Acrylamide in food: mechanisms of formation and influencing factors during heating of foods

Hans Lingnert¹, Spiros Grivas², Margaretha Jägerstad³, Kerstin Skog⁴, Margareta Törmqvist⁵ and Per Åman⁶

¹SIK, The Swedish Institute for Food and Biotechnology, Göteborg, Sweden; ²Department of Biosciences, Karolinska Institute and Södertörn University College, Stockholm, Sweden; ³Department of Food Science, University of Agricultural Sciences, Uppsala, Sweden; ⁴Department of Applied Nutrition and Food Chemistry, University of Lund, Sweden; ⁵Department of Environmental Chemistry, University of Stockholm, Sweden

Abstract

Background: In April 2002, the Swedish National Food Administration and a scientific group at the University of Stockholm jointly announced that they had shown acrylamide to be formed during the preparation of food and found it to occur in many foodstuffs. These new findings were clearly of concern to many types of industrial food processing as well as to home cooking. The Swedish Food Federation (Li) initiated and financed the formation of an expert committee to look into the chemical mechanisms. The present review is the final report of that expert committee.

Design: The study identified, examined and put together facts and present knowledge on reaction routes for acrylamide formation in food and causal connections to cooking and food processing conditions. The results are based on literature surveys, examination of the analytical data published by the Swedish National Food Administration and other follow-up studies, contacts with international scientific networks, and observations from food companies.

Results: The exact chemical mechanism(s) for acrylamide formation in heated foods is unknown. Several plausible mechanistic routes may be suggested, involving reactions of carbohydrates, proteins/amino acids, lipids and probably also other food components as precursors. With the data and knowledge available today it is not possible to point out any specific routes, or to exclude any possibilities. It is likely that a multitude of reaction mechanisms is involved. Acrolein is one strong precursor candidate, the origin of which could be lipids, carbohydrates or proteins/amino acids. Acrylamide is a reactive molecule and it can readily react with various other components in the food. The actual acrylamide level in a specific food product, therefore, probably reflects the balance between ease of formation and potential for further reactions in that food matrix. There are indications in support of that the Maillard reaction being an important reaction route for acrylamide formation, but lipid degradation pathways to the formation of acrolein should also be considered.

Conclusions: Reliable analytical methods to measure acrylamide in foods are available. Model studies are needed to identify precursors and reaction route(s) based on current hypotheses and to elucidate possible further reactions between acrylamide and other food components. Studies are needed to optimize formulation and processing conditions to minimize acrylamide levels, taking other product quality properties into consideration.

Keywords: Acrylamide, cooking, food processing, heated foods, Maillard reaction.

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Acrylamide: some basic facts

Synonyms: 2-propenamide, ethylene carboxamide, acrylic acid amide, vinyl amide, propenoic acid amide;
CAS no.: 79-06-1. Molecular mass: 71.09.
Acrylamide appears as a white crystalline solid, is odourless and has high solubility in water (2155 g l⁻¹ water). Its melting point is 84.5°C, and its boiling point (25 mmHg) 125°C (192.6°C at atmospheric pressure).

Acrylamide is a reactive chemical, which is used as a monomer in the synthesis of polyacrylamides used, for example, in the purification of water and in the formulation of grouting agents. Acrylamide is known as a component in tobacco smoke.

Acrylamide is primarily reactive through its ethylenic double bond. Polymerization of acrylamide occurs through radical reactions with the double bond. Acrylamide could also react as an electrophile by 1,4-addition to nucleophiles, e.g. SH- or NH₂-groups in biomolecules.

Acrylamide is metabolized in the body to glycidamide, a reactive compound formed through epoxidation of the double bond.

The toxicological effects of acrylamide have been studied in animal models. Exposure to acrylamide leads to DNA damage and at high doses neurological and reproductive effects have been observed. Carcinogenic action in rodents has been described, but carcinogenicity to humans has not been demonstrated in epidemiological studies, although it cannot be excluded. The International Agency for Research on Cancer (IARC) has classified acrylamide as “probably carcinogenic to humans” (group 2A). Neurological effects have been observed in humans exposed to acrylamide. The properties, use and toxic effects of acrylamide are reviewed by IARC (1) and the European Union (EU) (2).

Background: how was acrylamide formation in foods observed?

Reaction product from acrylamide observed in humans

Compounds that are reactive and therefore short lived in the body can be demonstrated through their stable reaction products (adducts) with biomacromolecules, e.g. haemoglobin (Hb) in blood. The adducts to Hb are accumulated during the life span of the protein (about 4 months in humans). Hb adducts are not indicators of toxic action but could be used for exposure measurements and calculation of intake.

A sensitive method for specific measurement of adducts to the N-terminal valines in the globin chains in Hb and for analysis by gas chromatography–mass spectrometry (GC-MS) has been developed and applied to a wide range of compounds (3, 4). Application of this methodology has shown that an adduct from acrylamide is formed by Michael addition to the ethylenic double bond to N-terminal valine (5). This adduct, N-(2-carboxamidomethyl)valine, has been measured in blood from acrylamide-exposed humans (5–8) and animals (9). In studies of occupational exposure it has been shown that the adduct also occurs in blood from persons without known exposure (6, 8), although at higher levels in smokers (since acrylamide occurs in cigarette smoke).

