Occurrence of acrylamide carcinogen in Arabic coffee *Qahwa*, coffee and tea from Saudi Arabian market

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The present work describes the outcomes of the assessment on acrylamide contents in a number of thermally treated foods (Arabic coffee *Qahwa*, coffee and tea) obtained from the Saudi Arabian markets. A total of 56 food samples of different brands and origin were studied, the amounts of acrylamide in Arabic coffee *Qahwa*, coffee and tea were obtained in the range of 10 to 682 μg kg\(^{-1}\). In comparison to coffee (152–682 μg kg\(^{-1}\)), the Arabic coffee *Qahwa* (73–108 μg kg\(^{-1}\)) and tea (10–97 μg kg\(^{-1}\)) contain lower amounts of acrylamide. Among the analyzed samples, the green tea contained low amounts of acrylamide ranged from 10 to 18 μg kg\(^{-1}\), and thus the green tea could be considered as a healthier hot drink. A great variation of acrylamide formation has been observed in these food products. This divergence may be due to the initial concentration of amino acids especially asparagines and reducing sugars in food products, in addition to roasting temperature and time, pH and water activity. The obtained data can also be used in epidemiological investigation to estimate the acrylamide exposure from nutritional survey.

In 2002, the University of Stockholm and the Swedish National Food Administration have jointly publicized that a number of food products thermally processed at elevated temperature contain fairly high amounts of acrylamide\(^1\). Since then, acrylamide gained a substantial awareness and it is known as process food contaminant\(^2\). The acrylamide is generated in the consequence of thermal treatment of foodstuffs via Maillard reaction, a reaction between the free amino acid mainly asparagines with reducing sugars at high cooking temperature\(^3\).

Previously, animal feeding investigation confirmed an association linking acrylamide in particular hemoglobin adducts and cooked animal foodstuffs\(^4\). To a certain extent, the long-term contact to acrylamide might cause harm to the nervous system including potential genetic and reproductive toxin in humans as well as in experimental animals\(^5\). In 1994, the International Agency for Research on Cancer (IARC) has classified acrylamide as ‘probably carcinogenic to humans’\(^6\). Recently, the European Food Safety Agency (EFSA) has proclaimed that the roasted coffee including other cooked foods for instance potatoes, crackers, cookies, toasted bread and breakfast cereals are the major source of acrylamide\(^7\), which contributes higher than 90% acrylamide intake through our daily diets. The concentrations of acrylamide in some food products for instance potato chips and coffee reach up to 3500 μg kg\(^{-1}\) or even higher than 4500 μg kg\(^{-1}\). These foodstuffs are significant from the opinion of contact estimation, either due to these foods holds a large amount of acrylamide compare to other food products or these food products are an element of the usual diet for the common inhabitants, therefore contain an elevated eating rates. Previously, the normal acrylamide ingestion through diet was described and found 1 μg kg\(^{-1}\) body weight day\(^{-1}\) and 4 μg kg\(^{-1}\) body weight day\(^{-1}\) for common inhabitants and higher consumers, respectively\(^8,9\).

In the recent years, the food lifestyles have significantly changed worldwide and gradually increasing eating of processed food products\(^10\). These food habits alteration are mainly apparent in juvenile populations whose lifestyles are allied with the consumption of high amounts of fast foods including Arabic coffee *Qahwa*, coffee and tea which supplies to the greater amounts of acrylamide ingestion\(^1\). In an attempt to assess the acrylamide...
of concentrations and recovery rates have been demonstrated in Tables 1, 2 and 3, where it can be observed that instance Arabic coffee Qahwa usually favor the formation of acrylamide.

From such types of food products.

The population concerning acrylamide carcinogen, as well as it can be utilized to approximate the ingestion of acrylamide exposed investigation will increase the consciousness in the Saudi Arabian inhabitants together with the global population concerning acrylamide carcinogen, as well as it can be utilized to approximate the ingestion of acrylamide from such types of food products.

