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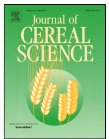
First report on the presence of Alloxan in bleached flour by Lc-Ms/Ms method

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FIRST REPORT ON THE PRESENCE OF ALLOXAN IN BLEACHED FLOUR BY LC MS/MS METHOD

- 3
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17	ABSTRACT
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19	In this work the presence of Alloxan in bread, pastry and cake bleached flour was investigate in order
20	to verify possible risk for consumers related to the use of chemicals for flour bleaching. A selective
21	UHPLC-MS/MS method has been developed and validated for the purpose. Alloxan is one of the
22	possible minor side products of oxidation after chemical bleaching of wheat flours, when several
23	chemical agents are used. One hundred and seventy-five flour samples were analyzed for Alloxan
24	determination. The validation of the method was performed in accordance with the ISO/IEC/EN 17025
25	for linearity, detection limit, quantification limit, accuracy, precision and ruggedness determination.
26	Satisfactory performances were obtained for the analyte, with a Limit of Detection (LOD) of 0.73 mg
27	kg ⁻¹ , a Limit of Quantification (LOQ) of 0.85 mg kg ⁻¹ and recovery values between 94% and 102%.
28	The present work report for the first time the presence of trace levels of Alloxan in 24% of the analyzed
29	samples, with mean values of 0,95±0,04 mg kg ⁻¹ . The presence of Alloxan was detected only in cake

30 flour samples. Further studies on toxicological levels of Alloxan are needed in order to evaluate 31 possible risks for consumers linked to the consumption of bakery products.

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Keywords: Flour; Alloxan; LC–ESI-MS/MS; validation procedure; bakery products

1. INTRODUCTION

Alloxan (2,4,5,6-tetraoxypyrimidine or 2,4,5,6-pyrimidinetetrone) is an oxygenated pyrimidine
synthesized by uric acid oxidation that can be found in hydrated form. Since 1943, it was found to
have toxicological effects on pancreatic β-cells leading to diabetogenic action, therefore (Jacobs, 1937;
Pincus, 2013) is commonly employee for the development of Type-I Diabetes Mellitus in animal
models. (Carvalho et al., 2003; Dunn et al., 1943; Goldner and Gomori, 1943; Lenzen and Panten,
1988; Saadia et al., 2005; Saleem Mir et al., 2013; Seifu et al., 2017; Shaw Dunn and Mcletchie, 1943;
Szkudelski, 2001; Webb, 1966).

As an analogous of the glucose, this molecule has two distinct effects on the pathology of diabetes: selective inhibition of glucose-induced insulin secretion by specific inactivation of glucokinase, (Dhanabal et al., 2007; Weaver et al., 1978; Webb, 1966; Zhou et al., 2017) and inducer of the formation of Reactive Oxygen Species (ROS), which can damage different cellular components through the oxidation of proteins, lipids and nucleic acids (Kim et al., 1994; Lenzen, 2008).

Alloxan also demonstrated to have a carcinogenic action in rats and fishes; furthermore, it can induce adenohypophysis cancer in mices (Suganuma et al., 1993). Mrozikiewiez et al. (1994) have found elevated levels of Alloxan in the blood of diabetic children with insulin-dependent diabetes mellitus, correlated to the onset of insulin-dependent diabetes mellitus (Mrozikiewicz et al., 1994). Alloxan is a minor product of the proteins oxidation, so, it may be produced during the bleaching processes of the alimentary flour for dough and colour improvement, becoming possible toxicant (Banu Shakila and Sasikala, 2012).

Flours obtained from freshly ground wheat have a pale yellow colour due to their carotenoid content. This flour produces sticky dough not easy to work and cook. During storage, the natural aging of the flour is due to a series of oxidative reactions involving carotenoids and sulfhydryl groups of proteins systems. The result is a white, soft and bulky flour, more suitable for the preparation of bakery products (Fennema, 1985).

To accelerate these natural processes, the food industry uses chemical methods (Joye et al., 2009) able to improve both the colour and the pasting properties. Bleaching agent commonly used are benzoyl peroxide, chlorine gas, chlorine dioxide, nitrosyl chloride and nitrogen oxides (Chittrakorn et al., 2014).

