HPLC determination of perfluorinated carboxylic acids with fluorescence detection

Ewa Poboży · Edyta Król · Lena Wójcik · Mariusz Wachowicz · Marek Trojanowicz

Received: 2 September 2010 / Accepted: 18 November 2010 / Published online: 12 December 2010
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Abstract Perfluorinated carboxylic acids (PFCAs) represent an important group of persistent perfluorinated organic compounds commonly determined in environmental and biological samples. A reversed-phase HPLC method was developed based on derivatization of the PFCAs with the commercially available fluorescent reagent 3-bromoacetyl coumarin. The method was optimized and this resulted in the efficient separation of PFCAs containing from 3 to 12 carbon atoms in molecule in 25 min run. To improve sensitivity, the preconcentration step has been optimized using Oasis-WAX and C18 sorbents for SPE. A 100-fold preconcentration is achieved by solid-phase extraction with the sorbent C18 Sep-PAK to result in limits of detection in the range from 43 to 75 ppt for the analytes examined, and in the application of the method of water analysis.

Keywords Perfluorinated carboxylic acids · HPLC · Fluorescence detection

Introduction

Perfluorinated compounds (PFCs) are a family of fluorine-containing compounds with a fully fluorinated carbon chains. Due to their unique properties such as resistance to high temperature, capacity to form water, stain, oil and grease repellent surface, they have found many applications [1]. In spite of introduction of PFCs in 1950s, the identification and quantification have been possible since late 1990-ties when analytical methods of sufficient sensitivity were available. The widespread occurrence of certain perfluorinated compounds in the environment, in animal as well as in human species, has attracted great attention in last years. Significant levels of PFCs were detected in various environmental samples and they are considered as new class of persistent organic pollutants [2].

Among different PFCs, the increasing of occurrence of perfluorinated carboxylic acids (PFCAs) in last decade has been reported in the environment in many countries. They are released into the environment by manufacturing processes, during production of fluoropolymers, and as residuals from products, or by degradation processes of other fluorinated compounds.

Global widespread of PFCAs is observed and they have been identified as well in environmental samples [3], as in animal tissues [4], and human blood [5]. The sources, fate and transport of PFCAs, with a specific focus on perfluorooctanoic acid (PFOA) in the environment (ocean water, freshwater, sediments and air) have been recently thoroughly reviewed [6]. Also extensive studies of PFOA toxicity for living organisms and human exposure sources have been carried out [7].

Determination of PFCAs in low trace level is an analytical challenge. Reliable methods of extraction, separation and identification in complex matrixes are necessary. The difficulty of the determination of PFCAs is very low concentration in the samples (ng-pg L$^{-1}$ or ng-pg g$^{-1}$) and their weak absorption in UV. A few analytical methods have been developed to determine PFCAs and most commonly employed are chromatographic ones [8].

For determination of PFCAs without derivatization, the liquid chromatography with the MS/MS detection and
Electrospary ionization is commonly used for analysis of environmental [9–11] and biological samples [12–14]. Ion chromatography with conductometric detection for separation of PFCAs having C3–8 has been also presented [15], with LODs in range 0.12–0.66 mg L\(^{-1}\), while with SPE preconcentration the limit of detection of 50 μg L\(^{-1}\) was reported.

Different approaches using solid-phase extraction methodology for preconcentration of PFCAs have been developed. For determination with LC-MS, the methodology of Moody et al. [16] employing C18 sorbents has been mainly employed by various authors, which is limited to long chain acids. Recently, also application of polymeric sorbents Oasis HLB and Oasis WAX mixed-mode weak anion-exchange reversed phase for PFCAs preconcentration has been presented [9, 17]. The acceptable recoveries of PFOA using both C18 (124%) and Oasis HLB (147%) were obtained but the lower background levels were observed for Oasis HLB cartridges [9]. Recovery of PFOA from water samples using Oasis HLB ranged from 33 to 98%, and it was above 50% for 80% of the samples analyzed, as reported by MacLachlan et al. [30].

For HPLC analysis of PFCAs with the fluorimetric detection Ohya et al. [18] reported derivatization with laboratory synthesized 3-bromoacetyl-7-methoxycoumarin (BrAMC), and conditions of carrying out reaction and separation were optimized for analysis of tissues. The developed method was applied for determination of PFCAs in rats liver and recoveries >90% were obtained for 50 mg homogenate spiked with 1 nM PFCAs. This method was also employed to study transport of PFCAs in organisms and their elimination [19], and to study toxicity of determined analytes [20].