In connection with studies of the leakage of acrylamide at the Hallandsvägen tunnel construction in Sweden, calculations of uptake of acrylamide and evaluation of cancer risk were performed (10). The calculations of uptake (from pharmacokinetic modelling and reaction kinetics) showed that the average background adduct level in unexposed controls would correspond to a daily intake by adults of about 100 μg acrylamide, and there was a preliminary indication that this background level could be associated with a considerable cancer risk. The estimated risk seemed to be higher than the risk from “background” exposure of other reactive compounds detected as adducts in persons without known exposure. In this situation it appeared urgent to find the source of the acrylamide adducts regularly observed in non-exposed persons. The occurrence of acrylamide in tobacco smoke and the findings of lower background levels in wild animals (Törnqvist et al., to be published) led to the hypothesis that acrylamide was formed in cooking.

Acrylamide formation during cooking: identification and quantification

The identification of acrylamide in heated foodstuffs originated from a hypothesis for which both direct and indirect proof was obtained. An animal feeding experiment was performed to test the hypothesis on acrylamide formation during heating of foodstuffs. A strong increase in the acrylamide Hb adduct level in blood from rats was observed when the animals were fed fried standard feed (11). In this context the identity of the observed Hb adduct was verified by tandem mass spectrometry (GC-MS/MS) through comparison with isotope-substituted standards (11).

A GC-MS method for analysis of acrylamide in water (based on bromination) was further developed for the analysis of acrylamide in the animal feed. The content of acrylamide in the fried feed...
was measured and found to be compatible with the increase in the acrylamide Hb adduct level in the rats.

In following experiments the effect of heating (frying, etc.) on the content of acrylamide in different foodstuffs was investigated (12). The GC-MS method that had been applied in the studies of animal feed was further improved and simplified. This method, based on a well-known procedure for analysis of acrylamide in water, involves bromination of the ethylenic double bond, the dibromopropionamide formed being the analyte. The bromination is performed at ambient temperature and at pH 1–3. The GC-MS analysis is performed at raised temperature. It was found desirable to confirm the results by a milder method for analysis of underivatized acrylamide. This was achieved by the development of a liquid chromatography–tandem mass spectrometry (LC-MS/MS) method. The analytical results obtained with these two methods (GC-MS and LC-MS/MS) are in full agreement. Analysis at different conditions with the two methods further supported the conclusion that acrylamide is the analyte. It was shown that acrylamide was formed in a temperature-dependent manner in food. Low contents of acrylamide were found in heated protein-rich foods (5–50 μg kg⁻¹) and high contents in carbohydrate-rich foods (100–4000 μg kg⁻¹), compared with undetectable levels in unheated or boiled foods.

The fact that in the above animal experiment the formed adduct levels tallied with the dietary intake of acrylamide as analysed in the heated feed, further supports the view that we are really dealing with acrylamide. Similar conclusions can be drawn from adduct levels in humans and human consumption on heated foods (12).

This work was carried out in a collaborative project between Stockholm University and AnalyCen Nordic AB, with development of methods and analysis of acrylamide in food in the latter laboratory. In view of the results obtained, showing high acrylamide contents in carbohydrate-rich commercial foods, the Swedish National Food Administration, in parallel work, developed an LC-MS/MS method for acrylamide analysis in food (13). They essentially verified the results and extended the study to a broader range of foodstuffs (http://www.slv.se), and also showed that there was good agreement between the analyses carried out at the two laboratories (13). The results from the studies were jointly announced in Stockholm on 24 April 2002.

Subsequent work verifying and extending the analyses of acrylamide in food, mostly analysed with LC-MS/MS, has been presented from several Food Authorities and other organizations in different countries (e.g. the UK, Norway, the Netherlands, Switzerland, Germany and the USA).

Chemical mechanisms for acrylamide formation

Food scientists and technologists have had an interest in acrylamide (and/or its derivatives, including polymers), its applications and its possible toxic effects for many years. For example, there are many reports on can coatings and food packaging, food additives (preservatives, artificial sweeteners, etc.) and acrylamide polymers of suitable quality with low residual acrylamide monomer levels that are used in, for example, the USA for treatment of poultry, potato, corn and other wastes, with the resulting concentrated solids used as components of blended animal feeds (14–19).

There are only a few earlier reports on the occurrence of acrylamide in food. For example, acrylamide has been reported to be present in plant material (potatoes, carrots, radish, lettuce, Chinese cabbage, parsley, onions, spinach and rice paddies) (20). In 1 g plant samples, 1.5–100 ng acrylamide could be detected. Acrylamide was also reported to occur in sugar (21). The origin of the detected acrylamide in these foods is not known; it may be exogenous.