Results and Discussion

Recently, the Joint FAO/WHO Expert Committee has described the explanatory statistics of acrylamide levels in various food products for the data collected between 2004 and 2009. The average levels of acrylamide in main foods were: coffee (3–68 μg kg⁻¹), potato chips (399–1202 μg kg⁻¹), cookies (169–518 μg kg⁻¹), French fries (159–963 μg kg⁻¹), and crisp bread and crackers (87–459 μg kg⁻¹). On the body weight basis, the average nutritional acrylamide ingestion for common inhabitants together with children, the acrylamide exposure in children are higher up to two folds relative to the adults population. On the basis of earlier findings of acrylamide in thermally processed carbohydrate-rich foods and its exposure to the human beings, it is highly essential to perform the screening of toxic acrylamide in such type of food products frequently available in Saudi diets.

For the analysis of acrylamide, the performance of the method (quality parameters) in terms of linearity, limit of detection (LOD), limit of quantification (LOQ) and precisions (run–to–run and day–to–day) were established. The linearity of the method was determined in the concentration range of 8–2000 μg L⁻¹ and excellent linearity (R² = 0.999) was obtained. The LOD (signal-to-noise ratio, 3:1) and LOQ (signal-to-noise ratio, 10:1) were calculated and obtained 2 μg L⁻¹ and 7 μg L⁻¹, correspondingly. To establish the run–to–run precision, five replicates of a known amount of acrylamide standard solution (600 μg L⁻¹) were analyzed on the same day. The relative standard deviation (RSD) was estimated from the five determined concentration levels, and the outcomes <2.0% were achieved. For the determination of day–to–day precision was determined by five replicates of same standard solution (600 μg L⁻¹) was analyzed over three successive days, and the RSD values <4% were attained. The outcomes of the run–to–run precision and day–to-day precision data showed good precision for ultra-performance liquid chromatography-tandem mass spectrometry (UPLC-MS/MS) determination of acrylamide in food products.

A total of 36 food products of different brands and origin were studied, the samples include Arabic coffee Qahwa, coffee and tea. So as to overcome the sample matrix effects, the quantification of acrylamide in food samples was carried out using isotope dilution method utilizing AA-D₃ as the labeled. The obtained results in terms of concentrations and recovery rates have been demonstrated in Tables 1, 2 and 3, where it can be observed that the acrylamide was identified in all of the analyzed food samples and concentrations reached up to 682 μg kg⁻¹. The recovery rates were achieved between 86% and 93% in all studied food samples. The acrylamide concentrations in Arabic coffee Qahwa have been illustrated in Table 1. The acrylamide was detected in all samples in the range of 73 to 108 μg kg⁻¹. The highest amount of acrylamide was identified in sample 7 (Maatouk) (108 μg kg⁻¹), while sample 8 (Ajyal) contains the lowest level (73 μg kg⁻¹). Apart from sample 7 (Maatouk), the other samples did not demonstrate a huge variation in acrylamide concentrations. Previously, a few studies relating to the determination of acrylamide in Arabic coffee samples have been reported. In particular, one Arabic coffee (light roasted) sample from the Saudi Arabia and two samples including Arabic coffee and Arabic coffee (dark) from the Egypt have been studied. The obtained acrylamide concentrations are found totally different from our results. In the Arabic coffee (light roasted) sample from the Saudi Arabia, the author has not detected any amount of acrylamide. Whereas, the studies performed in the Egypt, the author has detected acrylamide in both Arabic coffee (272 μg kg⁻¹) and Arabic coffee (dark, 480 μg kg⁻¹) samples. The significant differences in the results might be due to the different analytical methodologies had applied or the samples had been thermally processed at different roasting temperatures which favor the acrylamide formation.

| Sample | Type   | Country of origin | Acrylamide (μg kg⁻¹) ± SD | Recovery rates (%) |
|--------|--------|-------------------|---------------------------|-------------------|
| Sample 1 | Harrari | Ethiopia           | 74 ± 3                    | 89                |
| Sample 2 | Khaolani | Yemen              | 78 ± 3                    | 90                |
| Sample 3 | Baraeh | Yemen              | 82 ± 4                    | 92                |
| Sample 4 | Alasalah | Ethiopia          | 85 ± 4                    | 90                |
| Sample 5 | Anoosh | —                 | 79 ± 3                    | 91                |
| Sample 6 | Bayouni | Turkey            | 80 ± 3                    | 92                |
| Sample 7 | Maatouk | —                 | 108 ± 6                   | 90                |
| Sample 8 | Ajyal | —                 | 73 ± 3                    | 87                |
| Sample 9 | Yatoon | —                 | 89 ± 4                    | 92                |