The aim of this work was to develop the HPLC method for determination of perfluorinated carboxylic acids with fluorimetric detection based on the use of commercially available fluorophores and to develop the procedure for analysis of natural water samples. For investigation, PFCAs with 2 to 12 carbon atoms were chosen, which occur most commonly in environmental samples. The studies were focused on investigation of two coumarin fluorophores, namely 3-bromoacetoylcoumarin (3-BrAC) and 4-bromomethyl-7-methoxycoumarin (BrMMC), which are commercially available, and the latter one was already employed for derivatization of carboxylic acids for HPLC determination with fluorimetric detection of fatty acids [21, 22], okadaic acid and other shellfish poisoning toxins [23], and monofluoroacetate [24].

**Experimental**

**Materials and chemicals**

The following perfluorinated carboxylic acids (PFCAs) were purchased from Aldrich (Milwaukee, USA, [www.sigmaaldrich.com](http://www.sigmaaldrich.com)): perfluoroacetic acid (abbreviated as C2), perfluoropropionic acid (C3), perfluorobutyric acid (C4), perfluoropentanoic acid (C5), perfluorohexanoic acid (C6), perfluoroheptanoic acid (C7), perfluorooctanoic acid (C8 or PFOA), perfluorononanoic acid (C9), perfluorodecanoic acid (C10), perfluoroundecanoic acid (C11), and perfluorododecanoic acid (C12). Purity of all reagents was better than 95%. Methanol and acetonitrile (ACN), both of chromatographic purity, were obtained from Merck (Darmstadt, Germany, [www.merck.com](http://www.merck.com)). Ammonium solution (25%), acetone and sodium hydroxide were from Chempur (Pickary Slaskie, Poland, [www.chempur.pl](http://www.chempur.pl)). 4-bromo-methyl-7-methoxycoumarin (BrMMC) and 3-bromoacetyl-coumarin (3-BrAC) were purchased from Aldrich (Milwaukee, USA). Mili-Q (Millipore, USA) water was used throughout the all experiments.

Stock standard solutions of PFCAs of concentration 1,000 mg L\(^{-1}\) were prepared by dissolving them in acetonitrile. These solutions were stored in the dark at 4 °C. A series of working standards solutions were prepared by subsequent dilution of stock solutions with acetonitrile to reach concentrations range 0.025–0.5 mg L\(^{-1}\). The standard mixtures for preconcentration studies were obtained by dilution and mixing of stock solutions with water.

Solutions of coumarin derivatives (2.0 or 5.0 g L\(^{-1}\)) were prepared by dissolving appropriate amount of solid reagent in acetone. 3-BrAC dissolves easy in acetone but BrMMC needs long time dissolving in ultrasonic bath. The solutions were stored in the dark at 4 °C.

**Instrumentation**

The HPLC system Merck-Hitachi (Darmstadt, Germany) consisted of a system controller D-7000, an autosampler L-7250 (with sample volume loop 50 μL), a degasser, a gradient pump L-7100 and fluorescence detector L-7485. The detector was set at an excitation wavelength 306 nm and emission wavelength 420 nm for the derivatization with BrAC and at an excitation wavelength 330 nm and emission wavelength 390 nm for the derivatization with BrMMC. The analytical column was Phenomenex (Torrance, USA, [www.phenomenex.com](http://www.phenomenex.com)) Synsersi C18 (4 μm), 150×4.6 mm LD. Gradient elution with flow rate 1 mL min\(^{-1}\) was applied for separation. The gradient was hold at 100% eluent A for 5 min, then operated from 100% eluent A to 100% eluent B for 20 min, where eluent A was 60% ACN and eluent B 100% ACN.

Varian SPE manual system was used for preconcentration. Samples were pumped through cartridges with peristaltic pump model MS-4 REGLO (Ismatec, Switzerland).

