

M. LÖHMUS, Piret NIIDAS, M. LOPP, Ü. LILLE

QUANTITATIVE ANALYSIS OF PROSTACYCLIN BY HIGH PERFORMANCE LIQUID CHROMATOGRAPHY

Prostacyclin (prostaglandin (PG) I₂) was discovered by J. R. Vane and his co-workers in 1976 [1]. It appeared to be the strongest inhibitor of platelet aggregation described so far [1]. Also, PG_{I2} was further characterized as a potent vasodilator and cytoprotector [1, 2]. These properties make it an important candidate for therapeutic use. Unfortunately, PG_{I2} is chemically extremely unstable. In acid or neutral aqueous conditions, its cyclic enoether unit is hydrolyzed within a few minutes into 6-keto-PGF_{1α} [1, 3, 4]. Although under unhydrous conditions PG_{I2} Na-salt is stable in crystalline form, determination of the chemical purity of prostacyclin samples is obligatory for investigators studying the physiological activity of this compound.

The chemical instability of PG_{I2} gives rise to serious problems in its analysis. In solving these problems, G. T. Hill suggested the use of high performance liquid chromatographic method on reverse phase columns [5]. M. A. Wynalda, F. N. Lincoln and F. A. Fitzpatrick found an excellent mobile phase for eluting PG_{I2} Na-salt and its impurities. It consists of acetonitrile and water buffered at pH 9.3 with boric acid and sodium borate [6]. They used the external standard method that requires full-loop injections and regular peak shapes. The external standard method requires, before each series of sample analysis, injection of PG_{I2} Na-salt with high purity for comparison. This gives once more rise to some problems due to the instability of PG_{I2} Na-salt. Besides, the precision of the method is low [6]. Such a situation suggests the need for the development of an internal standard method for analyzing PG_{I2} Na-salt.

Two methods were tested. In the first method one internal standard and specially purified PG_{I2} Na-salt are used for calibration and in the second one two internal standards plus technical grade PG_{I2} Na-salt are employed.

Materials and methods

Apparatus. A DuPont HPLC system N 8845 with a Rheodyne Model 7125 Syringe Loading Sample Injector (maximum sample size 50 µl). A spectrophotometric detector was set at 205 nm and 0.64 AUFS. A recorder operated at 10 cm/h.

Reagents. Boric acid, sodium borate and sodium hydroxide (Reakhim, USSR) were used without purification. Acetonitrile (Reakhim) was redistilled. Bidistilled water was used. PG_{I2} Na-salt (99% by HPLC, mp. 164–166°) was prepared according to [7–10] and crystallized twice from water-acetonitrile (0.5:15). 6-keto-PGF_{1α} methyl ester was prepared in different batches according to [7]. Ester moiety was hydrolyzed by a 1N-NaOH solution in methanol. Biosynthetic PGE₁ (mp. 114–116°)

used also as an internal standard was tested to be at least 99% pure by HPLC [1].

Chromatographic conditions. Chromatography was performed at ambient temperature on a 4.0×220 mm Zorbax ODS column or 4.6×250 mm column prepacked with a Zorbax ODS. The columns were eluted with acetonitrile-water (85—87% of water, buffered at pH 9.3 with 0.01 M boric acid and 0.006 M sodium borate) at a flow rate of 0.8—1.0 ml/min.

Direct analysis with PGE₁ as an internal standard. Method A. In order to obtain the calibration curve, 5 different quantities of pure PGI₂ Na-salt in the range of 1.0—3.5 mg were weighed (W_I) and the precise amounts (1.0—3.5 mg) of PGE₁ (W_E) were added; the samples were dissolved in 300 μ l of water buffered at pH 10.1 with 0.013 M sodium borate and 0.023 M sodium hydroxide at 4°. In each case, 5 μ l of the sample was injected and the rest was kept at 4° between reinjections. Each sample injection was performed twice.

Analyses with PGE₁ and 6-keto-PGF_{1 α} as internal standards. Method B. In order to obtain the calibration curve, 11 different quantities of PGI₂ Na-salt in the range of 1.0—3.5 mg were weighed and the precise amounts (1.0—3.5 mg) of PGE₁ (W_E) were added. An aqueous solution of 6-keto-PGF_{1 α} was prepared in a concentration of 10 mg/ml, buffered at pH 10.1 with 0.013 M sodium borate and 0.023 M sodium hydroxide and kept at 4° before using. Thereupon, 8—100 μ l of the 6-keto-PGF_{1 α} solution (the weight of 6-keto-PGF_{1 α} (W_F)) was added to the weighed samples of PGI₂ Na-salt. The same buffer was added till the total volume of the sample of 300 μ l. In each case, 5 μ l of the sample was injected and the rest was kept at 4° between reinjections. Each sample injection was performed twice. Subsequently, PGI₂ Na-salt was converted into 6-keto-PGF_{1 α} by acidifying the samples with acetic acid up to pH 5—6. Then, 0.1 N sodium hydroxide was added at 4° up to pH 8—9 and 15 μ l of the sample was injected. The samples were stored and reinjected as described above.