The chlorine dioxide improves the properties of dough and makes the flour less yellow. In the United States, chlorine and hypochlorites are considered safe compounds for food processing, in particular chlorine falls within the food additive list of the Food and Drug Administration (Fukayama et al., 1986). In other Countries, the chlorine dioxide and the chlorine gases have been banished because of their possible toxic effects. (Joye et al., 2009). These compounds, in fact, destroy the conjugated double bonds and oxidise the thiol groups of the gluten proteins; the chlorine processing involves the breakage of hydrogen bonds and the cleavage of peptide bonds; it degrade also aromatic amino acids. (Joye et al., 2009; Kulp et al., 1985); these series of oxidation reactions may modify many flour components (Thomasson et al., 1995) and lead to the formation of toxic products, such as Alloxan. (Fukayama et al., 1986; "Idaho Observer: Bleaching agent in flour linked to diabetes," 2005).

The aim of this work is to prove the presence of Alloxan in bleached flour, given the absence of data in literature on the relative toxicity of this molecule in bakery products. A selective UHPLC–MS/MS method using precolumn derivatization was developed for the determination of Alloxan in flours. The method was validated by an in-house validation protocol according to ISO/IEC/EN 17025.

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2. MATERIAL AND METHODS

82 2.1 Sample collection

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A total of 175 bleached flour samples were collected from manufacturers and local market of Sicily (Southern Italy). All the samples considered in this study were randomly collected by choosing different texture and size of granulation: 62 bread flour (slightly coarse), 55 pastry flour and 58 cake flour (smooth and fine), respectively. The flours used for the validation have been collected from a farm that produces flour for personal use with ancient means of production. Flour samples that were not subjected to bleaching process were used as blank samples.

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90 2.2 Chemicals and reagents

91 Acetonitrile and formic acid 99.9% (LC-MS grade), water (HPLC gradient grade), were supplied from 92 VWR (VWR International PBI Srl Milan, Italy). Alloxan, Alloxazine, o-phenylenediamine, 93 hydrochloride acid 1M and sodium hydroxide 1M were purchased from Sigma (Sigma-Aldrich, Milan, 94 Italy). Standard stock solutions of Alloxan and o-phenylenediamine were prepared in hydrochloride 95 acid 0.1M (concentration of 1 mg mL⁻¹). Standard stock solution of Alloxazine was prepared in sodium hydroxide 0.1M at the concentration of 1 mg mL⁻¹. These solutions were stored at 2 to 8 °C up 96 97 to 3 months. Working stock solutions of Alloxan and Alloxazine were prepared in formic acid 0.1% 98 (v/v) in water at a concentration of 10 µg mL⁻¹ and stored at 2 to 8 °C up to 30 days. PTFE filters of 99 0.45 µm were purchased from Chromacol LTD (Thermo Fisher, Waltham, Massachusetts, USA).

- 101 2.3 Sample preparation
- 102 2 g of the homogenized flour samples were weighed in a polypropylene centrifuge tube and spiked
- 103 with 200 μ l of Alloxan working solution at 10 μ g mL⁻¹ in HCl 0.1 M to obtain a concentration of 1 mg
- 104 kg⁻¹. Every sample was mixed and allowed to rest for 15 min. Subsequently, 10 mL of hydrochloride
- acid 0.1M were added in the tube and then mixed for 1 min.
- The tube was vigorously centrifuged for 10 min at 3500 rpm; the supernatant was collected in a 50 mL
 polypropylene centrifuge tube that was filtered with filters of 0.45 μm.
- A 0.5 mL aliquot was added to 1.5 mL of 0.1% aqueous formic acid solution. This solution was spiked with 2 ml of o-phenylenediamine at 1 mg mL⁻¹. After a gentle stirring, an aliquot of 1 ml was transferred into vials and set at the appropriate temperature of 25°C for 24h, prior to LC–MS/MS analysis.
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113 2.4 Chromatographic conditions

