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# 1 **Stability of 5-methyltetrahydrofolate in fortified apple and carrot purées**

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## 20 **Abstract**

21 5-Methyltetrahydrofolate, the naturally abundant folate vitamer, has been proposed as an alternative to  
22 folic acid for fortification. However, it is less stable than folic acid. In a formate buffer (pH 3.5), folic  
23 acid was entirely preserved after heating the solution for 3 h at 80 °C. In contrast, 5-  
24 methyltetrahydrofolate was completely degraded in less than 15 minutes. As in the buffer, 5-  
25 methyltetrahydrofolate in apple or carrot purées degraded rapidly without the addition of ascorbic  
26 acid. By adding ascorbic acid, the stability could be increased, but the chosen amount was crucial. An  
27 excess of vitamin C compared to 5-methyltetrahydrofolate was not always sufficient for the complete  
28 protection of 5-methyltetrahydrofolate during 3 h at 80°C. Only by adding 2840 µmol/kg of ascorbic  
29 acid (equivalent to 500 mg/kg), 5-methyltetrahydrofolate seemed to remain stable. Degradation started  
30 after approximately 60 minutes when 570 µmol/kg of ascorbic acid (equivalent to 100 mg/kg) were  
31 added; after 120 minutes with 1420 µmol/kg (equivalent to 250 mg/kg). In addition, a temperature  
32 decrease to 70 °C or 60 °C did not increase the stability of 5-methyltetrahydrofolate.

33 **Keywords:** folate, fortification, food matrix, degradation, impacts

## 34 **1. Introduction**

35 Folic acid fortification is linked to a reduced risk of neural tube defects in newborns—and thus  
36 became mandatory in the United States and other countries for cereal products (Eichholzer, Tonz, &  
37 Zimmermann, 2006; Grosse & Collins, 2007). In addition, the insufficient intake of folates is  
38 associated with a higher colorectal cancer risk (Kim, 2003). Folic acid, a synthetic vitamer, is  
39 predominantly used for fortification and is transformed in the body into naturally occurring forms.  
40 When ingested in high doses, however, it also circulates in its unmetabolised form (Smith, Kim, &  
41 Refsum, 2008). Furthermore, folic acid fortification in excess (>1000µg/day) can mask the vitamin B<sub>12</sub>  
42 deficiency. Thus 5-methyltetrahydrofolate is naturally abundant form in the body, it has been proposed  
43 as an alternative fortificant (Pietrzik, Bailey, & Shane, 2010; Scaglione & Panzavolta, 2014).

44 Concerning its chemical structure, 5-methyltetrahydrofolate differs from folic acid by being reduced in  
45 positions N(5), C(6) and C(7), and N(8), while hydrogen in position N(10) is substituted by a methyl  
46 group (Delchier, Herbig, Rychlik, & Renard, 2016; Scott, Rebeille, & Fletcher, 2000). This structural  
47 difference has an enormous effect on stability. Compared to folic acid, 5-methyltetrahydrofolate is  
48 highly susceptible to oxidation. In the absence of oxygen, 5-methyltetrahydrofolate is completely  
49 stable (Delchier et al., 2014). The stability of 5-methyltetrahydrofolate under aerobic conditions  
50 decreases with rising temperatures (Indrawati et al., 2004; Nguyen, Indrawati, & Hendrickx, 2003;  
51 Oey, Verlinde, Hendrickx, & Van Loey, 2006) and depends on pH. It is more stable in a neutral rather  
52 than in an acidic medium (Indrawati et al., 2004; Liu et al., 2012). At the same pH, its stability  
53 depends additionally on the buffer solution used (Indrawati et al., 2004; Paine-Wilson & Chen, 1979).

54 The degradation of 5-methyltetrahydrofolate is accelerated by fructose but not by glucose (Verlinde et  
55 al., 2010). The impact is independent of the sugar concentration in the range from 1.6 mmol/L to 1.5  
56 mol/L and is linked to a new product that is formed by the glycation of the exocyclic amino group of  
57 5-methyltetrahydrofolate (Figure 1). This product is formed along with the three degradation products  
58 that are also generated in the absence of fructose (Figure 1). The acceleration of degradation by a  
59 fructose is counteracted by ascorbic acid, probably by reduction of the first oxidation product, 5-  
60 methyl-7,8-dihydrofolate.