Table 1. Amount of acrylamide and recovery rates in cooked Arabic coffee Qahwa. — not described on label; SD = standard deviation.
| Sample  | Country of origin | Acrylamide (μg kg⁻¹) ± SD | Recovery rates (%) |
|---------|-------------------|---------------------------|-------------------|
| Sample 1 | Brazil            | 682 ± 23                  | 92                |
| Sample 2 | Brazil            | 412 ± 17                  | 89                |
| Sample 3 | Switzerland       | 250 ± 12                  | 90                |
| Sample 4 | Malaysia          | 246 ± 12                  | 92                |
| Sample 5 | Ecuador           | 654 ± 22                  | 92                |
| Sample 6 | Brazil            | 502 ± 20                  | 90                |
| Sample 7 | Turkey            | 587 ± 22                  | 92                |
| Sample 8 | Turkey            | 305 ± 14                  | 93                |
| Sample 9 | Costa Rica        | 214 ± 10                  | 87                |
| Sample 10| Turkey            | 320 ± 15                  | 90                |
| Sample 11| Brazil            | 225 ± 14                  | 88                |
| Sample 12| Brazil            | 340 ± 15                  | 86                |
| Sample 13| Colombia          | 234 ± 12                  | 91                |
| Sample 14| Germany           | 560 ± 22                  | 92                |
| Sample 15| Switzerland       | 201 ± 10                  | 91                |
| Sample 16| Turkey            | 152 ± 8                   | 88                |
| Sample 17| Germany           | 247 ± 12                  | 89                |
| Sample 18| Germany           | 460 ± 18                  | 93                |
| Sample 19| Germany           | 475 ± 18                  | 93                |
| Sample 20| Germany           | 432 ± 17                  | 91                |
| Sample 21| Germany           | 360 ± 13                  | 87                |
| Sample 22| Netherlands       | 450 ± 18                  | 91                |
| Sample 23| France            | 368 ± 15                  | 91                |
| Sample 24| Switzerland       | 184 ± 10                  | 90                |
| Sample 25| Malaysia          | 430 ± 17                  | 92                |

Table 2. Amount of acrylamide and recovery rates in coffee products. *All samples were ground coffee beans type except sample 4 and 17 (freeze dried); SD = standard deviation.

| Sample  | Type             | Country of origin | Acrylamide (μg kg⁻¹) ± SD | Recovery rates (%) |
|---------|------------------|-------------------|---------------------------|-------------------|
| Sample 1| Black, Loose     | —                 | 84 ± 4                    | 91                |
| Sample 2| Black, Loose     | Spain             | 73 ± 3                    | 89                |
| Sample 3| Black Fine, Loose| —                 | 97 ± 6                    | 92                |
| Sample 4| Black, Loose     | —                 | 72 ± 3                    | 88                |
| Sample 5| Black, Loose     | —                 | 78 ± 4                    | 90                |
| Sample 6| Black, Loose     | —                 | 80 ± 5                    | 88                |
| Sample 7| Black, Loose     | —                 | 68 ± 3                    | 91                |
| Sample 8| Black, Bag       | —                 | 82 ± 4                    | 92                |
| Sample 9| Black, Bag       | Kenya             | 74 ± 4                    | 90                |
| Sample 10| Black, Bag      | UK                | 79 ± 4                    | 92                |
| Sample 11| Natural Green   | —                 | 18 ± 2                    | 91                |
| Sample 12| Green           | —                 | 23 ± 2                    | 89                |
| Sample 13| Pure Green      | —                 | 10 ± 1                    | 90                |
| Sample 14| Black, Bag      | Sri Lanka         | 79 ± 4                    | 91                |
| Sample 15| Green           | Sri Lanka         | 11 ± 1                    | 90                |
| Sample 16| Black, Loose    | Kenya             | 74 ± 3                    | 91                |
| Sample 17| Black, Bag      | —                 | 72 ± 3                    | 91                |
| Sample 18| Earl Grey, Bag  | UK                | 35 ± 2                    | 88                |
| Sample 19| Black, Bag      | —                 | 81 ± 4                    | 91                |
| Sample 20| Black, Bag      | Sri Lanka         | 78 ± 4                    | 92                |
| Sample 21| Black, Bag      | —                 | 62 ± 3                    | 92                |
| Sample 22| Black, Bag      | —                 | 85 ± 4                    | 89                |