114 LC analysis was carried out through a Thermo Fischer UHPLC system (Thermo Fisher Scientific, 115 California, U.S.A.) composed by an ACCELA 1250 LC pump and an ACCELA autosampler (Thermo 116 Fisher Scientific, California, U.S.A.). Chromatographic separation was obtained using a Thermo 117 Scientific Hypersil Gold PFP - UHPLC column (100mm x 2.1mm; 1.9µm). The LC eluents were water 118 (A) and acetonitrile (B) everyone containing 0.1% (v/v) of formic acid. The gradient (Table 1) was 119 initiated with 85% eluent A for 0.5 min, continued with linear variation to 5% A at 2 min; this 120 condition was maintained for 1 min. The system returned to 85% A in 0.5 min and was re-equilibrated 121 for 3.5 min. The column temperature was 30° C and the sample temperature was kept at 25° C. The 122 flow rate was 0.3 mL min⁻¹ and the injection volume 5 μ L.

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124 2.5 MS conditions

125 The mass spectrometer was a triple quadrupole TSQ Vantage (Thermo Fisher Scientific, California, 126 U.S.A.) in positive electrospray ionization mode (ESI+). Product ion scans of analyte was performed 127 by direct infusion (10μ l min⁻¹) of 1 µg mL⁻¹ Alloxazine standard solutions with the built-in syringe 128 pump through a T-junction, mixing with the blank column eluate (200μ l min⁻¹).

- ESI parameters optimized were as follows: capillary voltage, 4.5kV; capillary temperature, 310 °C; vaporizer temperature, 150 °C; sheath and auxiliary were fixed at 30 and 15 (arbitrary unit),
- respectively. The collision gas was argon at 1.5 mTorr and peak resolution of 0.7 FWHM was used on
- 132 Q1 and Q3. The scan time for each monitored transition was 0.05 s and the scan width was 0.05 m/z.
- 133 The collision energy parameters associated with the precursor and the product ions are given in Table

134 2. Acquisition data were recorded and elaborated using Xcalibur TM version 2.1.0.1139 software from Thermo Fisher Scientific (California, U.S.A.). The tune of the MS conditions for Alloxazine standard 135 136 was performed by direct infusion of 1 µg mL⁻¹ solution with the built-in syringe pump. It was be found 137 that the precursor ion with the most abundant signal are composed by the adduct [M+H+] in 138 electrospray positive mode. After, the chromatographic conditions were optimized by several 139 injections of Alloxazine standard solution at the concentration of 0.1 µg mL⁻¹ in order to test different 140 combinations of mobile phases. Then we found the best gradient condition, reported in the 141 experimental section of this paper, for the best symmetry and resolution of the peak. The spectrometric 142 determination was performed in MRM mode in order to obtain a better selectivity and sensitivity.

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144 2.6 Validation procedure

The method was validate by an in-house model, including determination of linearity, specificity, recovery, limit of detection (LOD), limit of quantification (LOQ), precision (repeatability and the within-laboratory reproducibility), accuracy and ruggedness. The validation was performed according to ISO/IEC/EN 17025 (2000). For the estimation of the validation parameters, we chose flour samples from companies operating in Sicily (Italy).

To verify the presence of Alloxan in the spiked sample, after extraction and derivatization, we compared the retention time and the relative abundance of the fragment with the signal of Alloxazine standard. The interpolation of the signal generated by the base peak in the calibration curve was used for the quantitative determination.

154 The instrumental linearity was calculated constructing calibration curve using Alloxazine commercial 155 standard; it represents the final derivatization product with o-phenylenediamine, which occurs at the 156 end of extraction. The calibration curve for the standard solution of Alloxazine was made with the 157 concentration levels of 0, 0.005, 0.010, 0.025, 0.050 and 0.100 μ g mL⁻¹ (including zero point). These 158 solutions were prepared in 0.1% aqueous formic acid solution. The calibration curve was built by 159 representing concentrations of Alloxazine against the corresponding peak The area. 160 selectivity/specificity was analyzed by testing 20 representative blank flour samples of different origin 161 in order to verify the absence of potential interfering compounds at retention time of the analyte. The 162 precision (repeatability and the within-laboratory reproducibility) and the trueness were calculated by 163 the determination of samples fortified at three levels (1 - 5 - 10 mg kg⁻¹), at beginning of the extraction 164 procedure. Ten aliquots were analyzed for each level, for three batches successively in a 3-week 165 period, giving a total of 90 replicates. The concentration of each replicate was determined using the 166 calibration curve prepared on the same batch. The precision was expressed as the RSD and

- 167 repeatability values calculated for each level (Table 3). The average recovery was estimated using
- 168 these matrix results.
- 169 The detection limit (LOD) was estimated on the basis of the results of ten replicates of flour sample 170 spiked at 0.5 mg kg^{-1} .