During the derivatization step, samples were heated using thermoblock RM-5 Labart (JW Electronic, Poland) equipped with thermostatically controlled heating.
Solid phase extraction procedures

PFCAs were preconcentrated using Oasis-WAX (30, 60 and 120 mg, 30 μm) and Sep-Pak C18 (500 mg) cartridges purchased from Waters (Milford, USA). Procedures with some modifications for Sep-Pak C18 were based on those described previously [16]. The Sep-Pak C18 cartridges were preconditioned with methanol (5 mL) followed by water (2 mL). Then standards solutions of PFCAs in water or natural water sample were passed through the cartridge with flow-rate 1 mL min⁻¹. After 15 min air drying, retained analytes were eluted with methanol (three 2.5 mL fractions). Each fraction was collected separately and evaporated under argon to dryness and next the residue dissolved in 2.5 mL or 1.0 mL of acetonitrile. The derivatization with coumarin derivatives proceeds better in acetonitrile than in methanol, but for elution methanol is preferred, so the evaporation and dissolving in acetonitrile was necessary.

For preconcentration with Oasis WAX cartridges, procedure presented by Tanyasu et al. [18] was applied. The cartridges were preconditioned with methanol containing 0.1 or 5% ammonia (4 mL) followed by methanol (4 mL) and water (4 mL). Then standard solutions of PFCAs in water were passed through the cartridge with flow-rate 1 mL min⁻¹. After 15 min air drying, the target analytes were eluted with methanol containing 0.1 or 5% ammonia (three 4.0 mL fractions). Each fraction was collected separately and evaporated under argon to dryness and next the residue dissolved in 2.5 mL or 1.0 mL of acetonitrile. Such obtained solutions were derivatized with coumarin derivatives.

In recovery study, 25, 250 and 1,000 mL solutions of concentrations 30, 3.0 and 0.3 μg L⁻¹, respectively, were preconcentrated. The final volumes of preconcentrated samples were 2.5, 2.5 and 1.0 mL and the preconcentration factors 10, 100 and 1,000 were achieved.

Derivatization procedure

PFCA standard solutions or extracts in acetonitrile were mixed with solution of coumarin derivatives (1:1, v/v) and heated in thermoblock in temperature 70 °C in closed, dark glass vials. Then after cooling, the obtained mixture was injected into HPLC system.

The reaction is inhibited by water hence using of dry vials and drying of cartridges after preconcentration is indispensable.

Analysis of surface water samples

Samples of natural surface waters were collected from various points in vicinity of chemical plant producing fluorinated polymers in town Tarnów, Southern Poland. As PFCAs may strongly adsorb on the glass surface, water samples were collected in polyethylene containers washed with methanol [26]. Samples were stored without preservation at 4 °C. Care was taken to avoid samples contact with PTFE.

Three 100 mL water samples were preconcentrated on Sep-Pak C18 columns with flow rate 1 mL min⁻¹. Analytes were eluted with 5 mL methanol and evaporated in argon atmosphere. Solid residues was dissolved in 1 mL ACN, and 0.7 mL solution was used for 1 h derivatization at 70 °C with 2 gL⁻¹ solution of 3-BrAC.
Results and discussion

Optimization of derivatization procedure

Among numerous reagents used in derivatization for application of fluorescence detection in HPLC, coumarin derivatives belong to the most sensitive fluorophores. As it was mentioned above they have been already widely employed for derivatization of carboxylic acids, which are non or slightly UV adsorbing hence fluorescence derivatization is often used to their determination. So far, one such attempt was published in the literature on application for HPLC of perfluorinated carboxylic acids, where laboratory synthesized 3-bromoacetyl-7-methoxycoumarin was employed, and limit of detection was given as 50 pM/50 mg of analyzed rat liver [18].

The yield of derivatization with coumarin derivatives, limit of detection and efficiency of separation of derivatized

Analysis of commercial preparations for shoe impregnation

Two commercial preparation were analyzed, aerosol from Kiwi, and liquid preparation from Boreal (Spain). Sample of aerosol (282.2 mg) was weighted in hermetically closed vessel, and then dissolved in acetonitrile (10 mL), whereas sample of liquid preparation was 100-fold diluted with acetonitrile. The 1 mL of acetonitrile solutions were taken for the derivatization.

