PAPER DETAILS

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A Facile HPLC-PDA Method for Simultaneous Determination of Paracetamol, Methyl Paraben, Sunset Yellow, and Carmosine in Oral Suspensions

Şule Dinç Zor*, Özlem Aksu Dönmez

Department of Chemistry, Faculty of Science and Arts, Yildiz Technical University, 34220 Davutpasa-Istanbul, Turkey

In the present study, a simple, fast, and accurate HPLC-PDA method was developed for the simultaneous determination of paracetamol (PAR), methylparaben (MP), sunset yellow (SSY) and carmoisine (CAR) in oral suspensions. The concentrations of colorants are less than with respect to those of active ingredient and this variation makes process of analysis troublesome. In the developed HPLC method, efficient chromatographic separation was achieved using reversed phase C18 column (4.6 mm x 150 mm x 5 μ m particle size) and phosphate buffer solution (pH = 6.5)-acetonitrile mobile phase with a flow rate of 1.6 mL/min in the gradient mode. The eluents were monitored via a PDA detector at 300, 254 and 230 nm. The mean retention times of PAR, MP, SSY and CAR were found to be 2.15, 4.42, 1.58 and 3.81, respectively. The proposed method was validated in accordance with ICH guidelines and it was seen that the method met all requirements in terms of linearity, precision, accuracy, and selectivity. The developed method was successfully applied for simultaneous determination of the studied compounds in two commercial oral suspension samples.

Keywords: Paracetamol, methylparaben, sunset yellow, carmoisine HPLC-PDA, oral suspension.

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*Corresponding author. E-mail: <u>sule_dinc@yahoo.com</u>. Tel: +90-2123834156.

INTRODUCTION

Paracetamol (N-acetyl-4-aminophenol) is widely used in pediatric syrup-suspension formulations as a pain-reliever and fever-reducer. It is easily available alone without a prescription or in combination with other drugs. Some preservatives and colorants are commonly used in these pediatric pharmaceutical formulations to prevent microbial growth and to improve appearance and color (1). In general, sodium benzoate, potassium sorbate or parabens such as methyl paraben or propyl paraben are used as a preservant system in the liquid formulations (2). Methyl paraben (methyl 4-hydroxybenzoate) is the most frequently used due to its broad antimicrobial spectrum and the fact that it does not modify the physical properties of the final products like taste, smell or color (3). According to some research reports, parabens exhibit estrogenic activity and their extreme usage could lead to some detrimental effects such as breast cancers and oxidative DNA damage (4-6). Owing to these concerns, many countries have put a ban to limit the use of parabens (7). Hence, determinations of these preservatives in pharmaceuticals are vitally necessary for both quality assurance and consumer safety.

Although the allowable amounts of synthetic colorants which could have toxicity are reduced by human health reasons, many synthetic colorants in stead of natural colorants are still widely used due to their low price, high effectiveness, and excellent stability in foods and pharmaceuticals (8-10). Since synthetic colorants could provoke allergic reactions including urticaria, dermatitis, and asthma and may give rise to hyperactive behavior in children, the use of synthetic colorants in many countries is strictly regulated under existing food laws (11,12). Thus, determination of colorants in food samples and pharmaceutical products is important to control the amount of use permitted and to ensure quality control. Also, in order to colorize pharmaceuticals, the mixture of two or three of them is used to create a hue corresponding with selected natural color. So, simultaneous determination of these colorants becomes difficult as the number of components in the mixtures increases (13). Furthermore, since the quantity of active compound is commonly higher than additives, there is a need of accurate, efficient, and fast analytical method for simultaneous quantification of ingredients and excipients (14,15). In this respect, we focused on simultaneous high performance liquid chromatography (HPLC) determination of the single drug, paracetamol, along with colorants, sunset yellow and carmoisine, and preservative, methyl paraben, in pediatric oral suspension samples that belong to the same pharmaceutical company, in this study. Many analytical methods, either for single or combination with other analytes in various matrices, have been described in the literature for determination of paracetamol (16,17), methyl paraben (18,19), sunset yellow (20,21) and carmoisine (22,23) by HPLC in particular. However, no method describes quantification of this drug and color and preservative additives simultaneously. There are some HPLC methods developed in order to quantify different active compounds and additives in association, which are non labor-intensive, without extraction step and with short analysis time (2, 13, 24-27).