Calculations. The peak areas were calculated by «Height \times Width at Half Height» technique [12].

Response factor of PGI₂ Na-salt versus PGE₁

$$f = \frac{A_I W_E}{A_E W_I} \quad (1)$$

where A_I and A_E are the peak areas of PGI₂ Na-salt and PGE₁, W_I and W_E are their weights, respectively.

Corrected PGE₁ peak area

$$A_E^{corr} = A_E - A_I \times k \quad (2)$$

where k is the area fraction of interfering impurity in PGI₂ Na-salt.

Formulas analogous to (1) and (2) may be used in calculations for 6-keto-PGF_{1 α} when PGI₂ Na-salt contains 6-keto-PGF_{1 α} .

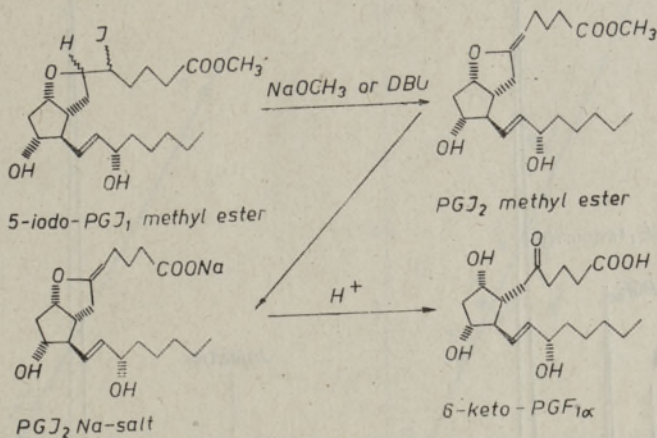
Real weight of PGI₂ Na-salt (the difference in molecular masses of 6-keto-PGF_{1 α} and PGI₂ Na-salt is neglected)

$$W_I^{real} = \left(\frac{R''_A}{R'_A} - 1 \right) W_F \quad (3)$$

where R'_A and R''_A are the ratios of peak areas of 6-keto-PGF_{1 α} to those of PGE₁ before and after acidification, respectively, W_F is the weight of 6-keto-PGF_{1 α} .

Results and discussion

PGI₂ Na-salt is synthesized from 5-iodo-PGI₁ methyl ester according to the following scheme:



The main impurities of PGI₂ Na-salt (technical grade) are 6-keto-PGF_{1\alpha}, an unidentified compound in an amount of 1–3.5% of PGI₂ Na-salt and inorganic salts (NaOH, NaI).

The internal standard method requires that the standard should elute near the peaks of interest and should be of high purity and chemically stable. We found that PGE₁ meets these requirements. It has an acceptable retention time under the chromatographic conditions used. The capacity factor k' , measured on a Zorbax ODS column for 6-keto-PGF_{1\alpha}, PGE₁, and PGI₂ Na-salt, was 0.9, 3.4 and 9.0, respectively (Fig. 1). The fact that PGE₁ elutes between 6-keto-PGF_{1\alpha} and PGI₂ Na-salt is also of importance. The purity of the crystalline PGE₁ (99% by HPLC) meets the analytical requirements.

It is well known that under basic conditions PGE₁ converts into PGB₁ via PGA₁ [13, 14]. By examining UV-spectra of PGE₁ at ambient temperature we found that at pH 9.3 less than 1% of PGE₁ is degraded into PGA₁ ($\lambda_{\max} = 217$ nm; $\epsilon = 10830$) [15] during 1 h. At pH 10.1 1.4% of PGE₁ was converted into PGA₁ in 30 min. The conversion of PGA₁ into PGB₁ ($\lambda_{\max} = 278$ nm; $\epsilon = 26800$) [16] proceeds much slower. Only 0.7% of PGE₁ can be converted into PGB₁ in 18 hs at pH 10.1. Also, by HPLC two injections from one sample under the above conditions were coincident with a precision of 1%. No trace amounts of PGA₁ or PGB₁ were detected.