61 Liu et al. (2012) and Oey et al. (2006) reported that ascorbic acid decreases the degradation of 5-  
62 methyltetrahydrofolate . Oey et al. (2006) related the concentration of ascorbic acid necessary to  
63 protect 5-methyltetrahydrofolate to the initial oxygen concentration. The concentration of ascorbic  
64 acid that led to a complete protection between 50 °C and 170 °C for 15 minutes was claimed to be  
65 twice as high as the molar initial oxygen concentration (Liu et al., 2012; Oey et al., 2006). However,  
66 the relationship can be supposed to be valid only in the analysed time range as ascorbic acid is also  
67 susceptible to degradation and thus, the protection effect will be lost once the ascorbic acid is  
68 degraded.

69 Almost all groups who studied the stability of 5-methyltetrahydrofolate, used model solutions: Paine-  
70 Wilson and Chen (1979) worked with McIlvaine, HCl/KCl and citrate buffers; Nguyen et al. (2003)

71 with phosphate buffer; Oey et al. (2006) with phosphate buffer; Verlinde et al. (2010) with water and  
72 Liu et al. (2012) with HEPES/CHES buffer. Mnkeni and Beveridge (1983) reported a faster  
73 degradation of 5-methyltetrahydrofolate in tomato and apple juices than in a citrate buffer. When  
74 comparing buffers and different foods, Indrawati et al. (2004) observed but did not explain different  
75 behaviours. Delchier et al. (2014) used spinach and green bean purées but only modulated oxygen.  
76 Therefore, the mutual influence of factors (pH, oxygen, ascorbic acid, sugars) with the complex  
77 composition and physico-chemistry of foods has hardly been taken into account. In particular,  
78 dissolved oxygen in real food products may also be consumed by other oxidisable compounds, which  
79 may change the relationship. For example, 500 mg/kg of ascorbic acid increase the stability of 5-  
80 methyltetrahydrofolate in carrot juice but not in asparagus when heat-treated for 40 minutes at 120 °C  
81 (Indrawati et al., 2004). The stability difference of 5-methyltetrahydrofolate in skim milk and soy milk  
82 has been linked to the different antioxidant capacities of the two food systems (Liu et al., 2012).

83 For the present study, apple purée and carrot purée were used as fortified food matrices since they are  
84 usually less heat-treated than cereal products and are well appreciated all over Europe. They are  
85 representative fruit and vegetable foods with different pHs, fructose contents, and antioxidants. The  
86 aim was to find suitable reheating conditions in the temperature range of 60-80 °C to impart high  
87 stability to 5-methyltetrahydrofolate in apple purée and carrot purée.

## 88 **2. Material and methods**

### 89 **2.1 Chemicals**

90 Formic acid, 2-(N-morpholino)-ethanesulfonic acid (MES) hydrate, 2,2'-Bipyridyl, ascorbic acid,  
91 trichloroacetic acid, DL-dithiothreitol, Na<sub>2</sub>HPO<sub>4</sub>, NaH<sub>2</sub>PO<sub>4</sub>·H<sub>2</sub>O, N-Ethylmaleimide, folic acid and  
92 citric acid monohydrate were purchased from Sigma Aldrich (Steinheim, Germany), and acetonitrile,  
93 methanol, *ortho*-phosphoric acid (85%), iron (III) chloride hexahydrate and ascorbic acid from VWR  
94 (Darmstadt, Germany). Applichem GmbH (Darmstadt, Germany) provided 2,3-Dihydroxybutan-1,4-  
95 dithiol; and [<sup>13</sup>C<sub>5</sub>]-5-methyltetrahydrofolate, [<sup>13</sup>C<sub>5</sub>]-folic acid, disodium hydrogen phosphate,

96 potassium dihydrogen phosphate, sodium acetate trihydrate and acetic acid were supplied by Merck  
97 (Darmstadt, Germany). Sodium hydroxide and sodium chloride were obtained from J.T. Baker  
98 (Deventer, the Netherlands) and ethanol from Fisher Scientific (Fair Lawn, NJ, USA). 5-  
99 Methyltetrahydrofolate was purchased from Schircks (Jona, Switzerland).