Fig. 2 Chromatograms obtained for blanks (a) and mixture of 0.5 mg L\(^{-1}\) PFCA standards (b) for derivatization with 2 gL\(^{-1}\) (A) and 5 gL\(^{-1}\) (B) solutions of 3-BrAC
analytes depend on numerous experimental factors [21–24]. In this study two different commercially available coumarin derivatives were examined as derivatization reagents for PFCAs, namely 3-BrAC and BrMMC. Using standard mixture containing PFCAs from C2 to C12 at 0.5 mg L$^{-1}$ concentration each, for both derivatives effect of reagent concentration used for derivatization, reaction time, temperature of reaction and kind of solvent were examined.

Comparison of two reagents used for derivatization was made in 70 °C, for reaction time 60 min and concentration of reagents 2 gL$^{-1}$. The obtained chromatograms with corresponding recordings for blanks are shown in Fig. 1. In similar period of time about 25 min a better separation of all analytes and more stable base-line was obtained for samples derivatized with 3-BrAC. Surface areas for recorded peaks with two coumarin derivatives are very similar. Much larger signal for excess of reagent recorded for BrMMC makes impossible the recording of signals corresponding to C2 and C3-PFCA. A much larger signal recorded for C6-PFCA with BrMMC is caused by unidentified solutes present in blank. For further investigations 3-BrAC was selected as the derivatizing reagent.

The signal magnitude for derivatized PFCAs depends to some degree on concentration of 3-BrAC taken for derivatization, which was examined for 2 and 5 gL$^{-1}$ concentration. As one can see from recorded chromatograms (Fig. 2), about double increase of the signal magnitude for larger concentration of coumarin reagent is observed, which is however associated with larger signals for blank. For further studies 2 gL$^{-1}$ concentration was used.

The effect of temperature used in derivatization reaction is illustrated by the histogram in Fig. 3A showing data obtained in the temperature range from 20 to 70 °C, for reaction time 60 min. Its significant influence is evident for all examined analytes. At the highest examined temperature of derivatization 70 °C also effect of derivatization time was investigated in the range from 20 to 120 min (Fig. 3B). The role of this parameter is smaller, and the decrease of signal magnitude at 120 min for most analytes was attributed to the loss of analytes during long time derivatization. As some loss of more volatile species can be expected also at highest examined temperature, as the most safe and efficient conditions of derivatization were selected 60 min derivatization at 70 °C.

Due to large differences in affinity of examined analytes to C18 stationary phase in reversed-phase system, their

| PFCA | Retention time (min) | Regression equations | Correlation coefficient | LOD (μg·l$^{-1}$) |
|------|----------------------|----------------------|------------------------|------------------|
| C2   | 6.5                  | y=336356 x           | 0.9997                 | 7.5              |
| C3   | 9.1                  | y=356820 x           | 0.9998                 | 5.5              |
| C4   | 11.3                 | y=640191 x           | 0.9997                 | 6.0              |
| C5   | 13.3                 | y=224169 x           | 0.9996                 | 6.8              |
| C6   | 15.0                 | y=354849 x           | 0.9995                 | 6.5              |
| C7   | 16.5                 | y=222639 x           | 0.9997                 | 6.6              |
| C8   | 17.9                 | y=190238 x           | 0.9998                 | 4.6              |
| C9   | 19.1                 | y=251250 x           | 0.9998                 | 4.3              |
| C10  | 20.2                 | y=111682 x           | 0.9997                 | 6.8              |
| C11  | 21.1                 | y=88769 x            | 0.9995                 | 6.5              |
| C12  | 22.0                 | y=95421 x            | 0.9998                 | 7.5              |
efficient separation in reasonable time period can be obtained for gradient elution, only. The additional factor that should be considered is the elimination of possible overlapping of signals for analytes with excess of reagent used for derivatization. For separation of mixture of PFCAs from C2 to C12 three different gradient programs were investigated, using 60% aqueous solution of ACN (A) and pure ACN (B). Initially, isocratic elution with eluent A during 5 min (Fig. 4a), 6 min (Fig. 4b) and 4 min (Fig. 4c) was used. Next linear gradient elution from 5 to 20 min (Fig. 4a), 6–18 min (Fig. 4b) and 4–15 min (Fig. 4c) was applied. As it is demonstrated by recorded chromatograms shown in Fig. 4 program (a) can be considered as the most appropriate.