764

This study aims a fast, simple, and sensitive HPLC method for the simultaneous determination of paracetamol, methyl paraben, sunset yellow and carmoisine in pediatric oral suspensions. Validation parameters for the current method were also tested according to the requirements of ICH guidelines.

MATERIALS and METHODS

Chemicals and Reagents

Reference standards of paracetamol (PAR), methyl paraben (MP), sunset yellow (SSY) and carmoisine (CAR) were obtained as gifts from a local pharmaceutical company (all purities \geq 99%). HPLC grade acetonitrile, methanol, orthophosphoric acid (H₃PO₄) and di-potassium hydrogen phosphate (K₂HPO₄) were purchased from Merck Chemicals (Germany). Ultrapure water was produced by a Milli-Q® Elix Water Purification System (Milford, MA, USA).

Instruments and Chromatographic Conditions

The HPLC system was a Shimadzu HPLC system LC-10AT VP equiped with a SIL-20AC autosampler and SPD-M10A VP photodiode array dedector (PDA). The chromatographic separations were performed on an Inertsil C18 column (4.6 mm x 150 mm x 5 µm particle size, GL Sciences, Japan). The mobile phase was made up of phosphate buffer (0.025 M, pH 6.5) and acetonitrile. Gradient elution conditions are given in Table 1. The flow rate of mobile phase was 1.6 mL/min at room temperature and injection volumes were 20 µL. The eluents were monitored in the range of 190 to 800 nm via a PDA dedector and the detections were carried out at 300 nm for PAR, at 254 nm for MP, at 230 nm for SSY and CAR. The run time was approximately 5 min and the total peak area was used for the quantification of each analyte.

Mobile Phase A	Mobile Phase B	Gradient Conditions	
Acetonitrile	Phosphate buffer (0.025 M, pH 6.5)	87%B 2.0 min, 70%B 3.0 min	

Table 1. Gradient elution conditions for the separation of analytes.

Preparation of Standard and Sample Solutions

Stock standard solutions of PAR (5000 μ g/mL) and MP (500 μ g/mL) were prepared in methanol by accurately weighting. Stock standard solutions of SSY and CAR (100 μ g /mL) were prepared in Milli-Q water. All stock solutions were stored at 4 °C and further dilutions to obtain calibration and other validation studies solutions were made in Milli-Q water.

Two kinds of marketed liquid pharmaceutical formulations (suspension) comprising an aqueous solution of paracetamol were purchased from local pharmacy shop in Istanbul, Turkey. 2.5 mL of Suspension I containing 250 mg PAR, 4 mg MP and an unknown amount of SSY in 5 mL was

accurately transferred into an 100 mL measuring flask, sonicated in ultrapure water, and the volume was then made up to the mark with the same solvent. Similarly, 5.0 mL of Suspension II containing 120 mg PAR, 5 mg MP and an unknown amount of CAR in 5 mL was accurately transferred into a 50 mL measuring flask, sonicated and diluted to its volume with ultrapure water. All standard and sample solutions were filtered through 0.45 micron membrane filter.

Method Validation

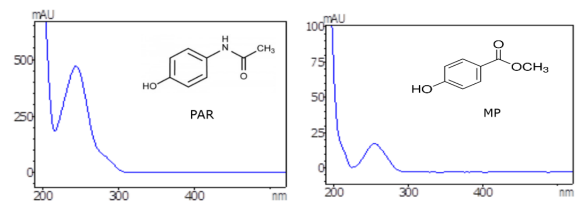
Method validation including selectivity, linearity, limit of detection and limit of quantification, precision and accuracy of the proposed method was performed according to ICH guidelines (28).