To the authors’ knowledge, no proposed or proven reaction routes for the formation of acrylamide during food processing have been published. Therefore, described below are the hypotheses that were found to be most relevant and probable in a food processing situation.

(A) Acrolein (2-propenal, CH₂=CH=CHO) is a three-carbon aldehyde and thus recalls the structure of acrylamide [CH₂=CH–C(O)─NH₂]. Further, acrolein is known to be formed by:

- transformation of lipids;
- degradation of amino acids and proteins;
- degradation of carbohydrates;
- the Maillard reaction (MR) between amino acids or proteins and carbohydrates.

Therefore, acrolein is a very probable precursor of acrylamide. Simple, fundamental chemical transformations (such as reaction with ammonia liberated from amino acids) can then convert acrolein (or a derivative of it) into acrylamide. The produc-
tion of acrylamide through the reaction of acrolein with ammonia can be anticipated.  
(B) Alternative formation mechanisms of acrylamide do not necessarily involve acrolein. For example, proteins and/or amino acids can, after a series of transformations, such as hydrolyses, rearrangements and decarboxylations, eventually lead to acrylamide.

Processes A and B are complicated and involve multistage reaction mechanisms which may also include free radical reactions to acrolein or acrylamide (23–25).

**Acrolein formation from lipids**

When oil is heated at temperatures above the smoke point, glycerol is degraded to acrolein, the unpleasant acrid black and irritating smoke (26–29). The formation of acrolein is known to increase with the increase in unsaturation in the oil and to lead to a lowering of the smoke point. The smoke point is higher for oils with a higher content of saturated fatty acids and lower content of polyunsaturated acids. The smoke points for some of the main oils and fats are: palm 240°C, peanut 220°C, olive 210°C, lard and copra 180°C, sunflower and soybean 170°C, corn 160°C, margarine 150°C and butter 110°C. Usually, the smoke starts to appear on the surface of heated oils before their temperature reaches 175°C. The oil is first hydrolysed into glycerol and fatty acids and then acrolein is produced by the elimination of water from glycerol by a heterolytic acid-catalysed carbonium ion mechanism followed by oxidation (30):

$$\text{CH}_2\text{(OH)}\text{CH(OH)}\text{CH}_2\text{(OH)} \rightarrow \text{CH}_2\text{CH=CHO}$$

Besides the above-mentioned mechanism for the formation of acrolein from acylglycerols, acrolein can also be produced as a result of oxidation of polyunsaturated fatty acids and their degradation products (31–34). Several aldehydic products (including malondialdehyde, C3–C10 straight chain aldehydes, and \(\alpha,\beta\)-unsaturated aldehydes, such as 4-hydroxynonenal and acrolein) are known to form as secondary oxidation products of lipids (35). Acrolein was also found to form in vivo by the metal-catalysed oxidation of polyunsaturated fatty acids, including arachidonic acid (36).

**Acrolein formation from amino acids, proteins and carbohydrates**

Several sources for the formation of acrolein are known. It may arise from degradation of amino acids and proteins (37, 38), degradation of carbohydrates (39), and the MR between amino acids or proteins and carbohydrates (40, 41). Many possible routes for the formation of this three-carbon aldehyde, taking the starting point from many different sugars or amino acids, may be proposed. Its formation from methionine by the Strecker degradation in the frame of the MR is one example. Alanine, with its three-carbon skeleton, has also been suggested as a possible source. However, fission reactions of longer carbon chains are common and well known, so at present there is no basis to give priority to any specific reaction routes.

**Formation of acrylamide through amino acid reactions not involving acrolein**

There are also numerous, plausible reaction routes by which amino acids (or proteins) may form acrylamide without going through acrolein. Within the frame of complex, multistage reaction mechanisms, involving hydrolyses, rearrangements, decarboxylations, deaminations, etc., many specific mechanistic pathways may be suggested. Decarboxylation and deamination of aspartagine, and transformations of dehydroalanine (formed from, e.g. serine or cysteine) are some examples of reaction routes that have been proposed. However, also in this case these can only be seen as possible examples, and similarly to above, there is no basis to give priority to any specific routes.

**Conclusion**

Since no systematic studies have been performed or reported, there is at present no evidence to indicate any specific reaction routes for acrylamide formation, or to exclude any possibilities. It is most likely that a multitude of reaction mechanisms is involved, depending on food composition and processing conditions.

**Further reactions of formed acrolein and acrylamide**

As mentioned above, acrolein can be converted into acrylamide by a series of fundamental reactions. However, both acrolein and acrylamide are reactive, because of their double bonds and the amino group of acrylamide. They can readily react further with other reactive groups present in the food matrix or formed during the heating process. For example, acrylamide can react with small reactive molecules, such as urea [CO(NH)\(_2\)] and form-
Acrylamide in foods

aldehyde (HCHO), or with glyoxal [(CHO)₂], aldehydes (RCHO), amines (R₂NH), thiols (RSH), etc. Furthermore, the products shown in the following scheme can even react further in the same mode of reaction:

These types of reactive functional group may also be found in macromolecules, such as proteins. (See adduct formation with valine in the globin chain of haemoglobin, described above. In haemoglobin adducts are formed not only with valine, but also with, e.g. cysteine.) The presence or absence of reactive groups (or its concentration) in the food matrix may thus be one explanation for differences in final acrylamide content in different food systems. The resulting acrylamide level may be due to a balance between formation and further reactions. The low acrylamide levels in heated meat products could, for instance, depend on adduct formation between acrylamide (or acrolein) and proteins.