Five operating factors were chosen for ruggedness study (performed at 5 mg kg⁻¹): adsorption time, centrifugation time and speed, concentration of hydrochloric acid in working solution, vortex time. The different factors and their levels were mixed in the Youden experimental plan (Youden and Steiner, 1975).

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176 2.7 Data collection

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All the data obtained from the LC-MS/MS analysis were collected on an Excel datasheet and then sorted by flour type in order to evaluate possible differences in Alloxan presence between bread, pastry and cake flour.

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3. RESULTS AND DISCUSSION

184 *3.1 Chromatography and derivatization studies*

Alloxan is a ketone with a low molecular weight and neutral functional groups. For these compounds, the fragmentation is difficult due to unstable transitions, so the ionization efficiency in ESI is low (Santa, 2011). The pre-column derivatization with o-phenylenediamine, which produces Alloxazine, has provided a better way to detect the analyte in question, due to a greater stability (Raghavamenon et al., 2009). Alloxazine is the result of reaction between a primary amine and carbonyl groups with formation of a product containing carbon nitrogen double bonds (Figure 1). The reaction is acid catalyzed by hydrochloric acid with elimination of two molecules of water.

192 The comparison between Alloxazine standard and derivatization product gave a full matching, for each the four identification points monitored with MRM conditions. An alloxan solution was admixed with 193 194 an excess of derivatizing agent (o-phenylenediamine), considering the stoichiometry as known and the 195 vield as unknown. The experiments were carried out evaluating the temperature and the reaction times. 196 The kinetic of reaction was studied at three different incubation temperatures: 4 - 25 and 50°C. 197 Furthermore, the instrumental results was collected at different time intervals: 5min - 15min - 30min -198 45min – 90min - 12h - 24h - 30h and 48h, in order to evaluating the reaction times. It was found an 199 increasing trend of the yield percentage, up to a maximum value in the 24h, following by a plateau in the next hours. The effect of the temperature on the yield was 4% at 4°C, 100% at 25 °C and 29% at 50
°C, respectively. All the results are shown in Figure 2.

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203 *3.2 Validation of the method*

Good validation parameters were obtained (linearity, selectivity/specificity, precision and accuracy,
 ruggedness, recovery, LOD and LOQ), according to ISO/IEC/EN 17025/2005.

The blank samples were fortified at three different concentrations: $1 - 5 - 10 \text{ mg kg}^{-1}$ with standard solution of Alloxan. Ten spiked samples, at each of the three levels, were analysed. The analysis of replicates (ten for each level) were repeated once a week, for a total of three weeks. Representative chromatogram of fortified sample is shown in Figure 3b. For the linearity, the calibration curve obtained for Alloxazine, constructed by plotting the peak area (y) versus concentration (x) of the analyte, giving a correlation coefficient (r²) of 0.9991. The areas used for the quantification are generated by the base peak signal only.

The Selectivity/Specificity test verified no interfering peaks near the retention time of the analyte. The retention time was 2.1 min for Alloxazine and the obtained peak is symmetrical.

The results of intraday and interday reproducibility are listed in Table 3. The overall precision of the assay expressed as RSD was less than 10% in the flour samples. Trueness was expressed as recovery rate and ranged from 94% to 102%. The recovery data for the low, medium and high levels are shown in Table 3. A detection limit (LOD) of 0.73 mg kg⁻¹ and a quantification limit (LOQ) of 0.85 mg kg⁻¹ were achieved by the method proposed.