RESULTS and DISCUSSION

Method Development

In order to achieve good chromatographic separation of the studied compounds, different solvents (water, acetonitrile and methanol) and buffer solutions (acetic acid/acetate and phosphate), ideal mobile phase proportion, pH and flow rate were duly studied. Satisfactory results were achieved by 87:13% (v:v) of phosphate buffer (0.025 M) pH 6.5 : acetonitrile for first 2.0 min and then 70:30% (v:v) of the same mobile phase for 2.0-5.0 min at flow rate of 1.6 mL/min in the gradient mode.

As the quantity of active ingredient in these pharmaceutical formulations is compared to those of colorants in particular, it is seen that there is an imbalance between the analytes. Generally, the amount of the colorants is less than the active compound and excipients. This variation makes the simultaneous chromatographic analysis of them along with other ingredients difficult. This difficulty can be overcome by choosing the appropriate wavelength for determination. Absorption spectra of the studied compunds can be seen in Figure 1. Detection wavelength was chosen at 300 nm for PAR, at 254 nm for MP, at 230 nm for SSY and CAR, accordingly.



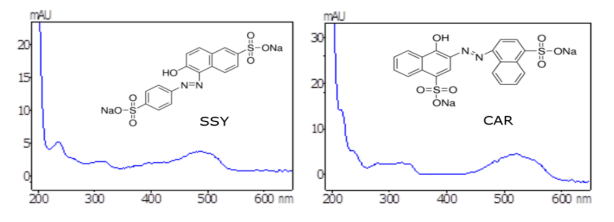
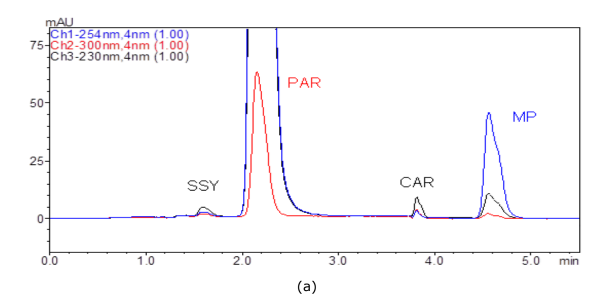
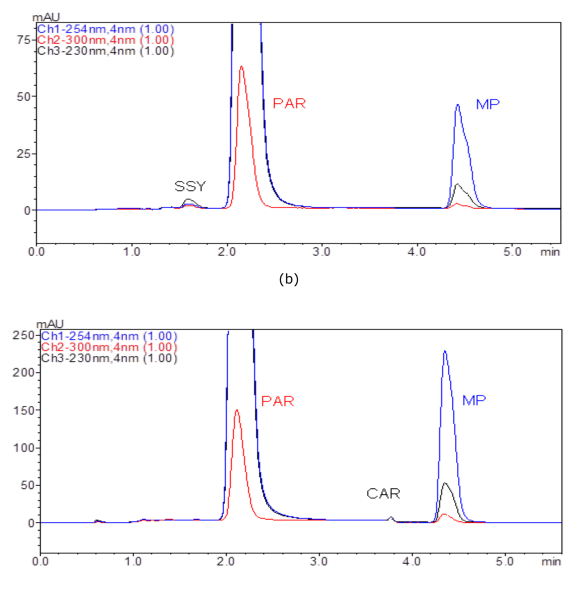


Figure 1. Absorption spectra of paracetamol (PAR), methyl paraben (MP), sunset yellow (SSY) and carmoisine (CAR) by PDA dedector.

Figure 2 shows the chromatogram obtained for the analytes in synthetic mixture and pharmaceutical formulations by the chromatographic conditions described above.





c)

Figure 2. a) Chromatogram of standard mixture solution containing 4 μ g/mL SSY, 1000 μ g/mL PAR, 2 μ g/mL CAR and 20 μ g/mL MP. b) Chromatogram of oral Suspension I, c) Chromatogram of oral Suspension II.