Calibration of PGI₂ Na-salt versus PGE₁ was performed by two different methods. In the first method (method A) the twice recrystallized PGI₂ Na-salt (Fig. 2) was used. The real peak areas of the weighed amounts of PGI₂ Na-salt and PGE₁ were measured. An assumption was made that the response of the detector to PGI₂ Na-salt and PGE₁ is linear to their amount in the investigated region. From the calibration curve obtained (Fig. 3, a) the response factor f_A of PGI₂ Na-salt versus PGE₁ on the UV-detector at 205 nm was calculated and found to be 3.65 ± 0.15 at a confidence level of 95%.

By the second method (method B), the peak areas of 6-keto-PGF_{1\alpha} (A_F) and PGI₂ Na-salt (A_I) were measured and the peak area of PGE₁ was calculated according to formula (2). For each sample the ratio of the peak areas of 6-keto-PGF_{1\alpha} to the peak area of PGE₁ (R'_A) was calculated. Analogously, these ratios (R''_A) were determined after acid-

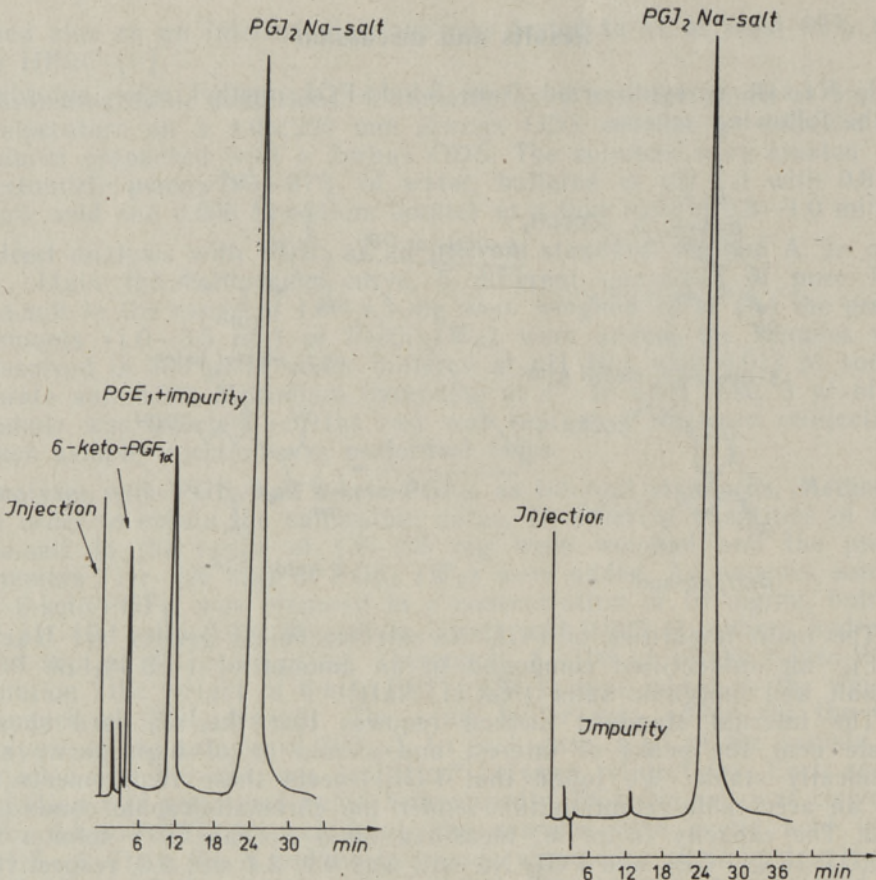


Fig. 1. Chromatogram of a mixture of 6-keto-PGF_{1 α} and PGE₁ and PGI₂ Na-salt (technical grade) used for calibration in method B. Chromatography was performed on a Zorbax ODS (4.6×250 mm) column using acetonitrile-water as mobile phase (13:87, v/v), buffered at pH 9.3 with 0.01 M boric acid and 0.006 M sodium borate. The flow rate was 1.0 ml/min. The detector was set at 205 nm and 0.64 AUFS. The recorder operated at 10 cm/h.

