# Rapid Detection of Dimethyl Yellow Dye in Curry by Liquid Chromatography-Electrospray-Tandem Mass Spectrometry

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## Abstract

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An accurate and rapid method, was devised for the identification and quantitation of dimethyl yellow dye in curry, based on liquid chromatography-tandem mass spectrometry interfaced with electrospray. Mass spectral acquisition was done in positive ion mode applying two fragmentation transitions to provide a high degree of selectivity. The extraction system provided a very high recovery (100.0% to 105.8%) and good results were obtained for the limit of detection (5  $\mu$ g/kg) and limit of quantitation (16  $\mu$ g/kg). The applicability of the method to identifing and quantifing the unauthorised dimethyl yellow dye in curry was demonstrated.

Keywords: azo dyes; dimethyl yellow; curry; LC-ESI-/MS/MS

Azo type dyes are most frequently studied for analytical food control due to their toxicity risk; the chromophoric azo group, under certain conditions, can be reduced to form confirmed or suspected carcinogenic aromatic amines (AHLSTROM et al. 2005). The commercialisation of food contaminated with hot chilli products coloured with azo dyes Sudan I, II, III, and IV led the EU in 2003 to adopt emergency measures (Commission Decision 2003). In April 2009, the Agence Fédérale pour la Sécurité de la Chaîne Alimentaire (AFSCA 2009a) in Belgium recalled curry contaminated with 4-(N,N-dimethylamino)-azobenzene, currently named Dimethyl yellow (C14H15N3, CAS No. 60-11-7). This dye has been classified as a category 2B carcinogen to humans by the International Agency for Research on Cancer. The Belgium alert was diffused in April 2009, the Rapid Alert System for Food and Feed (RASFF) determined the unauthorised Dimethyl yellow (> 0.50 mg/kg) to be in Madras curry powder from India.

Various analytical techniques and instruments have been employed for the analysis of synthetic co-

lours, such as spectrophotometry (NI & GONG 1997; SAYAR & ÖZDEMIR 1998; BERZAS et al. 1999; NI et al. 2001, 2009; VIDOTTI et al. 2005; PENMAN et al. 2006), stripping voltammetry (ALGHAMDI 2005a,b), capillary electrophoresis (Suzuкi et al. 1994; Liu et al. 1995; MASAR & KANIANSKY 1996; MASAR et al. 1996; Kuo et al. 1998; Chou et al. 2002; Del Giovine & PICCIOLI BOCCA 2003; DOSSI et al. 2007), high performance liquid chromatography with UV-VIS, UV-DAD or MS detectors (GREENWAY et al. 1992; Chen et al. 1998; Garrigos et al. 2002 Prado & GODOY 2002; KIRSCHBAUM et al. 2003; CALBIANI et al. 2004a,b; GARCIA-FALCON & SIMAL-GANDARA 2005; GIANOTTI et al. 2005; ZHANG et al. 2005; VI-DOTTI et al. 2006; KIRSCHBAUM et al. 2006; MA et al. 2006; Ertaș et al. 2007; Sun et al. 2007; Alves et al. 2008; NOGUEROL-CAL et al. 2008; PEREIRA et al. 2008; HARTIG et al. 2009).

The extraction recovery of various azo-dyes depends strongly on the analysed matrices and for banned azo-dyes is it very important to use analytical methods with very low limit of detection (LOD). In all cases, the LOD by HPLC/MS/MS for azo-dyes results as lower than those obtained by HPLC/UV-Vis. In addition, MS detection produces more specific structural information. This paper proposes very specific analytical conditions for the rapid detection of dimethyl yellow dye in spices with a LOD lower than the level generally proposed by official organisations for food safety. For example, the document published by AFSCA concerning the protection measures for dimethyl yellow cites the LOD =  $15 \mu g/kg$  as the analytical value useful to confirm the absence of this illegal dye (AFSCA 2009b).

# MATERIAL AND METHODS

**Chemicals.** HPLC-grade acetonitrile  $(CH_3CN)$  was obtained from J.T. Baker (Deventer, the Netherlands). Ethanol and ammonium formate were obtained from Merck (Darmstadt, Germany).

**Standards.** Dimethyl yellow standard (CAS Number 60-11-7) was obtained from Sigma Aldrich, Milan, Italy (Cod. Prod. 33129). A standard stock solution (680 mg/l) was prepared by dissolving 68 mg of dimethyl yellow standard powder in 100 ml CH<sub>3</sub>CN. First dilution (solution A): 1 ml stock solution was diluted to 100 ml with CH<sub>3</sub>CN to obtain a 6.80 mg/l solution. Second dilution (solution B): 2 ml solution A was diluted to 10 ml with CH<sub>3</sub>CN to obtain a 1.36 mg/l solution.