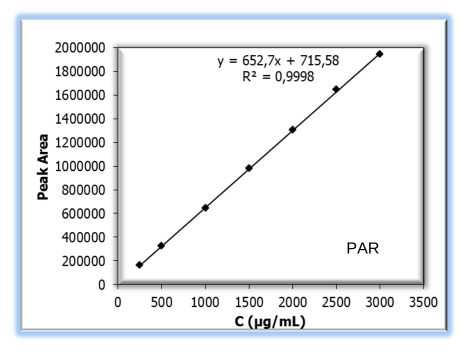
Method Validation

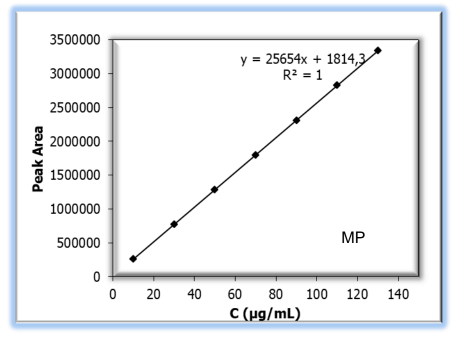
Selectivity: Selectivity was evaluated by the peak purity test using PDA detector. According to the results obtained, peak purity values were higher than 0.9990. Also no interferences were detected at retention times of the studied compounds in sample solutions, which shows that the developed method is selective (Figure 2).

Linearity: For linearity, seven different concentrations were chosen taking into account suspension contents. Hence, concentrations of the solutions were PAR 250 to 3000 μ g/mL, MP 10 to 110 μ g/mL, SSY 1 to 12 μ g/mL and CAR 1 to 7 μ g/mL. Each concentration of standard solutions was analyzed in triplicate and the mean values of peak areas were calculated and used

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for the calibration graph. Calibration curves of the studied compounds obtained by the proposed method can be seen in Figure 3. The linear regression equations for PAR, MP, SSY and CAR were found to be y = 652.7x - 715.58, y = 25654x + 1814.3, y = 10558x + 1532.5 and y = 15052x + 1742.6, respectively. The regression cofficients (R²) were found to be higher than 0.999, which indicates that the method has an acceptable degree of linearity.





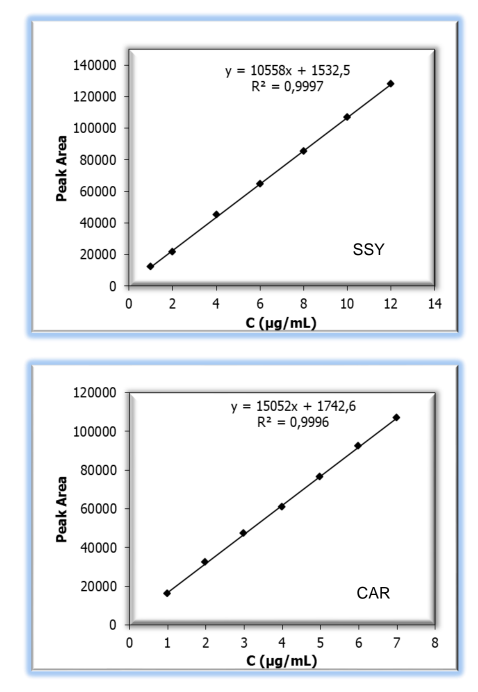


Figure 3. Calibration curves of PAR, MP, SSY and CAR.

Limits of Detection and Quantification (LOD and LOQ): Limits of detection (LOD) were calculated at a signal-to-noise ratio (S/N) of 3. Limits of quantification (LOQ) were calculated at a signal-to-noise ratio (S/N) of 10. The LOD was calculated to be 0.42, 0.20, 0.27 and 0.15 μ g/mL and the LOQ was calculated to be 1.35, 0.65, 0.80 and 0.46 μ g/mL for PAR, MP, SSY and CAR, respectively.