Factors with a possible influence on acrylamide formation

A couple of different chemical mechanisms for the formation of acrylamide has been outlined above. As long as the mechanisms are not confirmed, the influencing factors cannot be established. Thus, what is presented here are attempts to identify what factors would be of importance (regarding processing conditions or product composition) if a specific reaction route were the prevailing one. Specific emphasis is placed on the MR, since this reaction system involves many of the basic carbohydrate and amino acid reactions. Another major reaction in foods during processing, which could be of importance, is lipid hydrolysis followed by oxidation of the fatty acids.

Acrolein formation from lipids

Acrolein may be formed from the glycerol part of triglycerides or through oxidation of fatty acids. This means that factors favouring lipid hydrolysis as well as factors favouring lipid oxidation would promote acrolein formation. Temperature is an important factor for both of these reactions. Regarding hydrolysis, pH may also be of importance and high as well as low pH may be supposed to favour acrolein formation. Regarding oxidation, lipid composition is of key importance: the higher the degree of unsaturation, the lower the stability. Protection against oxygen and light will limit the oxidation, and pro-oxidants, such as metals, should be avoided. The protective effect of antioxidants should also be taken into account.

The Maillard reaction as the route for acrylamide formation

The MR has been proposed as a route for acrolein formation. The direct formation of acrylamide through amino acid transformations has also been proposed. These amino acid transformations also involve reactions common in the MR system.

Basics of the Maillard reaction

The MR is one of the most important chemical
reactions in food processing, with an influence on several aspects of food quality. Flavour, colour and nutritional value may be affected and certain reaction products have been noticed to be antioxidative, antimicrobial, genotoxic, etc. The practical applications of Maillard chemistry in food processing are, therefore, a matter of balance between favourable and unfavourable effects, and the aim of the food manufacturer is to find an optimum in this balance. This may be accomplished by influencing the main variables affecting the MR (42).

The MR takes place in three major stages and is dependent on factors such as concentrations of reactants and reactant type, pH, time, temperature and water activity. Free radicals and antioxidants are also involved (43).

The early stage (step 1) involves the condensation of a free amino group (from free amino acids and/or proteins) with a reducing sugar to form Amadori or Heyns rearrangement products. The advanced stage (step 2) means degradation of the Amadori or Heyns rearrangement products via different alternative routes involving deoxyosones, fission or Strecker degradation. A complex series of reactions including dehydration, elimination, cyclization, fission and fragmentation results in a pool of flavour intermediates and flavour compounds. Following the degradation pathway as illustrated schematically in Fig. 1, key intermediates and flavour chemicals can be identified.

One of the most important pathways is the Strecker degradation, in which amino acids react with dicarbonyls (formed by the MR) to generate a wealth of reactive intermediates. Typical Strecker degradation products are aldehydes, e.g. formaldehyde, acetaldehyde and possibly propenaldehyde (acrolein). Strecker degradation results in degradation of amino acids to aldehydes, ammonia and carbon dioxide (44) and takes place in foods at higher concentrations of free amino acids and under more drastic reactions, e.g. at higher temperatures or under pressure (45).

The final stage (stage 3) of the MR is characterized by the formation of brown nitrogenous polymers and co-polymers. While the development of colour is an important feature of the reaction, relatively little is known about the chemical nature of the compounds responsible. Colour compounds can be grouped into two general classes: low molecular weight colour compounds, which comprise two to four linked rings, and the melanoids, which have much higher molecular weights.

Review of factors influencing the Maillard reaction
Factors that are particularly important for the MR are the starting reactants, e.g. type of sugar and amino acid (protein), temperature, time and water activity. The presence of metal salts (pro-oxidants), and inhibitors, such as antioxidants and sulfate, may have an impact.
Starting reactants: reducing sugar and amino acids/proteins. The MR requires reducing sugars, i.e. sugars containing keto- or aldehydes (free carbonyl groups). The reactivity of different sugars can be summarized in the following way (46):

- The shorter the carbon chain of the sugar, the greater the lysine losses (MR).
- Pentoses are more reactive than hexoses and disaccharides in yielding brown colour.
- Aldoses are more reactive than ketoses both in aqueous solution model systems and at storage (low water content).
- Among isomeric sugars, stereochemistry is important. Thus, ribose is more reactive than xylose monitored as lysine losses.

All monosaccharides are reducing sugars. (Sugar alcohols do not participate in the MR.) Among the disaccharides all sugars except for sucrose are reducing sugars. In oligosaccharides and starch only the end-terminal monosaccharide is a reducing sugar. Starch and sugars, such as sucrose, lactose and maltose, can easily hydrolyse upon heating above 100°C at slightly acidic pH, resulting in the formation of monosaccharides (reducing sugars). Thus, thermal processing often results in a continuous supply of reducing sugar formed from complex carbohydrates.