*Spice enrichment with dimethyl yellow.* The spice used to verify the response of the adopted extraction method was curry; curry is one of the most complex matrices and potentially offers the most interference in analytical studies. Three

aliquots (2 g each) of the same curry sample, purchased in the market and not containing dimethyl yellow, were enriched with 10  $\mu$ l, 100  $\mu$ l, and 250  $\mu$ l of standard solution B, reaching dimethyl yellow concentrations of 6.8, 68.0, and 170.0  $\mu$ g/kg, respectively.

*Extraction method.* The spice (approx. 2 g of enriched curry) was treated with 50 ml  $CH_3CN$  in a 75 ml centrifuge tube. The mixture was shaken for 4 h (GFL 3006) and sonicated for 5 min in each hour. The solution was centrifuged for 10 min at 4000 rpm, filtered, and 10 µl f the extraxt were injected into the LC/MS/MS system.

LC-ESI (+)-MS/MS method. The experiments were carried out with a LC-ESI(+)-MS/MS system using an Agilent 1200 Series HPLC coupled to an Applied Biosystems API 2000 mass spectrometer (Toronto, Canada) equipped with a Turbo lonSpray Source (ESI). Chromatographic condi*tions* – A stainless steel column (Macherey-Nagel, Bethlehem, USA) EC/2 Nucleodur 100-5 C18 (150 mm × 2 mm, 110 Å pore diameter, 5.0 μm particle size) was used. Water/5mM ammonium formate (Phase A) and methanol/5mM ammonium formate (Phase B) were used for the mobile phase and the elution gradient was: from 40% to 85% B in 5 min, to 10% B in 5 min, then 100% B for 10 min and return to 40% B in 15 minutes. ESI(+) MS/MS method - To establish the appropriate multiple reaction monitoring (MRM) conditions, a test solution was produced by diluting the stock solution 1:1 with mobile phase (1:1 water/methanol, 5mM ammonium formate), to a concentration of 340 µg/l. The test solution was injected by continu-



Figure 1. (a) Mass spectrum of dimethyl yellow obtained with a single quadrupole, showing only one chromatographic peak corresponding to ion  $[M + H]^+ m/z = 226$ ; (b) daughter ions obtained with multiple reaction monitoring from ion precursor m/z = 226 (m/z = 105.2 and m/z = 133.1)



Figure 2. LC-ESI-MS/MS traces obtained from curry powder spiked with standard dimethyl yellow at three concentrations (6.8, 68.0, and 170.0  $\mu$ g/kg): (a) daughter ion *m*/*z* = 105.2; (b) daughter ion *m*/*z* = 133.1

ous infusion at 5 µl/min in +Q1 mode and in +MS<sup>2</sup> mode. The values of DP – Declustering Potential (25 V), FP – Focusing Potential (400 V), EP – Entrance Potential (12 V), CEP – Cell Exit Potential (9 V), CE – Collision Energy (27 V), and CXP – Cell Exit Potential (3 V) were adopted to obtain the 226.0  $\rightarrow$  105.2 transition. The same values, with the exception of DP (22 V) and CE (45 V), were used to obtain the 226.0 > 133.1 transition.

#### **RESULTS AND DISCUSSION**

The mass spectrum of dimethyl yellow, obtained with a single quadrupole, shows only one chromatographic peak corresponding to ion  $[M+H]^+$ at m/z = 226 (Figure 1a). The daughter ions produced with MRM from ion precursor m/z = 226are shown in Figure 1b (m/z = 105.2 and m/z =133.1). Figure 2a shows three peaks, corresponding to daughter ion m/z = 105.2, obtained from curry spiked with standard dimethyl yellow at three con-



centrations (6.8, 68.0, and 170.0 µg/kg); the peaks corresponding to daughter ion m/z = 133.1 appear in Figure 2b. The linear calibration curve shown in Figure 3 was obtained from the areas corresponding to the first transition (m/z = 105.2).

When considering the toxicity risk of dimethyl yellow in foods, the quantitative determination is not as important as its identification at a high level of sensitivity. More important, on the contrary, is it to know the LOD and LOQ levels. To determine these values, the calibration curve (Figure 3,  $R^2 = 0.9998$ ) was produced and LOD (5 µg/kg) and LOQ (16 µg/kg) were calculated from the intercept and slope as reported in Table 1. Concentrations higher than 170.0 µg/kg were not considered for the calibration curve because the quantitative determination of dimethyl yellow is not the aim of this paper for the reason explained before (not tolerance in food).