Table 3. Amount of acrylamide and recovery rates in tea products. —not described on label; SD = standard deviation; UK = United Kingdom.
Table 2 illustrates the amounts of acrylamide in roasted coffee of different brands and origin. The highest amounts of acrylamide were obtained in sample 1 (682 μg kg\(^{-1}\)) whereas, sample 16 comparatively produced low amount (152 μg kg\(^{-1}\)). This might have caused of high roasting temperature (160–210 °C) applied in sample 1 than sample 16. Formerly, a complete assessment on amounts of acrylamide in various food products has been reported\(^{19}\), particularly on coffee by the European Food Safety Authority\(^{21,23}\), and in few samples by Guenther \textit{et al.} and others\(^{22–24}\). The levels of acrylamide reached up to 1047 μg kg\(^{-1}\), which are similar to those reported by the Food and Drug Administration\(^{25}\). In concurrence with the results obtained, the amounts of acrylamide in sample 2, sample 18, sample 19, sample 22 and sample 25 are found to be average values of 443 μg kg\(^{-1}\), which are similar to those reported by the Food and Drug Administration\(^{25}\). Alternatively, sample 1, sample 5, sample 6, sample 7 and sample 14 are found to be slightly higher concentrations with an average of 597 μg kg\(^{-1}\) and found in good agreements with the values obtained in the similar study\(^{16,26–28}\). These higher values are most likely as a result of the extreme roasting conditions and it is very difficult to compare with earlier published literatures. The concentrations of acrylamide in remaining samples including sample 3, sample 4, sample 8, sample 9, sample 10, sample 11–16, sample 17, sample 23 and sample 24 ranged from 152 to 368 μg kg\(^{-1}\). These values are in good agreement to those obtained in the European Union database\(^{28}\), where the lowest and highest concentrations oscillated from 79 to 2955 μg kg\(^{-1}\). Relatively, these low amounts, considering the existence of substantial precursors amount, are most likely due to the moderate roasting temperature applied in the preparation of these food products\(^{29}\).

About tea, a few studies relating to the occurrence of acrylamide are obtainable, thus it was very essential to make the screening of acrylamide contents in such kind of food samples. In the present study, a total of 22 tea samples of two classes (black and green) were studied, the obtained concentrations has been demonstrated in Table 3. The amounts of acrylamide in studied samples were found <100 μg kg\(^{-1}\). From the obtained data, it has been observed that the green types of tea samples contained low levels of acrylamide (10–18 μg kg\(^{-1}\)), most likely the green tea were not dried out at elevated temperature. The black tea either loose or in bags comparatively comprise higher amounts of acrylamide ranged from 35 to 97 μg kg\(^{-1}\), this might be caused that these samples have been thermally processed at higher temperature between 100 °C and150 °C\(^{29}\). Our results are found in good agreement with those obtained in previous studies\(^{29,30}\) where the amounts of acrylamide in tea samples were also detected lower than 100 μg kg\(^{-1}\).

In comparison to coffee, the Arabic coffee \textit{Qahwa} and tea contain lower amounts of acrylamide and below the consumption limit as recommended by the European Commission\(^{31}\). Among these hot drink samples, the Arabic coffee \textit{Qahwa}, coffee and tea samples showed an important source of acrylamide and among these, tea especially green type contains lower amounts of acrylamide and can be considered as a heather drink followed by Arabic coffee \textit{Qahwa} and coffee. Moreover, green coffee is also recognized for a variety of health advantages which linked with threat diminution of various chronic diseases, for instance cardiovascular, cancer and diabetes\(^{32}\).