Most studies concerning the reactivity of amino acids have been performed on free amino acids in diluted aqueous solutions. The reactivity among the diamino acids increased with the length of the carbon chain. Among the amino acids studied lysine was most reactive. In proteins and peptides, only free amino groups can react, i.e. N-terminal α-amino groups and Ω-amino groups.

Temperature and time. The temperature dependence of chemical reactions is often expressed as the activation energy ($E_a$) in the Arrhenius equation. The higher the value of $E_a$, the more temperature dependent the reaction rate. Activation energy data for the MR have been reported within a wide range, 10–160 kJ mol$^{-1}$, depending on, among other things, water activity and pH and what effect of the reaction has been measured. The temperature dependence of the MR is also influenced by the participating reactants. The temperature effect is also affected by the other variables, and different aspects of the MR thus differ in temperature dependence (42).

Water. Water has both an inhibitory and an accelerating impact on the MR. Water acts partly as a reactant and partly as a solvent and transporting medium of reactants (reactant mobility). In the initial steps of the MR, 3 moles of water are formed per mole of carbohydrate. Thus, the reaction occurs less readily in foods with a high water activity ($a_w$) value. Water may depress the initial glucosylamine reaction, but enhance the deamination step later in the reaction.

The results from studies in model systems for optimal water concentration or $a_w$ (free water) or relative humidity (RH) vary markedly depending on selected reactants and how the MR is evaluated: as loss in lysine or browning intensity. Several studies have been performed, of which most claim the maximum $a_w$ to be between 0.3 and 0.7 (47). However, most data on the influence of $a_w$ are based on studies at relatively low temperatures (30–60°C). At higher temperature, more relevant to heat processes, considerably lower $a_w$ has been shown to be favourable to the MR (42).

The main explanation for an optimum reaction rate at an intermediate $a_w$ is that the reactants are diluted at the higher $a_w$, while at a lower $a_w$ the mobility of reactants is limited, despite their presence at increased concentrations.

pH. The MR itself has a strong influence on pH. Therefore, aqueous model systems based on reflux boiling of sugars and amino acids need to be buffered since the pH quickly drops from 7 to 5. Low pH values (<7) favour the formation of furfurals (from Amadori rearrangement products), while the routes for reductones and fission products are preferred at a high pH. However, the overall effect of pH is not clear cut, since the reactions take place by all three pathways. In unbuffered water solutions, pH increase during the MR and buffering with alkali have a catalytic effect.

The reactivity of different amino acids at various pH has been studied. Browning of a glucose solution upon heating was obtained first when pH exceeded 5 and it increased with increasing pH. The degree of browning varied with the position of the amino group. The function of pH is linked with specific reaction steps of the MR. Initially, only non-protonized forms of amino acids can form Schiff base. This explains the pronounced change in reactivity (monitored as browning) that happens when pH passes the isoelectric point of the amino group in the reacting amino acid. Thus, optimal pH for the MR varies with the system used and how the reaction is monitored (e.g. lysine losses or browning).

Inhibition of the Maillard reaction. Measures to inhibit the MR in cases where it is undesirable.
involves lowering the pH value, maintenance of the lowest possible temperatures and avoidance of critical water contents (moistures below 30%, during processing and storage), use of non-reducing sugars and addition of sulfite (45). The use of the inhibitor sulfur dioxide constitutes an important way of controlling the MR. It may combine with early intermediates. However, sulfite only delays colour formation and it is interesting to note that the colour formed in sulfite-treated systems is less red and more yellow than in untreated systems.

Maillard reactions and food processing

In exploiting the MR, the key target for the food industry is to understand and harness the reaction pathways enabling the improvement of existing products and the development of new products. While it would be easy to assume that this means the generation of flavour and colour, not all Maillard products endow positive characteristics to foods and ingredients. The positive contributions of the MR are flavour generation and colour development. The negative aspects are off-flavour development, flavour loss, discoloration, loss of nutritional value and formation of toxic Maillard reaction products (MRPs). In applying the MR, there are challenges that are common to the food industry, independent of the type of the product. These challenges can be classified: maintenance of raw material quality, maintenance of controlled processes for food production, maintenance of product quality and extension of product shelf-life (42, 43).

Flavour/aroma. The most common route for formation of flavours via the MR comprises the interaction of α-dicarbonyl compounds (intermediate products in the MR, stage 2) with amino acids through the Strecker degradation reactions. Alkyl pyrazines and Strecker aldehydes belong to commonly found flavour compounds from the MR. For example, low levels of pyrazines are formed during the processing of potato flakes when the temperature is less than 130°C, but increase 10-fold when the temperature is increased to 160°C, and decrease at 190°C, probably owing to evaporation or binding to macromolecules. The aroma profile varies with the temperature and the time of heating. At any given temperature–time combination, a unique aroma, which is not likely to be produced at any other combination of heating conditions, is produced. Temperature also affects the development of aroma during extrusion cooking.

