by SEC. In particular, the peak for the largest sizes shifts from 37 nm for NR1 to 40 nm for NR2. As explained above, it was not possible to determine the $R_h$ PDF from the SEC data. However, we note that the ranking in the SEC elution curves is consistent with the TDA data, since NR2 is eluted before NR1, indicative of a larger size.

From the $R_h$ distribution obtained by method 2, it is also possible to determine a weight-average $R_h$ by integration on the whole distribution (see Eq. 9). We exclude the contribution of the small molecules, $R_h < 3$ nm, consistent with the subtraction performed in method 1. The results are presented in Table 1: we find a difference of less than 3% between the two methods. This demonstrates the excellent agreement between the two different ways to process the Taylorograms, thereby confirming the robustness of the methods of analysis presented here.

<Figure 6>

4 Conclusions

We have demonstrated that TDA can be used in non-aqueous media (cyclohexane in the present study) for the analysis of PI in industrial and natural rubbers. Because TDA is implemented in open capillary tubes, it is much less influenced by undesirable interactions as compared to SEC, where microaggregates from NR samples (hydrodynamic radii ~40-50 nm)
may result in abnormal elution curves at large elution times. By leveraging on the constrained regularized linear inversion (CRLI) method, not only does TDA give access to the weight average hydrodynamic radius, it also provides the full hydrodynamic radius distribution. While under some conditions SEC data can yield the PDF of $R_h$, the present work shows that TDA has a broader applicability and a superior ability to resolve the various modes contributing to the sample size distribution in a large range of hydrodynamic radii from ~1 to 100 nm. Given the strengths of the method and the advantages of modern TDA in its miniaturized version (low sample volume ~nL, low solvent consumption, fast analysis (a few min), absolute determination, simple instrumentation and straightforward methodology), there is no doubt that TDA holds great potential for quality control and production monitoring of PI in the rubber industry. Possible future work could be to investigate new capillary coatings and/or capillary i.d. to improve the peak symmetry, especially in the case of NR analysis.

References


**Figures and Captions**

**Figure 1.** Taylorgrams obtained for PI standards in cyclohexane before (A) and after substraction of the contribution due to small molecules (B). Experimental conditions: fused silica capillary 40 cm (30 cm to the detector) x 50 µm i.d. Mobile phase: cyclohexane. Hydrodynamic injection: 0.3 psi, 3 s. Mobilization pressure: 1.0 psi. Temperature: 25 °C. Samples: PI standards 4.5, 111, 361, and 766 × 10³ g/mol at 1 g/L in cyclohexane. Detection at 200 nm. The normalized taylorgram is presented (S(t)/S(t₀)).
Figure 2: Taylorgrams obtained for industrial PI in cyclohexane before (A) and after substraction of the contribution due to small molecules (B). Same conditions as in Figure 1.
Figure 3. Taylorgrams obtained for natural rubbers (NR) in cyclohexane before (A) and after subtraction of the contribution due to small molecules (B). Same conditions as in Figure 1.
**Figure 4:** Log-log plot of the TDA hydrodynamic radius (method 1, after subtraction of the small molecule contribution) versus the molar mass for the PI standards, PI industrial and for NR. The dotted line is a power law fit for the PI standards. See Table 1 for the numerical values.
A. DRI Signal vs. Elution Time (min)