Precision: Precision of the method is assessed by the estimate of the relative standard deviation (RSD) with respect to both repeatability and intermediate precision. For repeatability, three different concentrations of standard solutions were analyzed in triplicate on the same day.

Intermediate precision (n=4) was performed on three different days. Acceptable RSD% values (<3.38% RSD) as shown in Table 2 were obtained.

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Compound	Actual	Repeatability	Intermediate precision
	Concentration	Mean ± %RSD	Mean ± %RSD
	(µg/mL)	(n =3) (µg/mL)	(n=4) (µg/mL)
PAR	1000	990.10 ± 1.52	993.75 ± 1.49
	1500	1479.33 ± 0.95	1487.00 ± 1.66
	2000	1992.67 ± 1.65	1991.75 ± 1.26
MP	20	20.47 ± 1.97	19.19 ± 2.45
	30	29.5 ± 1.70	29.83 ± 1.82
	40	39.6 ± 2.91	40.20 ± 2.96
SSY	2	2.07 ± 3.38	1.98 ± 2.80
	4	3.96 ± 3.26	4.08 ± 2.27
	6	5.97 ± 1.94	6.06 ± 2.27
CAR	1	0.98 ± 2.00	1.05 ± 2.52
	2	1.90 ± 2.80	1.95 ± 3.05
	3	3.10 ± 2.50	3.15 ± 2.65

Table 2. Repeatability and intermediate precision	n values of PAR, MP, SSY and CAR.
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Accuracy

Accuracy of the method was expressed as % recovery obtained by standard addition method at three different levels. Good recovery data for accuracy are displayed in Table 3. The recovery values obtained also imply that there is no matrix effect on the determination of analytes by the proposed method.

ion Amount Added (μg/mL)	Amount Found (μg/mL) ± %RSD (n=5) 487.0 ± 1.03	Mean Recovery (%)
	487.0 ± 1.03	
1000		97.40
1000	1002.67 ± 0.75	100.27
1500	1496 ± 0.68	99.73
10	9.98 ± 0.76	99.80
20	19.80 ± 1.35	99.00
30	29.12 ± 2.00	97.07
2	1.97 ± 2.18	98.50
4	3.95 ± 2.50	98.75
6	6.05 ± 2.20	100.83
1	1.05 ± 2.44	105.00
2	1.98 ± 1.96	99.00
	3.08 ± 2.15	
	20 30 2 4 6 1	$\begin{array}{cccc} 20 & 19.80 \pm 1.35 \\ 30 & 29.12 \pm 2.00 \\ \hline 2 & 1.97 \pm 2.18 \\ 4 & 3.95 \pm 2.50 \\ \hline 6 & 6.05 \pm 2.20 \\ \hline 1 & 1.05 \pm 2.44 \\ \hline 2 & 1.98 \pm 1.96 \\ \end{array}$

Table 3. Accuracy studies of PAR, MP, SSY and CAR.

Application of the Method to Commercial Oral Suspensions

After method optimization and validation, the developed method was successfully applied for the simultaneous determination of PAR, MP, SSY and CAR in oral suspension samples produced by the same pharmaceutical company. The quantitative results of the analysis are summarized in Table 4. Found values close to 100% demonstrate the applicability of the method for control of the liquid formulations in quality control laboratories. So these formulations containing the studied compounds can be analyzed with the same HPLC method in a short time.

	Ingredients	Labeled amount (mg/5 mL)	Amount found (mg/5 mL)	Recovery (%)
	PAR	250	250.6	100.24
Commercial Oral Suspension I	MP	4	4.05	101.32
	SSY	-	0.35	-
Commercial Oral Suspension II	PAR	120	121.2	101.0
	MP	5	5.11	102.29
	CAR	-	0.075	-

Table 4. Analysis of marketed samples by the HPLC method.