The measuring range, studied in this work, is included between 0.85-10.0 mg kg⁻¹; representing the range of concentration between the lowest limit, the LOQ, and the highest limit (the validation point with the highest concentration). The expanded uncertainty of the method was estimated having regard to every contributions of the important elements (weights, volumes, repeatability, standards, calibration uncertainty). The obtained values are shown in Table 3. The ruggedness study, performed at 5 mg kg⁻¹, confirmed that the tested factors are not critical.

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227 *3.3 Samples analysis*

Alloxan trace levels were found in 42 (24%) of the analyzed samples, with mean values of $0,95\pm0,04$ mg kg⁻¹ and a range between 0.88 and 1.02 mg kg⁻¹. The positive samples were confirmed by repeated analysis. The identification of the target compound was based on RT with a window of \pm 3 times the SD value of the RT, and MS² productions of the characteristic ion. The quantification was performed using the signal of base peak only. The chromatograms of a positive flour sample and spiked sample 233 are shown in Figure 3a and 3b, respectively. The presence of Alloxan was found only in cake flour, 234 suggesting possible correlations with the use of chemical substances such as chlorine gas and chlorine 235 dioxide for smooth and fine cake flour production. Bleaching compounds tightens the protein 236 molecules in the flour, enabling it to carry more than its weight in sugar and fat. Therefore, most of the 237 cake flour are bleached in order to improve their baking performance and responding to a wide request 238 for production (about 140 million pounds of flour each day in USA). The use of chemical oxidizing 239 agents and bleaches were developed to produce quick aging of wheat flour (48 hours), instead the 240 natural conditions that require several months (Fennema, 1985). Chlorine gas used as bleaching agent may reacts with some proteins in the flour (including the gluten) producing Alloxan as a by-product 241 242 (Cohen, 2010). High-gluten flours, such as the cake flour examined in this work, may contain 5 to 8 g 243 of gluten per 100 g of flour, suggesting a possible presence of Alloxan due to reaction with chlorine gas 244 and other chemicals.

245 Furthermore, the presence of these compounds could lead to a series of oxidation reactions that may 246 modify many flour components such as the pterins (Socaciu, 2007). Pterins share with flavins properties such as radical formation, participation in redox chains, photosensitizing capacity and 247 248 absorption of near-ultraviolet light (Galland and Senger, 1988). The total amount of pterins depends on 249 the plant species, on the developmental stage, and on external factors; good sources of these 250 compounds are legumes and wheat germ (Rébeillé et al., 2006). Several of these compounds on 251 oxidation with chlorine and methanol yield 5-methoxyuramil-7-oxalic acid methyl ester together with 252 glycol monomethyl ether, and the ester on hydrolysis gives oxalic acid and Alloxan (Engineers, 2005).

253 Although there are different studies on the acute effects of Alloxan in rats and other organisms at 254 concentrations higher than what was found in this study (50-100 mg kg⁻¹) (Bakırel et al., 2008; Lenzen, 255 2008), very little is known about the chronic effects of Alloxan to specific concentrations. Vadlamudi 256 et al. (1982) showed depressed left ventricular pressure and positive and negative dP/dt development in 257 Witsar rats after 100, 180 and 360 days of treatment with Alloxan (Vadlamudi et al., 1982), while a 258 preliminary study conducted by de Olivera et al. (2005) showed blood and tissue alterations after 90 259 days of Alloxan treatment (De Oliveira et al., 2005). So further studies are needed in order to evaluate 260 the chronic effects of this molecule, the timing of dose and the toxic levels for human health. In United 261 States, wheat flour is normally bleached with chlorine gas prior to its use in baking cakes. Chemical 262 treatments and chemical additives have become suspect and alternative methods to avoid such 263 treatments are needed. Many European countries ban the use of chemical bleaching and oxidation 264 chemicals and other additives in bread completely. The Environmental Protection Agency (EPA) 265 identifies chlorine gas as a flour-bleaching, aging and oxidizing agent that is a powerful irritant,

dangerous to inhale, and lethal (Cohen, 2010). An alternative to the chlorination method is to subject
the flour to specified temperatures for limited periods of time, this process does not pose a potential
hazard to the health of those who consume the products (Hanamoto and Bean, 2005).