B. $R_h P_R(R_h)$ for Industrial PI

C. $R_n P_R(R_n)$ for Industrial PI
**Figure 5**: Elution profiles obtained by SEC (RI signal, A) and $R_h$ distributions obtained by SEC-MALS (B) or TDA using *method 2* (C), for the 3 industrial PI samples. Experimental conditions: SEC in tetrahydrofuran, two Waters HMW6E columns (20 µm, 300 mm x 7.8 mm I.D.) maintained at 45°C, flow rate of 0.65 mL.min⁻¹; TDA conditions as in Figure 2. The insert in Figure 2C is an expanded view. $R_h$ distributions obtained by SEC-MALS are based on the assumption that $R_d/R_h = 1.78$. 
Figure 6: Elution profiles obtained by SEC (RI signal, A) and $R_h$ distributions obtained by TDA using method 2 (B) for two NR samples. Experimental conditions: SEC, see Figure 5; TDA, as in Figure 3. Insert in Figure 6B is an expanded view.
**Table 1:** Number- and weight-average molar masses ($M_n$, $M_w$), hydrodynamic radius ($R_h$ +/− one standard deviation) for natural and synthetic PI. $M_n$, $M_w$ are determined by SEC-MALS, except for the PI standards*. $R_h$ is determined by TDA, using both method.

<table>
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<th>Type</th>
<th>$M_n$ (kg.mol$^{-1}$) (1)</th>
<th>$M_w$ (kg.mol$^{-1}$) (1)</th>
<th>$\langle R_{h/\text{TDA}} \rangle$ (nm)$^b$ Method 1</th>
<th>$\langle R_{h/\text{CRLI}} \rangle$ (nm)$^b$ Method 2</th>
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<td></td>
<td></td>
<td></td>
<td>with subtraction$^c$</td>
<td>without subtraction$^d$</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>Integration for $R_h$ &gt; 0$^f$</td>
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<td>PI Standard</td>
<td>766</td>
<td>766</td>
<td>27.9 ± 0.7</td>
<td>27.3 ± 0.6</td>
</tr>
<tr>
<td>PI1040</td>
<td>PI Standard</td>
<td>1040</td>
<td>1040</td>
<td>28.9 ± 0.05</td>
<td>27.4 ± 0.2</td>
</tr>
<tr>
<td>IR307</td>
<td>PI Industrial</td>
<td>1079</td>
<td>1986</td>
<td>40.1 ± 0.5</td>
<td>38.9 ± 0.5</td>
</tr>
<tr>
<td>IR309</td>
<td>PI Industrial</td>
<td>652</td>
<td>1935</td>
<td>35.0 ± 0.1</td>
<td>33.6 ± 0.2</td>
</tr>
<tr>
<td>IR2200</td>
<td>PI Industrial</td>
<td>584</td>
<td>1271</td>
<td>32.4 ± 0.3</td>
<td>31.1 ± 0.3</td>
</tr>
<tr>
<td>NR1</td>
<td>NR Industrial</td>
<td>595</td>
<td>902</td>
<td>31.7 ± 1.5</td>
<td>29.8 ± 1.4</td>
</tr>
<tr>
<td>NR2</td>
<td>NR Industrial</td>
<td>711</td>
<td>1282</td>
<td>36.2 ± 0.4</td>
<td>34.3 ± 0.5</td>
</tr>
</tbody>
</table>

* For standard polyisoprene, all $M_n$ and $M_w$ were given by the supplier.

* Three repetitions in average

$^c$ Average $R_h$ obtained by integration of the taylorgram after subtraction of the small molecule contribution (method 1 applied to Figures 1B, 2B, 3B).

$^d$ Average $R_h$ obtained by integration from the whole taylorgrams including the small molecule contribution (method 1 applied to Figures 1A, 2A, 3A).

$^e$ Average $R_h$ obtained by method 2 (CRLI, eq. (9)) by integration of the $P(R)$ distribution for $R_h$ higher than 3 nm.

$^f$ Average $R_h$ obtained by method 2 (CRLI, eq. (9)) by integration of the whole $P(R)$ distribution.
Highlights

- Sizing natural or synthetic polyisoprenes by miniaturized Taylor dispersion analysis
- Full hydrodynamic radius distribution of polyisoprenes from Taylor dispersion analysis
- Taylor dispersion analysis has a better size-resolution compared to SEC
- No abnormal elution of the microaggregates in Taylor dispersion analysis
- Short analysis time (a few min) and low sample and solvent consumption