Colour. The coloured products of the MR are of two types: the high molecular weight macromolecule materials commonly referred to as the melanoidines, and the low molecular weight coloured compounds, containing two or three heterocyclic rings (48). Colour development increases with increasing temperature, time of heating, increasing pH and intermediate moisture content \(a_w = 0.3–0.7\). In general, browning occurs slowly in dry systems at low temperatures and is relatively slow in high-moisture foods. Colour generation is enhanced at \(\text{pH} > 7\). Of the two starting reactants, the concentration of reducing sugar has the greatest impact on colour development. Of all the amino acids, lysine makes the largest contribution to colour formation and cysteine has the least effect on colour formation.

Antioxidative capacity. There are several reports on the formation of antioxidative MRPs in food processing. The addition of amino acids or glucose to cookie dough has been shown to improve oxidative stability during the storage of the cookies. Heat treatment of milk products before spray drying has been reported to improve storage stability, as has heat treatment of cereals (42).

The antioxidant effect of the MRP has been extensively investigated (49). It has been reported that the intermediate reductone compounds of MRP could break the radical chain by donation of a hydrogen atom: MRP was also observed to have metal-chelating properties and retard lipid peroxidation. Melanoidines have also been reported to be powerful scavengers of reactive oxygen species (50). Recently, it was suggested that the antioxidant activity of xylose–lysine MRPs may be attributed to the combined effect of reducing power, hydrogen atom donation and scavenging of reactive oxygen species (51).

Nutritive value. Loss of protein quality is often associated with the MR, especially in cereal products and milk powder produced by heat treatment. Usually the essential amino acid having an extra free amino group, e.g. lysine, is most vulnerable. If the essential amino acid is also the nutritionally limiting amino acid, the influence of the MR on the protein quality is substantial. This is not a problem in cooking meat and fish, since these food items are very rich in protein. Loss of protein quality in terms of nutritional value is a more serious problem for heat treatment and dehydration, especially of cereals, milk and their mixtures (breakfast cereals, gruels, bread, biscuits), since carbohydrates
dominate over proteins in these food items and the proteins levels are also generally low.

Toxic effects. The possibilities that MRPs could be mutagenic and/or carcinogenic were explored with the Ames test around 20–25 years ago. In general, weak genotoxicity/mutagenic activities were found for known MRPs. Most attention over the past few decades has been paid to the food mutagens found in the crust from cooked meat and fish. Chemically, these compounds belong to a class of heterocyclic amines, currently amounting to around 20 different species. Most of them have been classified as possible food carcinogens (group 2B) by the IARC, based on long-term studies on rodents. The precursors of the heterocyclic amines are free amino acids and, for more than half of the 20 species, also creatine (a natural energy metabolite present in muscle cells only). Reducing sugars up to equimolar amounts compared with amino acids and/or creatine enhance the yields of heterocyclic amines markedly.

Thus, the MR and/or pyrolysis have been claimed to be important mechanisms for the formation of these heterocyclic amines, where Strecker aldehydes, pyrazines or pyridines and creatine have been suggested to play an important role. The yields of these food-borne carcinogens increase with time and temperature, especially from 150°C and above. The highest concentrations of heterocyclic amines are found in the crust of pan-fried, grilled or barbecued meat and fish. In addition, gravies prepared from dried meat juice collected from pan residues or oven roasting could be rich in heterocyclic amines. Pro-oxidants, water activity in the optimal range for the MR and high temperatures (200–400°C) enhance their yield. The average daily exposure for heterocyclic amines is around 0.5 µg per person, with a range between 0 and 20 µg. Antioxidants, an excess of carbohydrates, cooking temperatures below 200°C and moisture contents above 30% reduce the occurrence of heterocyclic amines. Moreover, heterocyclic amines rarely occur in plant foods, even during well-done cooking (52).

To the authors’ knowledge, no reports in the literature so far have studied acrylamide formation linked with the MR.

Conclusions and ideas from data and observations presented so far

No systematic studies of acrylamide content in food products have yet been published. The Swedish National Food Administration reported results from their analyses on 23 April 2002, with some new results added on 26 April (http://www.slv.se/HeadMenu/livsmedelsverket.asp). In most cases the figures reported referred only to one single, randomly selected package of each specific product.


In all studies single, randomly selected samples were analysed. The main original results reported by the Swedish National Food Administration were confirmed by the later studies, but it must be concluded that substantial variations are found within a given food group and in cases when repeated analyses of the same product have been performed considerable variations have been found between samples. This makes the data premature as a basis for conclusions on the mechanisms for acrylamide formation. An attempt to summarize the results published so far is presented in Table 1.

Table 1, presenting hitherto reported acrylamide data of heat-treated foods, can only give indications as to which factors are important in acrylamide formation. Table 1 indicates that high temperatures are needed for the acrylamide to form. No acrylamide formation has so far been demonstrated at temperatures below 100°C and it is probable that the products reported have reached temperatures well above this level. There are several examples of exaggerated acrylamide formation on overheating. Fried products of plant origin seem to give the highest concentrations, but frying fat is not a prerequisite for acrylamide formation.