Therefore, the precision data were evaluated to confirm the efficiency of the detection method and the relative standard deviation for repeatability (RSDr) was determined from NMKL procedure

Figure 3. Calibration curve obtained using the area counts corresponding to the first transition  $(m/z \ 226 \rightarrow 105.2)$  at three concentrations (6.8, 68.0, and 170.0 µg/kg

	mg/kg(x)	Area counts $(y_i)$	$y_t = b \times x + a$	$y_i - y_t$		$(y_i - y_t)^2$
1	0.007	2 600	2 679	-79		6 238
2	0.068	8 070	7 944	126		15 930
3	0.170	16 700	16 747	-47		2 231
		a = 2.074.8	<i>b</i> = 86308.3			
	$S(y_i - y_t)^2$	$S_{y/x}$	LOD (mg/kg)	LOQ (mg/kg)	R	$R^2$
	24 399	156	0.005	0.016	0.9999	0.9998

Table 1. Limit of detection (LOD) and limit of quantitation (LOQ) of the LC-ESI-MS/MS method for curry samples calculated from the calibration curve shown in Figure 3

 $S_{v/x} = \sqrt{[(\Sigma(y_i - y_i)^2/(n-2)]]};$  LOD = 3 ×  $S_{v/x}/b$  limit of detection; LOQ = 9 ×  $S_{v/x}/b$  limit of quantification





INCR\_CURRY\_GIALLO DIMETILE\_0.017 MG/KG\_PROVA\_1 - ... Area: 3520. commts Height: 2.14e+002 cps RT: 11.2 min



INCR\_CURRY\_GIALLO DIMETILE\_0.017 MG/KG\_PROVA\_2 - GIALLO Area: 3650. counts Height: 1.80e+002 tps RT: 11.3 min



Figure 4. LC-ESI-MS/MS traces obtained from: (a) curry powder (blank matrix), (b) and (c) duplicate curry powder samples spiked with 17.0  $\mu$ g/kg dimethyl yellow

No. 5 (Nordic Committee on Food Analysis 1997). The suggested method was applied to duplicate analyses of dimethyl yellow in each sample and was applied to 10 different curry samples enriched with different amonts of dimethyl yellow, measured on different days. The RSDr value for 10 samples of curry was 2.7%.

Considering the calibration curve was produced from the data obtained by matrix enrichment, the recovery corrections are included in the method. To verify the high recovery also at lower concentrations of dimethyl yellow near the LOQ value, two identical additions were made to the curry sample not containing dimethyl yellow: 25 µl of standard solution B was added to 2 g of matrix, resulting in a 17 µg/kg concentration of dimethyl yellow. Figure 4 shows three chromatograms, corresponding to the blank matrix (Figure 4a) and the duplicate samples of the enriched matrix (Figures 4b and c, with area values of 3520 and 3650). The recovery values correspond to 100.0% and 105.8%, respectively, and the efficiency in the identification of dimethyl yellow at the concentration levels of the LOQ order were confirmed.

# CONCLUSION

The LC-ESI(+)-MS/MS fast method proposed seems to be adequate to detect the banned dimethyl yellow azo-dye since the detection limits are lower than obtained with other methods. The extraction method adopted for curry, the matrix showing more interferences than other spices, produced high recovery and good repeatability data; more over, the LOD and LOQ values proved to provide unambiguous identification as requested for food matrices where the presence of the specific azo dye is banned. The experiences were developed in particular to ameliorate the data quality in correspondence with LOD and LOQ levels, the quantitative determination at higher levels being not useful for food safety control.