To display the results, the UPLC-MS/MS chromatograms of acrylamide in Arabic coffee \textit{Qahwa} (sample 1, Harrari), coffee (sample 1) and tea (sample 3, Black Fine) have been illustrated in Figs 1, 2 and 3, respectively. The tremendous sensitivity, excellent peak symmetry and with no any interfering peaks arising at the elution time of target analyte was achieved.
Conclusions

The contents of acrylamide in 56 food samples including Arabic coffee Qahwa, coffee and tea of different brands and origin were investigated. The acrylamide was detected in all of the analyzed samples and the concentrations ranged from 10 to 682 μg kg\(^{-1}\). In comparison to coffee, the Arabic coffee Qahwa and tea contain lower amounts of acrylamide. Additionally, the green tea contained low contents of acrylamide ranging 10 to 18 μg kg\(^{-1}\) and on the basis of acrylamide amounts we can assume green tea as a healthier hot drink. The excellent recovery rates (up to 93%) and quality parameters (linearity, LOD, LOQ and precisions) of the method were achieved. The outcomes revealed that the concentration of acrylamide in cooked foods show a big difference between diverse food
groups and brands. This variance may be due to the initial concentration of amino acids especially asparagines and reducing sugars in food products, in addition to the roasting temperature and time, pH and water activity. The obtained data can also be used in epidemiological investigation to estimate the acrylamide exposure from nutritional survey.

**Materials and Methods**

**Chemicals and reagents.** Acrylamide (assay ≥ 99.8%), acrylamide–2,3,3–D3 (AA–D3, isotopic purity 98%) (Fig. 4), potassium hexacyanoferrate (assay ≥ 99.0%) and zinc acetate dihydrate (assay ≥ 98%) were purchased from Sigma–Aldrich (Sigma–Aldrich Chemie GmbH, Steinheim, Germany). Acetonitrile and methanol of HPLC–grade were purchased from Merck (Darmstadt, Germany). A Milli–Q water purification device, model advantage A10 from Millipore Corporation (Bedford, USA) was used to purify the water. Acrylamide stock solutions of 1 mg mL−1 and AA–D3 of 1 mg mL−1 were prepared with Milli–Q water. To get a series of standard solutions, the stocks solutions were diluted with Milli–Q water. Relating to the linearity, the calibration curve for the quantification of acrylamide in foods by UPLC-MS/MS was constructed and found linear over the range of 8–2000 μg L−1. All solutions were refrigerated at −4 °C until analysis. To get the Carrez I and II solutions, potassium hexacyanoferrate (10.6 g) and zinc acetate dihydrate (24 g) were individually dissolved in 100 mL of Milli–Q water.

Solid phase extraction (SPE) columns Strata™ –X–C polymeric strong cation, 200 mg, 6 mL was obtained from Phenomenex (Torrance, USA) and Isolute ENV+, 200 mg, 3 mL was purchased from Biotage (Uppsala, Sweden). PTFE syringe filters (0.45 μm) were supplied from Macherey–Nagel GmbH (Düren, Germany). Nitrocellulose membrane, pore size 0.45 μm was purchased from Sigma-Aldrich (Sigma-Aldrich Chemie GmbH, Steinheim, Germany).

**Sample treatment.** The cooked food samples (Arabic coffee Qahwa, coffee and tea) were purchased from a local supermarket (Riyadh, Saudi Arabia). The food samples were blended and homogenized by means of Microtron® MB 800 (Kinematica AG, Littau, Switzerland), ultra-turrax T25 digital (IKA®-WERKE GmbH, Staufen, Germany) and coffee grinder CML-1000MKII (Stardust, Osaka, Japan), followed by passing through 250 μm sieve. Finally, the sieved samples were homogenized, bottled and labelled. Each bottle was packed with approximately 40 g of food sample and stored in a refrigerator at 4 °C until SPE procedures.