CONCLUSION

A novel, facile, rapid, and efficient reversed phase HPLC-PDA method without any extraction stage was developed for the simultaneous quantification of PAR, MP, SSY and CAR in oral suspensions which the ingredients are present in variable concentrations. The most important advantage of this method is to analyze PAR, MP, SSY and CAR at the same time as there is no method in the literature for simultaneous determination of these compounds. The developed method has a good resolution between all analytes with a short analysis time below 5 min. In addition, according to validation study results, this method is linear, precise, accurate, sensitive, and selective. So the proposed method can be used for routine analysis of these compounds in similar pharmaceutical products.

COMPLIANCE WITH ETHICAL STANDARDS

The authors declare that there is not an unethical situation.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

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Declared none.

REFERENCES

1. Sorouraddin MH, Saadati M, Mirabi F. Simultaneous determination of some common food dyes in commercial products by digital image analysis. Journal of Food and Drug Analysis. (2015); 23(3): 447-452.

2. Grosa G, Del Grosso E, Russo R, Allegrone G. Simultaneous, stability indicating, HPLC-DAD determination of guaifenesin and methyl and propyl-parabens in cough syrup. Journal of Pharmaceutical and Biomedical Analysis. 2006;41:798-803.

3. Baranowska I, Wojciechowska I, Solarz N, Krutysza E. Determination of preservatives in cosmetics, cleaning agents and pharmaceuticals using fast liquid chromatography. Journal of Chromatographic Science. 2013;52:88-94.

4. Yang J, Li Y, Gong W, Wang C, Liu B, Sun C. Simultaneous determination of six parabens in foods by matrix liquid-phase dispersion extraction combined with high-performance liquid chromatography. Food Analytical Methods. 2014;7:1693-702.

5. Routledge EJ, Parker J, Odum J, Ashby J, Sumpter JP. Some alkyl hydroxy benzoate preservatives (parabens) are estrogenic. Toxicology and Applied Pharmacology. 1998;153: 12-19.

6. Shanmugam G, Ramaswamy BR, Radhakrishnan V, Tao H. GC–MS method for the determination of paraben preservatives in the human breast cancerous tissue, Microchemical Journal. 2010; 96: 391–396.

7. Wang W, Wang Y, Zhang J, Chu Q, Ye J. Simultaneous determination of electroactive and non-electroactive food preservatives by novel capillary electrophoresis with amperometric detection. Analytica Chimica Acta. 2010; 678:39-43.

8. Pagáčiková D, Lehotay J. Determination of synthetic colors in meat products using highperformance liquid chromatography with photodiode array detector. Journal of Liquid Chromatography & Related Technologies. 2015;38:579-583.

9. Martin F, Oberson JM, Meschiari M, Munari C. Determination of 18 water-soluble artificial dyes by LC–MS in selected matrices. Food Chemistry. 2016;197:1249-1255.

10. Davletbaeva P, Chocholouš P, Bulatov A, Šatínský D, Solich P. Sub-1 min separation in sequential injection chromatography for determination of synthetic water-soluble dyes in pharmaceutical formulation. Journal of Pharmaceutical and Biomedical Analysis. 2017; 143: 123-129.

11. The European Parliament, Regulation (EC) No 1333/2008 of the European Parliament and of The Council of 16 December 2008 on food additives, Official Journal of the European Union, No 1333/2008.

12. EFSA (European Food Safety Authority) (2009) The EFSA Journal 7: 1328-67.

13. Aksu Dönmez Ö, Aşçı B, Dinç-Zor Ş, Aslan Çakır A. Simultaneous quantitative analysis of ephedrine HCl, guaifenesin, and some synthetic additives in syrups by RP-HPLC using Box-Behnken design. Latin American Journal of Pharmacy. 2018; 37: 85-94 (2018).