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4. CONCLUSIONS

272 According to our knowledge, this work represents the first report on the presence of Alloxan in cake 273 bleached flour, suggesting a potential risk for consumers due to the application of chlorine gas and 274 other chemicals for baking cakes. The reported UHPLC-MS/MS method was found very sensitive and 275 accurate for the determination of Alloxan in wheat flour starting from 0.85 mg kg⁻¹. The method was successfully applied to the analysis of 175 real samples, with the aim of verify the presence of this 276 277 contaminant. The results obtained show that the flour bleached with chlorine dioxide and chlorine gas may contain Alloxan as a minor product of a series of oxidation reactions. As a pilot study, further 278 279 studies are needed with a larger number of flour samples, in order to understand the real risk for the 280 consumers.

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REFERENCES

- Bakırel, T., Bakırel, U., Keleş, O.Ü., Ülgen, S.G., Yardibi, H., 2008. In vivo assessment of antidiabetic
 and antioxidant activities of rosemary (Rosmarinus officinalis) in alloxan-diabetic rabbits. J.
 Ethnopharmacol. 116, 64–73. doi:10.1016/j.jep.2007.10.039
- Banu Shakila, M., Sasikala, P., 2012. Alloxan in refined flour: A Diabetic concern. Int. J. Adv. Innov.
 Res. 1, 204–209.
- Carvalho, E.N. de, Carvalho, N.A.S. de, Ferreira, L.M., 2003. Experimental model of induction of
 diabetes mellitus in rats. Acta Cir. Bras. 18, 60–64. doi:10.1590/S0102-86502003001100009
- Chittrakorn, S., Earls, D., MacRitchie, F., 2014. Ozonation as an alternative to chlorination for soft
 wheat flours. J. Cereal Sci. 60, 217–221. doi:10.1016/j.jcs.2014.02.013
- Cohen, S., 2010. Diabetes without Drugs: The 5-Step Program to Control Blood Sugar Naturally and
 Prevent Diabetes Complications. Rodale.
- De Oliveira, C.A.M., Luciano, E., De Mello, M.A.R., 2005. The role of exercise on long-term effects
 of alloxan administered in neonatal rats. Exp. Physiol. 90, 79–86.
- doi:10.1113/expphysiol.2004.028241

- Dhanabal, S.P., Mohan Maruga Raja, M.K., Ramanathan, M., Suresh, B., 2007. Hypoglycemic activity
 of Nymphaea stellata leaves ethanolic extract in alloxan induced diabetic rats. Fitoterapia 78,
 288–291. doi:10.1016/j.fitote.2007.02.009
- Dunn, J.S., Kirkpatrick, J., McLetchie, N.G.B., Telfer, S.V., 1943. Necrosis of the islets of Langerhans
 produced experimentally. J. Pathol. Bacteriol. 55, 245–257. doi:10.1002/path.1700550302
- Engineers, N.B. of C.&, 2005. The Complete book on Natural Dyes & Pigments. ASIA PACIFIC
 BUSINESS PRESS Inc.
- 305 Fennema, O.R., 1985. Food additives, 2nd ed. Marcel Dekker Inc, New York.
- Fukayama, M.Y., Tan, H., Wheeler, W.B., Wei, C.I., 1986. Reactions of aqueous chlorine and chlorine
 dioxide with model food compounds. Environ. Health Perspect. 69, 267–274.
- Galland, P., Senger, H., 1988. The Role of Pterins in the Photoreception and Metabolism of Plants.
 Photochem. Photobiol. 48, 811–820. doi:10.1111/j.1751-1097.1988.tb02896.x
- Goldner, M.G., Gomori, G., 1943. ALLOXAN DIABETES IN THE DOG. Endocrinology 33, 297–
 308. doi:10.1210/endo-33-5-297
- 312 Hanamoto, M.M., Bean, M.M., n.d. Process for improving baking properties of unbleached flour.
- Idaho Observer: Bleaching agent in flour linked to diabetes [WWW Document], 2005. URL
 http://proliberty.com/observer/20050718.htm (accessed 3.2.17).
- ISO/IEC 17025:2005 General requirements for the competence of testing and calibration laboratories,
 n.d.
- Jacobs, H.R., 1937. Hypoglycemic Action of Alloxan. Proc. Soc. Exp. Biol. Med. 37, 407–409.
 doi:10.3181/00379727-37-9590P
- Joye, I.J., Lagrain, B., Delcour, J.A., 2009. Use of chemical redox agents and exogenous enzymes to
 modify the protein network during breadmaking A review. J. Cereal Sci. 50, 11–21.
 doi:10.1016/j.jcs.2009.04.001
- Kim, H.-R., Rho, H.-W., Park, B.-H., Park, J.-W., Kim, J.-S., Kim, U.-H., Chung, M.-Y., 1994. Role of
 Ca2+ in alloxan-induced pancreatic β-cell damage. Biochim. Biophys. Acta BBA Mol. Basis
 Dis. 1227, 87–91. doi:10.1016/0925-4439(94)90111-2
- Kulp, K., Olewnik, M., Bachofer, C., 1985. Functional effects of chlorinated flour on cookie spread
 and quality of sugar snap cookies. Tech Bull Am Inst Bak. 7, 1–9.
- Lenzen, S., 2008. The mechanisms of alloxan- and streptozotocin-induced diabetes. Diabetologia 51,
 216–226. doi:10.1007/s00125-007-0886-7
- Lenzen, S., Panten, U., 1988. Alloxan: history and mechanism of action. Diabetologia 31, 337–342.
 doi:10.1007/BF02341500