**Sample extraction method.** Formerly ground and homogenized 2 g of subsamples were weighed into falcon tube (50 mL) followed by adding 10 mL water and 93 μL of internal standard AA–D3 (10 μg mL−1) into it. Falcon tubes containing sample were shaken for one hour on a tube rotator (Stuart, Staffordshire, United Kingdom). Subsequently, a HERMLE centrifuge, type Z32HK (Hermle Labortechnik GmbH, Wehingen, Germany) was used to centrífuge the sample tubes at 5000 rpm for 30 minutes. For the precipitation of co-extractives in the sample matrix, the clear sample supernatant was moved into another falcon tube and treated with 500 μL of Carrez I and II solutions, respectively. Subsequent centrifugation of the treated samples was performed at 5000 rpm for 3 min. Then, an aliquot (3 mL) of aqueous sample solution was filtered through a syringe filter (nitrocellulose membrane, pore size 0.45 μm) and loaded onto Strata™ –X–C polymeric strong cation SPE column which was connected online with Visiprep™ vacuum manifold (Supelco, Gland, Switzerland). Subsequently, the Strata™ –X–C column was eluted with water (3 mL) and the obtained eluent was loaded onto Isolute ENV + SPE column and eluted with a mixture of methanol:water (1 mL, 60:40, v/v). The sample extracts were gently evaporated to the final volume of 400 μL using Visidry™ vacuum manifold (Supelco, Gland, Switzerland) under nitrogen stream. Finally, the samples were filtered through PTFE syringe filters (0.45 μm) and moved into amber auto sampler glass vials for UPLC-MS/MS determination.

For the quantification of acrylamide in food samples, the isotope dilution method (a method of measuring the amount of chemical compounds in high matrix samples) has been employed. This method comprises the addition of known quantities of isotopically-enriched compound to the unknown analyzed samples, utilizing AA–D3 as the labeled compound was used to quantify the acrylamide in food samples. A response curve was constructed of area response ratio for m/z 55/58 (Table 4) versus the quantity of acrylamide injected with a constant quantity of [AA–D3]. This procedure offers more accurate results due to a correction of both extraction effectiveness and changes in the device response. At the beginning of the sample extraction procedure the known amount of AA–D3 standard solution was added to the food samples which allowed the quantification of the target compound33,34.

**Ultra-performance liquid chromatography-tandem mass spectrometry.** The amount of acrylamide in food products was most frequently determined by liquid chromatography-tandem mass spectrometry.
Table 4. MRM conditions applied with the triple quadrupole mass spectrometric system.* Dwell time (0.025 s) in both cases.

| Compound            | Precursor ion [M + H]^+ (m/z) | Product ion (m/z) | Collision energy (eV) | Product ion (m/z) | Collision energy (eV) |
|---------------------|-------------------------------|------------------|-----------------------|------------------|-----------------------|
| Acrylamide          | 72                            | 55               | 48                    | 44               | 43                    |
| Acrylamide-2,3,3-D_3 | 75                            | 58               | 52                    | 47               | 47                    |

(LC–MS/MS). Specifically, the LC–MS/MS has been considered a very compatible technique to the analysis of polar and non-volatile substances for instance acrylamide. In nature, the acrylamide is not sufficiently volatile and cannot be quantitatively analyzed by gas chromatography-mass spectrometry. Lately, an innovative technique based on UPLC-MS/MS has been applied for the analysis of acrylamide in food products. The UPLC-MS/MS technique offers rapid quantitative method in addition to excellent quality parameters for instance excellent linearity, precisions, selectivity and sensitivity. For the determination of acrylamide in such types of highly matrixes samples, the use of triple quadrupole mass spectrometry system functioning in multiple reaction monitoring (MRM) mode is highly suggested due to its high selectiveness. In the present study, the acrylamide in various food products were analyzed quantitatively determined by Waters Acquity® UPLC method outfitted with a thermostat column compartment, quaternary pump, vacuum degasser and thermostat auto sampler (Milford, USA). The reversed phase Waters Acquity® BEH C18 analytical column (50 mm × 2.1 mm i.d. and particle size, 1.7 μm) was used (Milford, USA) to get the separation of both acrylamide and AA-D3 analytes. The most favorable separation was attained in isocratic elution mode with mobile phase, methanol (10%) and 0.1% of formic acid in Milli-Q water (89.90%). The mobile phase flow rate and sample volume was 300 μL and 5 μL, correspondingly. In order to get the maximum instrumental sensitivity, at every ten samples injection, the BEH C18 analytical column was rinse with a mixture of methanol (50%) and water (50%). The UPLC instrument was outfitted with ion source (electrospray ionization, ESI) and mass analyzer (Quattro Premier triple quadrupole) (Micromass, Milford, USA). To get the highest instrumental response relating to the determination acrylamide and AA-D3, the mass spectrometric system (MS) was functioned in positive ionization mode and data were attained MRM mode. The ESI source working parameters were as follows: Cone voltage, 48 V; capillary voltage, 3.5 kV; desolvation temperature, 350°C; source temperature, 120°C; desolvation gas flow rate, 600 L h⁻¹ and cone gas flow rate, 60 L h⁻¹. For cone and collision gases, nitrogen gas of high purity (99.99%), produced by a nitrogen generator (model NM30LA, Peak Scientific, Inchinnan, United Kingdom) and argon were used, correspondingly. To supply the primary vacuum to the MS instrument, an Oerlikon rotary pump (model SOGEVACSV40 BI, Cedex, France) was used. The MS/MS conditions for instance dwell times, collision energy voltages and the precursors and daughter ions associated to the chosen transitions of acrylamide and AA–D3 are demonstrated in Table 4. For the estimation of acrylamide and AA–D3, the most abundant daughter ions were chosen nevertheless, for the confirmation of acrylamide and AA–D3, the second-most abundant daughter ions were selected. The Waters MassLynx V4.1 software (Milford, USA) was used for the data acquisition.