14. Lokhnauth JK, Snow NH. Determination of parabens in pharmaceutical formulations by solid-phase microextraction-ion mobility spectrometry. Analytical Chemistry. 2005;77:5938-5946.

15. Rebbeck C, Hammond R, Wong J, Nair L, Raghavan N, Hepler D, Campbell W, Lynn, R. Solid-phase extraction and HPLC analysis of methylparaben and propylparaben in a concentrated antibiotic suspension. Drug Development and Industrial Pharmacy. 2006; 32:1095-1102.

16. Bosch ME, Sánchez AR, Rojas FS, Ojeda CB. Determination of paracetamol: Historical evolution. Journal of Pharmaceutical and Biomedical Analysis. 2006;42:291-321.

17. Dinç Ş, Aksu Dönmez Ö, Aşçı B, Bozdoğan AE. Chromatographic and chemometricsassisted spectrophotometric methods for the simultaneous determination of allobarbital, adiphenine hydrochloride, and paracetamol in suppository. Journal of Liquid Chromatography & Related Technologies. 2014;37:560-571.

18. Tzanavaras P, Karakosta T, Rigas P, Themelis D, Zotou A. Isocratic liquid chromatographic determination of three paraben preservatives in hygiene wipes using a reversed phase coreshell narrow-bore column. Open Chemistry. 2012;10:1459-1463.

19. Kumar S, Mathkar S, Romero C, Rustum AM. Development and validation of a single RP-HPLC assay method for analysis of bulk raw material batches of four parabens that are widely used as preservatives in pharmaceutical and cosmetic products. Journal of Chromatographic Science. 2011;49: 405-411.

20. Rovina K, Acung LA, Siddiquee S, Akanda JH, Shaarani SM. Extraction and analytical methods for determination of sunset yellow (E110)—a Review. Food Analytical Methods. 2017;10:773-787.

21. Rejczak T, Tuzimski T. Application of high-performance liquid chromatography with diode array detector for simultaneous determination of 11 synthetic dyes in selected beverages and foodstuffs. Food Analytical Methods. 2017;10: 3572-3588.

22. Khanavi M, Hajimahmoodi M, Ranjbar AM, Oveisi MR, Ardekani MRS, Mogaddam G. Development of a green chromatographic method for simultaneous determination of food colorants. Food Analytical Methods. 2012;5:408-415.

23. Yamjala K, Nainar MS, Ramisetti NR. Methods for the analysis of azo dyes employed in food industry–a review. Food Chemistry. 2016;192:813-824.

24. Hasan N, Chaiharn M, Toor UA, Mirani ZA, Sajjad G, Sher N, Aziz M, Siddiqui FA Development, validation and application of RP-HPLC method: Simultaneous determination of antihistamine and preservatives with paracetamol in liquid formulations and human serum. Open Medicinal Chemistry Journal, (2016); 10:33-43.

25. Abd El-Hay SS, Mohram MS. Development and validation of new RP-HPLC method for simultaneous determination of methyl and propyl parabens with levetiracetam in pure form and pharmaceutical formulation. Chromatography Research International. 2016.

26. Weshahy SAEF, Yaaqob MS, Morcos MN, Hassan DW, Youssef NF. Simultaneous determination of levodropropizine, methylparaben, and propylparaben in oral co-formulated syrup by RP-HPLC method. Journal of the Chilean Chemical Society. 2015;60: 2729-2733.

27. Ali MS, Ghori M, Khatri AR. Stability indicating simultaneous determination of domperidone (DP), methylparaben (MP) and propylparaben by high performance liquid chromatography (HPLC). Journal of Pharmaceutical and Biomedical Analysis. 2006; 41:358-365.

Zor,Dönmez. JOTSCA. 2018;5(2) 763-74

28. International Conference on Harmonization (ICH) Q2 (R1): Validation of Analytical Procedures-Test and Methodology, Geneva, Switzerland, 2005.