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### References

- AFSCA (2009): Communique AFSCA: rappel de curry sous forme d'épice ou de mélange d'épices 01 avril 2009. Agence Fédérale pour la Sécurité de la Chaîne Alimentaire. Available at http://www.favv-afsca.be/ communiquesdepresse/documents/2009-04-01 curry fr.pdf (accessed 2 April 2009)
- AFSCA (2009): Contamination au jaune de methyl. Agence Fédérale pour la Sécurité de la Chaîne Alimentaire. Available at http://www.afsca.be/denreesalimentaires/jaunemethyle/ (accessed on 8 March 2010)
- ALVES S.P., BRUM D.M., ANDRADE E.C.B. DE NETTO A.D.P. (2008): Determination of synthetic dyes in foodstuff by high performance liquid chromatography with UV-DAD detection. Food Chemistry, **107**: 489–496.
- ALGHAMDI A.H. (2005a): A square-wave adsorptive stripping voltammetric method for the determination of Amaranth, a food additive dye. Journal of AOAC International, **88**: 788–793.
- ALGHAMDI A.H. (2005b): Determination of Allura red in some food samples by adsorptive stripping voltammetry. Journal of AOAC International, **88**: 1387–1393.
- AHLSTROM L.H., SPARR ESKILSSON C., BJORKLUND E. (2005): Determination of banned azo-dyes in consumer goods. Trends in Analytical Chemistry, 24: 49–56.
- BERZAS J.J., RODRIGUEZ-FLORES J., VILLASENOR LLE-RENA M.J., RODRIGUEZ FARINAS N. (1999): Spectrophotometric resolution of ternary mixtures of Tartrazine, Patent Blue V and Indigo Carmine in commercial products. Analytica Chimica Acta, **391**: 353–364.
- CALBIANI F., CARERI M., ELVIRI L., MANGIA A., PISTARA L., ZAGNONI I. (2004a): Development and in-house validation of a liquid chromatography – electrospraytandem mass spectrometry method for the simultaneous determination of Sudan I, Sudan II, Sudan III, and Sudan IV in hot chilli products. Journal of Chromatography A, **1042**: 123–130.
- CALBIANI F., CARERI M., ELVIRI L., MANGIA A., ZAG-NONI I. (2004b): Accurate mass measurements for the confirmation of Sudan azo-dyes in hot chilli products

by capillary liquid chromatography-electrospray tandem quadrupole orthogonal-acceleration time of flight mass spectrometry. Journal of Chromatography A, **1058**: 127–135.

- CHEN Q., MOU S., HOU X., RIVIELLO J.M., NI Z. (1998): Determination of eight synthetic food colourants in drinks by high-performance ion chromatography. Journal of Chromatography A, **827**: 73–81.
- CHOU S.S., LIN Y.H., CHENG C.C., HWANG D.F. (2002): Determination of synthetic colours in soft drinks and confectioneries by micellar electrokinetic capillary chromatography. Journal of Food Science, **67**: 1314–1318.
- Commission Decision of 20 June 2003 on emergency measures regarding hot chilli and hot chilli products (notified under document number C (2003) 1970) (2003/460/EC). Official Journal of the European Union L 154 (21 June 2003): 114.
- DEL GIOVINE L., PICCIOLI BOCCA A. (2003): Determination of synthetic dyes in ice-cream by capillary electrophoresis. Food Control, **14**: 131–135.
- Dossi N., Toniolo R., Pizzariello A., Susmel S., Perennes F., Bontempelli G. (2007): A capillary electrophoresis microsystem for the rapid in-channel amperometric detection of synthetic dyes in foods. Journal of Electroanalytical Chemistry, **601**: 1–7.
- ERTAȘ E., ÖZER H., CESARETTIN A. (2007): A rapid HPLC method for determination of Sudan dyes and Para Red in red chilli pepper. Food Chemistry, **105**: 756–760.
- GARCIA-FALCON M.S., SIMAL-GANDARA J. (2005): Determination of food dyes in soft drinks containing natural pigments by liquid chromatography with minimal clean-up. Food Control, **16**: 293–297.
- GARRIGOS M.C., RECHE F., MARIN M.L., JIMENEZ A. (2002): Determination of aromatic amines formed from azo colorants in toy products. Journal of Chromatography A, **976**: 309–317.
- GIANOTTI V., ANGIOI S., GOSETTI F., MARENGO E., GENN-ARO M.C. (2005): Chemometrically assisted development of IP-RP-HPLC and spectrophotometric methods for the identification and determination of synthetic dyes in commercial soft drinks. Journal of Chromatography & Related Technologies, **28**: 923–937.
- GREENWAY G. M., KOMETA N., MACRAE R. (1992): The determination of food colours by HPLC with on-line dialysis for sample preparation. Food Chemistry, **43**: 137–140.
- HARTIG L., HOENICKE K., VON CZAPLEWSKI K (2009): Detection von 6 sudan dyes, dimethyl yellow and para red in spices and sauce with HPLC/MS/MS. Available at http://www3.appliedbiosystems.com/cms/groups/

psm\_marketing/documents/generaldocuments/cms\_ 042197.pdf (accessed 1 June 2009)