References

1. Swedish National Food Administration, Information about Acrylamide in Food. (2002). https://www.livsmedelsverket.se/en/sok/3q=acrylamide (Date of access: 01/11/2016).
2. Messas, M. & Morales, F. J. Acrylamide in commercial potato crisps from Spanish market: Trends from 2004 to 2014 and assessment of the dietary exposure. Food Chem. Toxicol. 81, 104–110 (2015).
3. Mottram, D. S. & Wedzicha, B. L. Acrylamide is formed in the Maillard reaction. Nature 419, 448–449 (2002).
4. Tareke, E. et al. Acrylamide: a cooking carcinogen? Chem. Res. Toxicol. 13, 517–49 (2000).
5. LoPachin, R. M. Jr. & Lehning, E. J. Acrylamide-induced distal axon degeneration: a proposed mechanism of action. Neurotoxicology 15, 247–259 (1994).
6. Dearfield, K. L. et al. Acrylamide: a review of its genotoxicity and an assessment of heritable genetic risk. Mutat. Res. 330, 71–99 (1995).
7. Costa, L. G. et al. Comparative studies on the neuro- and reproductive toxicity of acrylamide and its epoxide metabolite glycidamide in the rat. Neurotoxicology 13, 219–224 (1992).
8. IARC, International Agency for Research on Cancer, Monographs on the Evaluation of Carcinogenic Risks to Humans: Some Industrial Chemicals No. 60. IARC, Lyon, France (1994). http://monographs.iarc.fr/ENG/Monographs/vol60/vol60-16.pdf (Date of access: 01/11/2016).
9. EFSA, European Food Safety Authority, Draft scientific opinion on acrylamide in food (2014). http://www.efsa.europa.eu/en/consultations/call/140701.pdf (Date of access: 12/10/2016).
10. Olmez, H. et al. A survey of acrylamide levels in foods from the Turkish market. J. Food Compos. Anal. 25, 564–568 (2008).
11. Results on acrylamide levels in food from monitoring years 2007–2009 and exposure assessment, Scientific Report of EFSA, EFSAJ. Food Chem. Toxicol. 48, 2113 (2011).
12. Delgado-Andrade, C. Maillard reaction products: some considerations on their health effects. Clin. Chem. Lab. Med. 52, 53–60 (2014).
13. Gilbert, P. A. & Khokhar, S. Changing dietary habits of ethnic groups in Europe and implications for health. Nutr. Rev. 66, 203–215 (2008).
14. Wyka, J. et al. Estimation of dietary exposure to acrylamide of Polish teenagers from an urban environment. Food Chem. Toxicol. 75, 151–155 (2015).
15. U.S. Food and Drug Administration, Exposure Assessment for Acrylamide (2006). http://www.cfsan.fda.gov/dms/acrydata.html (Date of access: 10/05/2016).
16. IRMM, Institute for Reference Materials and Measurements, Acrylamide Level Monitoring Database (2008). http://www.irmm.jrc.be/html/activities/acrylamide/database.htm (Date of access: 10/10/2016).
17. FAO/WHO, 2011, Joint FAO/WHO Expert Committee on Food Additives: Evaluation of certain contaminants in food report from seventy-second meeting (Rep. No. WHO technical report series; No. 959).
18. EI-Assouli, S. M. Acrylamide in Selected Foods and Genotoxicity of Their Extracts. *Egypt Public Health Asso.* 84, 371–392 (2009).
19. Atwa, M. A. *et al.* Acrylamide levels in heat-treated Egyptian foods. *J. Food Dairy Sci.* 1, 69–84 (2010).
20. Bagdonaitė, K., Derler, K. & Murkovic, M. Determination of acrylamide during roasting of coffee. *J. Agric. Food Chem.* 56, 6081–6086 (2008).
21. European Food Safety Authority (EFSA). Update on acrylamide levels in food from monitoring years 2007 to 2010. *EFSA Journal* 10, 2938 (2012).