Fig. 2. Chromatogram of PGI₂ Na-salt (standard grade) used for calibration in method A. Chromatographic conditions are as in Fig. 1.

ifying the samples. From the increase in the peak areas ratio the amount of PGI₂ Na-salt (W_1) in the sample was calculated according to formula (3). By method B the response factor of PGI₂ Na-salt versus PGE₁ (f_B) was found to be 4.3 ± 1.1 at a confidence level of 95%. The response factor of 6-keto-PGF_{1 α} versus PGE₁ was found to be 0.68 ± 0.20 at a confidence level of 95%.

As can be seen from Fig. 3, method A gives a calibration curve which is in accordance with experimental data. Method B, on the contrary, gives a calibration curve, with a great deviation from experimental data. Consequently, method A is superior to method B in the analysis of PGI₂ Na-salt samples. By the statistical treatment of the deviation of experimental points we found that it cannot be subjected to normal distribution. In addition we found that between the experimental points 1—5 and 9—11 (Fig. 3, b) (these points correspond to different 6-keto-PGF_{1 α} batches) a systematic error occurs. This suggests a

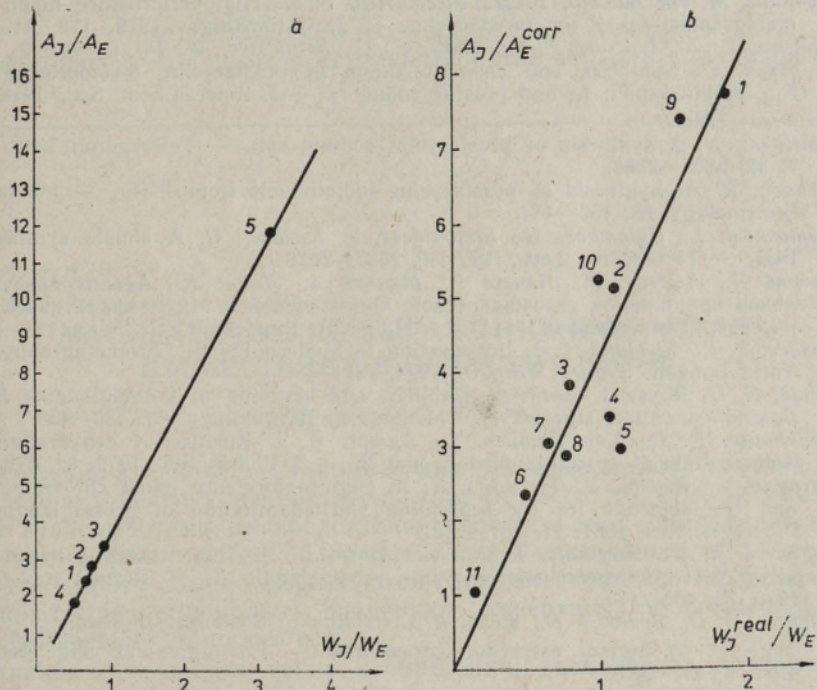


Fig. 3. Calibration curves of PGI_2 Na-salt. *a* — method A, *b* — method B.

conclusion that the purity of 6-keto-PGF_{1 α} in different batches is different due to difficulties in the preparation, extraction and purification of this compound [8, 17]. The conditions under which 6-keto-PGF_{1 α} can be used for constructing a calibration curve require further study.

Conclusions

1. The internal standard method using PGE₁ as an internal standard for determining the purity of PGI_2 Na-salt was developed.
2. The direct calibration method where a solution of pure PGI_2 Na-salt was used (method A) is superior to the method via 6-keto-PGF_{1 α} (method B).
3. PGI_2 Na-salt response factor to PGE₁ at 205 nm is 3.65 with a standard deviation of 0.05. It provides sufficient precision for determining the purity of PGI_2 Na-salt.

REFERENCES

1. Moncada, S., Gryglewski, R., Bunting, S., Vane, J. R. An enzyme isolated from arteries transforms prostaglandin endoperoxides to an unstable substance that inhibits platelet aggregations. — Nature (London), 1976, 263, 663—665.
2. Vane, J. R. Bioassay-Abenteuer auf dem Weg zum Prostacyclin (Nobel-Vortrag). — Angew. Chem., 1983, 95, 782—794.
3. Pace-Asciak, C. Isolation, structure and biosynthesis of 6-ketoprostaglandin F_{1 α} in the rat stomach. — J. Amer. Chem. Soc., 1976, 98, 2348—2349.
4. Chiang, Y., Kresge, A. J., Cho, M. J. Acid catalyzed hydrolysis of prostacyclin: origin of the unused lability. — J. Chem. Soc. Chem. Commun., 1979, N 2, 129—130.
5. Hill, G. T. High-performance liquid chromatographic determination of prostacyclin. — J. Chromatogr., 1979, 176, 407—412.