Figures of merit of developed method

In optimized conditions of derivatization the determinations of PFCAs were carried out in concentration range from 0.025 to 0.5 mg L$^{-1}$ in injected solutions. For 50 μL sample volume satisfactory linearity of response was obtained and limits of detection (for S/N=3) in the range from 5 to 10 μg L$^{-1}$ (Table 1).

In the study of repeatability of derivatization procedure with 3-BrAC in optimized conditions, for 5 repetitions carried out in 5 consecutive days, RSD values for peak areas for all analytes were below 1%. No significant differences in the efficiency of the separation were also observed. Derivatized analytes were stable in time, as after 2 days of storage recorded chromatograms were practically the same as those recorded immediately after derivatization. No additional by-products after 2 days storage were observed.

Optimization of SPE preconcentration procedure

The detection limits obtained in developed procedure are too high for direct determination of PFCAs at level present
in natural waters [16, 25]. As this application is one of the targets of this work, the next step in development of the complete procedure was the optimization of the solid-phase extraction method for the preconcentration of analytes from diluted natural samples. Based on vast literature on the application of different sorbents for this purpose [26], two commercially available sorbents were selected for this purpose: Sep-PAK C18 and Oasis-WAX. Besides the optimization of elution mode for retained species on the sorbent bed, another essential factor was to eliminate traces of water from extracts, which interferes in derivatization reaction.

For sorbent Sep-PAK C18 optimum pH range is from 3 to 7 and it can be used for preconcentration of hydrophobic solutes. Mixtures of PFCAs containing 30, 3.0 or 0.3 μg L⁻¹ each were preconcentrated from sample volumes 25, 250 and 1,000 mL, respectively, in order to obtain well measurable amounts of analytes in the final extract, to examine the recovery of preconcentration. The derivatization with coumarin derivatives proceeds with larger yield in acetonitrile than in methanol, hence in the first attempt ACN was tested as eluent of retained analytes, however for this solvent much lower recoveries were observed than for methanol. In alternative procedure, elution was carried out with methanol, which then was evaporated in the argon stream, and solid residue was dissolved in ACN for further derivatization.

Because of strong retention of examined analytes on C18 sorbent bed, the recovery of elution was investigated using three 2.5 mL fractions of methanol. The obtained data for 100-fold preconcentration using 500 mg sorbent bed are shown in Fig. 5. The satisfactory recovery (above 80%) with single portion of methanol was obtained for C4, C7 and C9 PFCAs, only. All data represents mean of 3 parallel measurements. The total recovery obtained by means of three fractions of eluting methanol was satisfactory for all examined analytes from C3 to C12 PFCA, except C2 where much lower total recovery was found. The reproducibility of this procedure was also satisfactory, as the RSD was obtained for various analytes in the range from 0.2 to 1.4%.

For 1000-fold preconcentration from 1 L sample containing 0.3 mg L⁻¹ of each PFCA, satisfactory total recovery above 90% was obtained for C4 to C6 PFCAs, only, whereas for other examined analytes it was much lower, and non-acceptable for practical applications.

Similar studies were carried out also for Oasis-WAX cartridges. As according to manufacturer this sorbent can be used for partly polar analytes with pKₐ values about or larger than 1.0, it seems to be suitable for examined analytes, where retention is based on both hydrophobic and ion-exchange interactions. Investigation of the role of amount of sorbent taken for preconcentration and level of ammonia in methanol was investigated, however, a satisfactory elution (recovery better than 80%) was observed for few PFCAs, only, namely C4, C5, C6 and C9. Amount of sorbent used for preconcentration affects obtained recovery but in used conditions where 100-fold preconcentration was carried out from 250 mL samples containing 3 μg L⁻¹ of each PFCA doubling amount of sorbent from 60 to 120 mg did not affect recovery of analytes. Elution with three consecutive portions of 4.0 mL methanol containing ammonia for 100-fold preconcentration, also does not improve total recovery. Based on obtained experimental data C18 sorbent Sep-PAK was considered as more suitable for the preconcentration of trace amounts of PFCA from environmental samples.