22. European Commission (EC). Council Directive 98/83/EC of November 1998 on the quality of water intended for human consumption. OJEC L330, 32–54 (1998).
23. European Food Safety Authority (EFSA). Results on acrylamide levels in food from monitoring years 2007–2009 and Exposure assessment. *EFSA Journal* 9, 2133 (2011).
24. Guenther, H. *et al.* Acrylamide in coffee: review of progress in analysis, formation and level reduction. *Food Addit. Contam.* 24, 60–70 (2007).
25. U.S. Food and Drug Administration, survey data on acrylamide in food: individual food products (2006). http://www.fda.gov/food/foodborneillnesscontaminants/chemicalcontaminants/ucm053549.htm (Date of access: 29/10/2016).
26. Andrzejewski, D. *et al.* Analysis of coffee for the presence of acrylamide by LC-MS/MS. *J. Agric. Food Chem.* 52, 1996–2002 (2004).
27. Alves, R. C. *et al.* Acrylamide in espresso coffee: Influence of species, roast degree and brew length. *Food Chem.* 119, 929–934 (2010).
28. Ölmez, H. *et al.* A survey of acrylamide levels in foods from the Turkish market. *J. Food Compos. Anal.* 21, 564–568 (2008).
29. Liu, J. *et al.* Quantitative analysis of acrylamide in tea by liquid chromatography coupled with electrospay ionization tandem mass spectrometry. *Food Chem.* 108, 760–767 (2008).
30. Mizukami, Y. *et al.* Analysis of acrylamide in green tea by gas chromatography–mass spectrometry. *J. Agric. Food Chem.* 54, 7370–7377 (2006).
31. European commission, recommendation of 10.1.2011 on investigations into the levels of acrylamide in food (2011). https://ec.europa.eu/food/sites/food/files/safety/docs/cs_contaminants_catalogue_acrylamide_recommendation_10012011_food_en.pdf (Date of access: 29/09/2016).
32. Rashidinejad, A., Birch, E. J. & Everett, D. W. Antioxidant activity and recovery of green tea catechins in full-fat cheese following gastrointestinal simulated digestion. *J. Food Compos. Anal.* 48, 13–24 (2016).
33. Bermudo, E. *et al.* Determination of acrylamide in foodstuffs by liquid chromatography ion-trap tandem mass–spectrometry using an improved clean-up procedure. *Anal. Chim. Acta* 559, 207–214 (2006).
34. Bermudo, E. *et al.* Liquid chromatography coupled to tandem mass spectrometry for the analysis of acrylamide in typical Spanish products. *Talanta* 76, 389–394 (2008).
35. Naushad, M., Khan, M. R. & Alothman, Z. A. History and Introduction of UPLC/MS. In: Naushad, M., & Khan, M. R., editors, *Ultra Performance Liquid Chromatography Mass Spectrometry Evaluation and Applications in Food Analysis*, Chapter 1. CRC Press, Taylor & Francis Group, New York, pp. 1–15 (2014).
36. Zhang, Y. *et al.* An improved method validation for rapid determination of acrylamide in foods by ultra-performance liquid chromatography combined with tandem mass spectrometry. *J. Chromatogr. A* 1142, 194–198 (2007).

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**Author Contributions**

M.R.K. wrote the manuscript and researched data. Z.A.A. and M.N. reviewed/edited manuscript. A.K.A., I.H.A. and M.S.A. contributed to the experimental works. S.M.A. reviewed the manuscript and data. All authors reviewed the manuscript.

**Additional Information**

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