- KIRSCHBAURN J., KRAUSE C., PFALZGRAF S., BROCKNER H. (2003): Development and evaluation of an HPLC-DAD method for determination of synthetic food colorants. Chromatographia, **57**(S): 115–119.
- KIRSCHBAUM J., KRAUSE C., BRÜCKNER H. (2006): Liquid chromatographic quantification of synthetic colorants in fish roe and caviar. European Food Research and Technology, **222**: 572–579.
- KUO K.L., HUANG H.Y., HSIEH Y.Z. (1998): High-performance capillary electrophoretic analysis of synthetic food colourants. Chromatographia, **47**: 249–256.
- LIU H., ZHU T., ZHANG Y., QI S., HUANG A., SUN Y. (1995): Determination of synthetic colourant food additives by capillary zone electrophoresis. Journal of Chromatography A, **718**: 448–453.
- MA M., LUO X., CHEN B., SU S., YAO S. (2006): Simultaneous determination of water-soluble and fat-soluble synthetic colorants in foodstuff by high-performance liquid chromatography-diode array detection – electrospray mass spectrometry. Journal of Chromatography A, **1103**: 170–176.
- MASAR M., KANIANSKY D. (1996): Determination of synthetic dyes in food products by capillary zone electrophoresis in a hydrodynamically closed separation compartment. Journal of Capillary Electrophoresis, **3**: 165–171.
- MASAR M., KANIANSKY D., MADAJOVA V. (1996): Separation of synthetic food colourants by capillary zone electrophoresis in a hydrodynamically closed separation compartment. Journal of Chromatography A, **724**: 327–336.
- NI Y., GONG X.F. (1997): Simultaneous spectrophotometric determination of mixtures of food colorants. Analytical Chimica Acta, **354**: 163–171.
- NI Y., QI M., КОКОТ S. (2001): Simultaneous spectrophotometric determination of ternary mixture of Tartrazine, Sunset Yellow and Ponceau 4R by H-point standard addition method. Analytical Letters, **34**: 2585–2596.
- NI Y., WANG Y., ΚΟΚΟΤ S. (2009): Simultaneous kitetic spectrophotometric analysis of five synthetic food colorants with the aid of chempmetrics. Talanta, **78**: 432–441.
- NOGUEROL-CAL R., LÓPEZ-VILARINO J.M., FERNÁN-DEZ-MARTINEZ G., BARRAL-LOSADA L., GONZÁLEZ-RODRIGUEZ M.V. (2008): High-performance liquid chromatography analysis of ten dyes for control of

safety of commercial articles. Journal of Chromatography A, **1179**: 152–160.

- Nordic Committee on Food Analysis (1997): Estimation and expression of measurement uncertainty in chemical analysis. NMKL Procedure No. 5.
- PENMAN K.G., HALSTEAS C.W., MATTHIAS A., DE Voss J.J., STUTHE J.M.U., BONE K.M., LEHMANN R.P. (2006): Bilberry adulteration using the food dye amaranth. Journal of Agricultural and Food Chemistry, **54**: 7378–7382.
- PEREIRA ALVES S., MARES BRUM D., CASTELLO BRANCO DE ANDRADE E., DUARTE PEREIRA NETTO A. (2008): Determination of synthetic dyes in selected foodstuffs by high performance liquid chromatography with UV-DAD detection. Food Chemistry, **107**: 489–496.
- PRADO M.A., GODOY H.T. (2002): Validation of the methodology to determine synthetic dyes in food and beverages by HPLC. Journal of Chromatography & Related Technologies, **25**: 2445–2472.
- SAYAR S. ÖZDEMIR Y. (1998): First-derivative spectrophotometric determination of ponceau 4R, sunset yellow and tartrazine in confectionery products. Food Chemistry, **61**: 367–372.
- SUN H.W., WANG F.C., L. AI L.F. (2007): Determination of banned 10 azo-dyes in hot chili products by gel permeation chromatography-liquid chromatography – electrospray ionization-tandem mass spectrometry. Journal of Chromatography A, **1164**: 120–128.
- SUZUKI S., SHIRAO M., AIZAWA M., NAKAZAWA., SASA K., SASAGAWA H. (1994): Determination of synthetic food dyes by capillary electrophoresis. Journal of Chromatography A, **680**: 541–547.
- VIDOTTI E.C., CANCINO J.C., OLIVEIRA C.C., ROLLEM-BERG M.D.C.E. (2005): Simultaneous determination of food dyes by first derivative spectrophotometry with sorption onto polyurethane foam. Analytical Science, **21**: 149–153.
- VIDOTTI E.C., COSTA W.F., OLIVEIRA C.C. (2006): Development of a green chromatographic method for determination of colorants in food samples. Talanta, **68**: 516–521.
- ZHANG Y.P., ZHANG Y.J., GONG W.J., COPALAN A.I., LEE K.P. (2005): Rapid separation of Sudan dyes by reverse-phase high performance liquid chromatography through statistically designed experiments. Journal of Chromatography A, **1098**: 183–187.

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