The data indicate strongly that acrylamide formation is mainly a surface phenomenon. This has also been verified by other data presented to the authors by companies. This implies that water activity may be an important factor, although there are strong links between temperature and water activity in a frying or baking process.
From the analyses reported it is very difficult to draw conclusions on variations between different plant raw materials. Corn products are possibly lower in acrylamide content than comparable products made from potato or other cereals. However, it is difficult to explain these possible differences or draw conclusions on reaction mechanisms, since detailed data on chemical composition (reducing sugars, specific amino acids, etc.) are lacking. This is a general difficulty. There are insufficient details on chemical composition to be able to suggest which are the important precursors and formation mechanisms.

Some data and observations, however, indicate that acrylamide formation is increased by increased concentration of (reducing) sugar in the raw materials or ingredients. This strongly supports the MR mechanisms. The MR hypothesis is also supported by several other observations, parallel to browning, the influence of temperature and water activity, etc. In fried products the proposed route via acrolein formed from lipids should also be considered. The relatively high temperatures combined with low water activity which favour acrylamide formation, also favour free radical reactions. Thus, antioxidants and other free radical scavengers or quenchers could act as inhibitors.

The MR occurs wherever non-enzymic browning is induced by heat treatment, e.g. extrusion cooking, roasting, popping, baking, pan-frying, deep-fat frying, barbecuing and autoclavage. Most unprocessed foods contain the necessary starter reactants, i.e. amino acids/proteins and reducing sugar. Conventionally cooked foods are subjected to a relatively high temperature for a relatively long time, and the surface of the food dries out to give a crust with a low $a_w$, favouring the MR.

It is also clear from the composition of cereals and potato (see below), and most other raw food material, that they contain all necessary precursors, e.g. protein/free amino acids, carbohydrates/sugars and lipids, to initiate both MRs and lipid oxidation or other degradation routes during processing and storage. The extent to which these foods contain the optimal proportion of precursors and modifying components in terms of enhancers or inhibitors for acrylamide formation remains to be established.

In this context one must also bear in mind the high potential for acrylamides to be consumed through further reactions with other components in the food product. Similarly, acrolein or other precursors could react with other food components and take reaction routes not leading to acrylamide. Consequently, the final acrylamide level in a food product may be due to the balance between formation and further reactions, controlled by the chemical composition of that specific food. Low acrylamide levels may thus be a result of further reactions (or altered reaction routes) in that specific food matrix. As already speculated, the low acrylamide levels demonstrated in meat products could be a result of adduct formation of acrylamide with proteins or other components. All of the reaction mechanisms mentioned so far (MR, lipid hydrolysis and oxidation, etc.) are also known to proceed in meat systems.

### Table 1. Acrylamide data of heat-treated foods

<table>
<thead>
<tr>
<th>Product group</th>
<th>Acrylamide concentration (µg kg$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Typical range</td>
</tr>
<tr>
<td>Potato crisps</td>
<td>600–2000</td>
</tr>
<tr>
<td>French fries$^a$</td>
<td>300–700</td>
</tr>
<tr>
<td>Pan-fried potatoes</td>
<td>250–300</td>
</tr>
<tr>
<td>Biscuits and crackers</td>
<td>100–600</td>
</tr>
<tr>
<td>Popcorn$^b$</td>
<td>400</td>
</tr>
<tr>
<td>Crispbreads</td>
<td>50–400</td>
</tr>
<tr>
<td>Coffee (powder)</td>
<td>200</td>
</tr>
<tr>
<td>Breakfast cereals</td>
<td>50–250</td>
</tr>
<tr>
<td>Corn crisps</td>
<td>100–200</td>
</tr>
<tr>
<td>Soft breads</td>
<td>&lt; 30–50</td>
</tr>
<tr>
<td>Meat and fish products</td>
<td>&lt; 30–50</td>
</tr>
<tr>
<td>Pizza, pancakes, waffles,</td>
<td>&lt; 30</td>
</tr>
<tr>
<td>scrambled egg</td>
<td></td>
</tr>
<tr>
<td>Raw, boiled or mashed</td>
<td></td>
</tr>
<tr>
<td>potatoes</td>
<td></td>
</tr>
<tr>
<td>Pasta</td>
<td>&lt; 30</td>
</tr>
<tr>
<td>Wheat and rye flour,</td>
<td>&lt; 30</td>
</tr>
<tr>
<td>rice, oat flakes</td>
<td></td>
</tr>
<tr>
<td>Vegetarian schnitzel,</td>
<td>&lt; 30</td>
</tr>
<tr>
<td>cauliflower gratin</td>
<td></td>
</tr>
<tr>
<td>Dried fruit$^b$</td>
<td>&lt; 30</td>
</tr>
<tr>
<td>Beer (alcohol free)$^b$</td>
<td>&lt; 30</td>
</tr>
</tbody>
</table>

$^a$ For deliberately overcooked samples, values as high as 12,800 µg kg$^{-1}$ have been reported.