- 331 Mrozikiewicz, A., Kiełczewska-Mrozikiewicz, D., Łowicki, Z., Chmara, E., Korzeniowska, K.,
- Mrozikiewicz, P.M., 1994. Blood levels of alloxan in children with insulin-dependent diabetes
 mellitus. Acta Diabetol. 31, 236–237. doi:10.1007/BF00571958
- Pincus, G., 2013. Recent Progress in Hormone Research: The Proceedings of the Laurentian Hormone
 Conference. Elsevier.
- Raghavamenon, A.C., Dupard-Julien, C.L., Kandlakunta, B., Uppu, R.M., 2009. Determination of
- alloxan by fluorometric high-performance liquid chromatography. Toxicol. Mech. Methods 19,
 498–502. doi:10.3109/15376510903334862
- Rébeillé, F., Ravanel, S., Jabrin, S., Douce, R., Storozhenko, S., Van Der Straeten, D., 2006. Folates in
 plants: biosynthesis, distribution, and enhancement. Physiol. Plant. 126, 330–342.
 doi:10.1111/j.1399-3054.2006.00587.x
- Saadia, S.A., Khan, H.A., Ghazanfar, A.S., Saadatullah, K., 2005. Alloxan induced diabetes in rabbits.
 Pak. J. Pharmacol. 22, 41–45.
- Saleem Mir, M., Maqbool Darzi, M., Musadiq Khan, H., Ahmad Kamil, S., Hassan Sofi, A., Ahmad
 Wani, S., 2013. Pathomorphological effects of Alloxan induced acute hypoglycaemia in rabbits.
 Alex. J. Med. 49, 343–353. doi:10.1016/j.ajme.2013.03.007
- Santa, T., 2011. Derivatization reagents in liquid chromatography/electrospray ionization tandem mass
 spectrometry. Biomed. Chromatogr. 25, 1–10. doi:10.1002/bmc.1548
- Seifu, D., Gustafsson, L.E., Chawla, R., Genet, S., Debella, A., Holst, M., Hellström, P.M., 2017.
 Antidiabetic and gastric emptying inhibitory effect of herbal Melia azedarach leaf extract in
- rodent models of diabetes type 2 mellitus. J. Exp. Pharmacol. 9, 23–29.
- doi:10.2147/JEP.S126146
- Shaw Dunn, J., Mcletchie, N.G.B., 1943. EXPERIMENTAL ALLOXAN DIABETES IN THE RAT.
 The Lancet, Originally published as Volume 2, Issue 6265 242, 384–387. doi:10.1016/S01406736(00)87397-3
- 356 Socaciu, C., 2007. Food Colorants: Chemical and Functional Properties. CRC Press.
- 357 Suganuma, N., Kikkawa, F., Seo, H., Matsui, N., Tomoda, Y., 1993. Poly (adenosine diphosphate-
- ribose) synthesis in the anterior pituitary of the female rat throughout the estrous cycle: Study of
 possible relation to cell proliferation and prolactin gene expression. J. Endocrinol. Invest. 16,
 475–480. doi:10.1007/BF03348885
- Szkudelski, T., 2001. The mechanism of alloxan and streptozotocin action in B cells of the rat
 pancreas. Physiol. Res. 50, 537–546.