Analytical applications of developed method

Two different examples were examined to illustrate the possible applications of the developed method. In the first one, PFCAs content was measured in two preparations for shoe impregnation, which did not require preconcentration step. Following the procedure given in Experimental, in acetonitrile solution obtained from aerosol preparation Kiwi, three PFCAs have been determined, namely C3, C8 and C9 PFCA at level 0.137, 0.107 and 0.215 mg L⁻¹, which corresponds to 4.85, 3.79 and 7.62 μg g⁻¹ of analyzed samples. In liquid preparation Boreal, only C5-PFCA was

| Method                        | Sample                     | LOD                                         | Reference       |
|-------------------------------|----------------------------|---------------------------------------------|-----------------|
| GC-MS                         | Harbor seawater Waste water| 0.02–0.75 μg L⁻¹ (with SPME preconcentration from 5 mL samples) | [27]            |
| HPLC with conductivity detection | Industrial process liquids | 0.12–0.66 mg L⁻¹ and 50 μg L⁻¹ (with SPE preconcentration from 100 mL samples) | [15]            |
| HPLC with fluorescence detection | Liver homogenate            | 50 pmol/50 mg tissue                        | [18]            |
| LC-MS/MS                      | Seawater                   | 1.8–5.2 pg L⁻¹ (with SPE from 1 L sample)    | [9]             |
| HPLC with fluorescence detection | Surface water              | 43–75 ng L⁻¹ (with SPE preconcentration from 100 mL samples) | This work       |

SPE solid-phase extraction; SPME solid-phase micro extraction.
determined at 0.85 mg L\(^{-1}\) level. The example recorded chromatogram for analyzed samples is shown in Fig. 6A.

A complete procedure with 100-fold preconcentration with C18 Sep-PAK 500 mg cartridge was employed for determination of PFCAs in surface water sample collected from industrial region close to Tarnów city in Southern Poland. Chromatograms recorded for blank, standard mixture of PFCAs, and analyzed sample are shown in Fig. 6B. Based on retention times, two PFCA analytes were identified, namely C7 and C8 (PFOA) at concentrations 1.0, 0.63 ppb, respectively. Recoveries 93.2% and 99.3% for C7 and C8 in natural water were obtained.

Conclusions

The environmental importance of perfluorinated organic compounds widely discussed in recent decade requires a continuous improvement of the analytical methods for routine use. Although they are commonly determined in advanced analytical laboratories by LC/MS with satisfactory selectivity and limits of detection, there is also a need for methods suitable for more common use in e.g. local environmental laboratories. Such a role can be played by the reversed-phase HPLC method with fluorescence detection developed in this work, which is demonstrated for the determination of PFCAs, a widely proliferated group of perfluorinated organic compounds. The application of commercial derivative of coumarin allows relatively simple derivatization with well repeatable formation of derivatized products. Developed gradient method of elution allows efficient separation of whole range of PFCAs from C3 to C12. With obtained limits of detection in the range from 4.3 to 7.5 ppb it can be used for the control of content of these species in household preparations or cosmetics.

Applications in environmental analysis e.g. for surface water analysis require additional preconcentration of trace analytes. Satisfactory results were obtained with 100-fold preconcentration using commercial solid-phase extraction cartridges C18 Sep-PAK, with recoveries at least 80%. This allows obtaining LODs values in the range 43–75 ppb, which is satisfactory for environmental applications for monitoring of these analytes. In comparison to the previous work on application of HPLC with fluorimetric detection for analysis of rats liver samples [18], in this work the commercial derivatizing reagent was employed and developed method was optimized for trace analysis of environmental samples. Taking into account of a current wide demand for monitoring of perfluorinated compounds in environmental samples, the developed method can be the easier available alternative compared to costly and required highly trained personnel LC/MS methods for routine environmental applications.

As it is shown by comparison of LOD values reported for different chromatographic methods (Table 2), only application of LC/MS allows the obtained better limits of detection, however LC/MS instrumentation still is too expensive for common application in environmental monitoring. The LOD values obtained in this work illustrate its possible application in routine environmental analysis for PFCA determination [17] with use of much simpler and more commonly employed instruments. Such possibility was not so far investigated with HPLC using fluorescence detection for environmental samples. Similar method with different fluorophore was developed only for analysis of biological material [18].

Acknowledgements The authors thank Dr. Bogdan Szostek (E.I. DuPont De Nemours & Co. Inc., Wilmington, DE, USA) for his advice and assistance during the conduct of experiments and manuscript preparation. We acknowledge also financial support from DuPont’s Center for Collaborative Research and Education, Europe under the European University Support Program.

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