6. Wynalda, M. A., Lincoln, F. H., Fitzpatrick, F. A. High-performance liquid chromatographic assay for prostacyclin. — J. Chromatogr., 1979, **176**, 413—417.
7. Johnson, R. A., Lincoln, F. H., Nidy, E. G., Schneider, W. P., Thompson, J. L., Axen, U. Synthesis and characterization of prostacyclin, 6-ketoprostaglandin $F_{1\alpha}$, prostaglandin I_1 , and prostaglandin I_3 . — J. Amer. Chem. Soc., 1978, **100**, 7690—7705.
8. Whittaker, N. A synthesis of prostacyclin sodium salt. — Tetrahedron Lett., 1977, N° 32, 2805—2808.
9. Johnson, R. A. Synthesis of prostacyclin sodium salt from PGF₂. — Methods in Enzymology, **86**, 459—464.
10. Tömösközi, I., Galambos, G., Simonidesz, V., Kovács, G. A simple synthesis of PGI₂. — Tetrahedron Lett., 1977, **30**, 2627—2628.
11. Самель Н., Лыхмус М., Алисте Р., Мянник А., Лилле Ю. Анализ простагландинов при помощи газожидкостной, тонкослойной и высокоэффективной жидкостной хроматографии. — Изв. АН ЭССР. Хим., 1981, **30**, № 3, 199—207.
12. Snyder, L. R., Kirkland, J. J. Introduction to Modern Liquid Chromatography. New York; London; Sydney; Toronto, 1974, 431—444.
13. Stehle, R. G. Physical chemistry, stability, and handling of prostaglandins E_2 , F_2 , D_2 and I_2 : critical summary. — Methods in Enzymology, **86**, 436—458.
14. Monkhouse, D. C., Van Campen, L., Aguiar, A. J. Kinetics of dehydration and isomerization of prostaglandins E_1 and E_2 . — J. Pharm. Sci., 1973, **62**, 576—580.
15. Terragno, A., Rydzik, R., Terragno, N. A. High-performance liquid chromatography and UV detection for the separation and quantitation of prostaglandins. — Prostaglandins, 1981, **21**, 101—112.
16. Nugteren, D. H., Beertuis, R. K., Van Dorp, D. A. The enzymic conversion of all-cis 8,11,14-eicosatrienoic acid into prostaglandin E_1 . — Recueil trav. chim., 1966, **85**, 405—419.
17. Mitchell, M. D., Brunt, J. D., Webb, R. Instability of 6-keto-prostaglandin $F_{1\alpha}$ when subjected to normal extraction procedures. — Prostaglandins and Medicine, 1981, **6**, 437—440.

Academy of Sciences of the Estonian SSR,
Institute of Chemistry

Received
Feb. 29, 1984

M. LOHMUS, Piret NIIDAS, M. LOPP, Ü. LILLE

PROSTATSÜKLIINI KVANTITATIIVNE ANALÜÜS KÖRGEFEKTHIIVSEL VEDELIKUKROMATOGRAAFIA MEETODIL

On välja töötatud prostatsükliini määramise kvantitatiivne meetod, kasutades sisestandardit. On võrreldud kahte meetodit: A — sisestandardiks oli PGE₁; B — sisestandarditeks olid 6-keto-PGF_{1α} ja PGE₁.

Meetod A andis piisava täpsusega tulemused. 205 nm juures oli PGI₂ detekteerimise tundlikkus $3,65 \pm 0,15$ korda suurem kui PGE₁-l.

M. ЛЫХМУС, Пирет НИЙДАС, М. ЛОПП, Ю. ЛИЛЛЕ

КОЛИЧЕСТВЕННЫЙ АНАЛИЗ ПРОСТАЦИКЛИНА С ПОМОЩЬЮ МЕТОДА ВЫСОКОЭФФЕКТИВНОЙ ЖИДКОСТНОЙ ХРОМАТОГРАФИИ

Разработан количественный метод определения чистоты натриевой соли простациклина с использованием внутреннего стандарта, в качестве которого служили ПГЕ₁ (метод А) и 6-кето-ПГF_{1α}, ПГЕ₁ (метод Б). Метод А дал результаты удовлетворительной точности. При 205 нм чувствительность детектирования натриевой соли ПГI₂ превышала чувствительность детектирования ПГЕ₁ в $3,65 \pm 0,15$ раза.