- Thomasson, C.A. (Kansas S.U., Miller, R.A., Hoseney, R.C., 1995. Replacement of chlorine treatment
 for cake flour. Cereal Chem. USA.
- Vadlamudi, R.V.S.V., Rodgers, R.L., McNeill, J.H., 1982. The effect of chronic alloxan- and
 streptozotocin-induced diabetes on isolated rat heart performance. Can. J. Physiol. Pharmacol.
 60, 902–911. doi:10.1139/y82-127
- Weaver, D.C., McDaniel, M.L., Naber, S.P., Barry, C.D., Lacy, P.E., 1978. Alloxan Stimulation and
 Inhibition of Insulin Release from Isolated Rat Islets of Langerhans. Diabetes 27, 1205–1214.
 doi:10.2337/diab.27.12.1205
- Webb, L., 1966. ENZYME AND METABOLIC INHIBITORS,. Academic Press., New York and
 London.
- 373 Youden, W.J., Steiner, E.H., 1975. Statistical Manual of the AOAC, 5th ed. AOAC
- 374 INTERNATIONAL.
- 375 Zhou, W., Wei, L., Xiao, T., Lai, C., Peng, M., Xu, L., Luo, X., Deng, S., Zhang, F., 2017.
- Diabetogenic agent alloxan is a proteasome inhibitor. Biochem. Biophys. Res. Commun. 488,
 400–406. doi:10.1016/j.bbrc.2017.05.065

Time (min.)	Component A (%)	Component B (%)			
0.0	85	15			
0.5	85 15				
2.0	5	95			
3.0	5	95			
3.5	85	15			
7.0	85	15			
Table 1. Gradient profile for the determination of Alloxan.					

Analyte Parent ion		Transition 1Collision energy (e	
		169.9ª	19
Alloxazine	215.1	Transition 2	Collision energy (eV)
		144.1	22

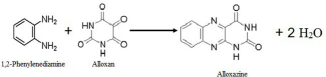
abundant product ion.

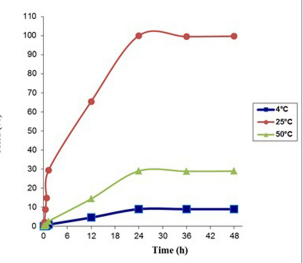
	Precision		Trueness	
Fortification level (mg kg ⁻¹)	Intra-day analysis RSD (%) (n=10)	Inter-day analysis RSD (%) (n=30)	Recovery (%)	Uncertainty (mg kg ⁻¹)
1.0	6.0	9.9	94	0.2
5.0	4.6	8.9	102	0.5
10.0	3.1	7.0	99	1.1

Table 3. Precision and trueness study at the three validation levels. *n*: number of determinations.

- **Fig.1.** Chemical Formula of Alloxan (2,4,5,6- tetraoxypyrimidine) and Alloxazine (Benzo[g]pteridine-
- 388 2,4, (1H,3H)-dione): Schematic representation of derivatization with 1,2-Phenylenediamine.389
- Fig. 2 Graphic representation of the yield percentage of the derivatization reaction: kinetic of reactionstudied as a function of the time at three different temperatures.
- 392
- Fig. 3. Chromatograms of a real positive wheat flour sample (a) and spiked wheat flour sample (b) with relative m/z ratios.
- 395
- 396

- In this work the presence of Alloxan in bleached flour was reported for the first time;
- A reliable UHPLC–MS/MS method was carried out according to EN/ISO 17025;
- Alloxan trace levels were found in 24% of the cake flour analyzed samples;
- Bleached flour may contain Alloxan as a minor product of a series of oxidation reactions.





Yield (%)