$^b$ One single value.

### Raw material composition

The raw material studied so far has been mainly cereals and potato, and as shown in Table 1, heat-treated products from these materials contain the highest concentrations of acrylamides, in several samples exceeding 500 µg kg$^{-1}$. These plant materials are storage organs containing large quantities of starch, protein and cell-wall materials as well as lipids, ash, polyphenols and a large number of low molecular weight compounds such as sugars and
free amino acids. It is well known that there exists a large variation in chemical composition in most plant materials. This variation is dependent on both genetic and environmental factors. The main mechanisms for acrylamide formation discussed so far are related to fat degradation and reactions involving sugars, amino acids and proteins, not the least the MR. Other components such as starch and dietary fibre may be involved by modulating the processing conditions, such as the water activity.

On a dry matter basis, wheat contains 60–73% starch, 9–16% crude protein, 2–3% fat, 2–5% sugars (glucose, fructose and fructo-oligosaccharides as well as maltose in germinated products) and 1–2% ash. The crude protein contains all the common amino acids and also significant amounts of free amino acids. Rye and dehulled oats contain less starch but more dietary fibre, especially water-soluble and viscous dietary fibre. Oats also have a higher content of fat, which in this cereal is also present in the starchy endosperm. Rye has a higher content of sugars, especially fructo-oligosaccharides. Corn has higher starch and fat contents, but a lower content of dietary fibre than wheat.

Analytical methods: can we trust the data?
The methods used to analyse acrylamide in foods were described briefly in the background section. GC-MS methods and LC-MS/MS methods have been used. Analyses have now been performed by a number of laboratories (using somewhat different methods). When the same type of product has been analysed in several countries, generally good agreement has been obtained between the results. Good correlation has also been demonstrated when identical samples have been analysed by different methods at different laboratories (13). This strongly supports that the analytical methods are reliable, that it really is acrylamide that is measured, and that the reported data can be trusted. Further support comes from the finding that Hb adduct levels in animals tallied with acrylamide intake as analysed in the feed (11). Similar conclusions can be drawn from adduct levels in humans and human consumption of heated foods (12).

The same conclusion was reached by the Food and Agriculture Organization/World Health Orga-


Conclusions
The exact chemical mechanisms of acrylamide formation in heated foods are not known. Several plausible mechanistic routes may be suggested, involving reactions of carbohydrates, proteins/amino acids, lipids and probably also other food components as precursors. With the data and knowledge available today it is not possible to point out any specific routes, or to exclude any possibilities. A multitude of reaction mechanisms is probably involved, depending on food composition and processing conditions. Acrolein is one strong precursor candidate. Current data indicate that the MR may be an important reaction route for acrylamide formation, but lipid degradation pathways to the formation of acrolein should also be considered.

Acrylamide is a reactive molecule and it can readily react with various other components in the food. The actual acrylamide level in a specific food product therefore probably reflects the balance between ease of formation and the potential for further reactions in that food matrix.

More research is needed before any firm conclusions can be drawn concerning precursors, reaction route(s) and conditions for acrylamide formation in terms of reactants, time, temperature, pH, water activity, etc. More data regarding acrylamide levels in a broader range of food products are also needed. Reliable analytical methods to measure acrylamide in foods are available.

Research needs: suggestions for further studies

- Model studies, based on current hypotheses, to identify chemical mechanisms for acrylamide formation (precursors, reaction conditions, possible inhibitors, etc.);
- model studies to elucidate possible further reactions between acrylamide and other food components;
- kinetic studies of acrylamide formation in model systems;
- studies in foods/food models: influence of processing parameters; influence of ingredients/possible precursors;
optimization of formulation and processing conditions to minimize acrylamide levels, taking into consideration other product quality properties;
continued mapping of acrylamide and acrolein in different foods;
development of simple methods to measure acrylamide in foods.

Could anything be done while awaiting the final answers?
The knowledge is still too limited to draw conclusions regarding cooking practices during industrial processing or food preparation at home. More answers from further research are needed. In the meantime the only obvious practical advice would be to avoid overheating. As long as the chemical mechanisms remain unknown, further practical recommendations are difficult to make. However, there are some indications that the MR could be involved, suggesting that factors such as levels of free (reducing) sugars and amino compounds in raw materials and ingredients should be taken into consideration. Although there is little available information on the possible influence of lipids, an interim piece of advice would be to be aware of this possibility and to control lipid degradation and oxidation in frying oils as far as possible, until more knowledge is obtained about this.

Postscript
During the publishing of this article the first reports on the chemical mechanisms of acrylamide formation were presented. They all showed that acrylamide can be formed by a reaction between amino acid and sugar, and identified asparagine as a main amino acid.


References


Hans Lingnert
SIK, The Swedish Institute for Food and Biotechnology
Box 5401
SE-402 29 Göteborg
Sweden
E-mail: hl@sik.se