## 100 **2.2 Fortification and thermal treatment**

101 The formate buffer was attained by diluting formic acid (10 mL/L) with distilled water and adding  
102 sodium hydroxide to the solution until a pH of 3.5. Apple purée (brand: ALNATURA) and carrot  
103 purée (brand: HIPPI) without additives were purchased at a supermarket in Freising, Germany. In order  
104 to attain a concentration of 4.0  $\mu\text{mol/kg}$  of 5-methyltetrahydrofolate and folic acid, the dilution steps  
105 were carried out as follows: The 5-methyltetrahydrofolate and folic acid were pre-dissolved in a  
106 phosphate buffer (pH 7.0) and diluted with distilled water to a volume of 100 mL; Subsequently, 10  
107 mL of this solution were added to 40 g of apple purée and carrot purée and thoroughly vortexed.  
108 Homogeneity was tested by comparing the recovery amount in two aliquots of two independently  
109 prepared fortification mixtures. Amber glass vials (4 mL) were filled with 2 mL of a formate buffer  
110 solution containing 4.0  $\mu\text{mol/kg}$  of 5-methyltetrahydrofolate, or up to the corresponding height in the  
111 case of fortified apple and carrot purées, and closed with screw caps. Thermal treatments were carried  
112 out in a stirred silicon oil bath. Temperatures at the bottom and in the middle were verified and  
113 showed a homogenous temperature distribution over time. After taking the tubes out of the silicon oil  
114 bath, the tubes were immediately deep-frozen ( $\leq -18\text{ }^\circ\text{C}$ ). Before analysis, glass vials were put in a  
115 water bath for rapid thawing. The first point of every time curve was omitted from the heat treatment.

## 116 **2.3 Folate analysis**

117 The stable isotope dilution assay of Ringling and Rychlik (2013) was employed with the modification  
118 that only the vitamer 5-methyltetrahydrofolate was analysed and  $^{13}\text{C}$ - instead of  $^2\text{H}$ -labelled standards  
119 were taken for quantification. Shortly after an equilibration of the isotopic marked internal standards  
120 with the sample, the extraction of folates followed. Before analysis by HPLC-MS/MS, the samples  
121 were purified.

122 Instrumental conditions were identical. A HyperClone Reversed Phase Column C18 (Phenomenex,  
123 Aschaffenburg, Germany) was used for separations. Values were first calculated as calcium salt. The  
124 percental aberration of a fixed standard concentration of 200  $\mu\text{g}/100\text{ g}$  was then calculated and was on  
125 average  $7 \pm 7\%$ . Each point of a time curve was subsequently corrected by the percental aberration of  
126 the first value to the standard concentration, to avoid parallelism of kinetics. The values were then  
127 transferred to their actual 5-methyltetrahydrofolate content by taking into consideration the calcium  
128 proportion, and converted to their molar concentration by respecting a molar mass of 459 g/mol.

#### 129 **2.4 Ascorbic acid analysis**

130 Ascorbic acid quantification was carried out by means of the colorimetric method of Stevens, Buret,  
131 Garchery, Carretero, and Causse (2006). Modifications were the following: approximately 500 mg of  
132 the food sample were taken for extraction, and absorption was measured at 525 nm on a  
133 spectrophotometer (Safas Xenius, Monaco).

### 134 **3. Results and discussion**

#### 135 **3.1 Natural folate concentration, extraction yield and test of homogeneity**

136 Apples and carrots are food matrices that contain inherently low amounts of folates. Raw apples do  
137 not contain any measurable folates amounts and raw carrots only contain 19  $\mu\text{g}/100\text{ g}$  of folates  
138 (USDA, 2016). The amount of 5-methyltetrahydrofolate monoglutamate in the purchased apple purée  
139 was under the limit of quantification for the analytical method, which is 0.22  $\mu\text{g}/100\text{ g}$  (Ringling &  
140 Rychlik, 2013). The concentration of 5-methyltetrahydrofolate in carrot purée was  $2.5 \pm 0.1\ \mu\text{g}/100\text{ g}$   
141 which was in the range listed on the USDA platform for raw carrot (19  $\mu\text{g}/100\text{ g}$ , USDA, 2018).  
142 Natural amounts of 5-methyltetrahydrofolate in both matrices were thus negligible compared to the  
143 target fortification level of 185  $\mu\text{g}/100\text{ g}$  corresponding to 4.0  $\mu\text{mol}/\text{kg}$ . Folic acid is a synthetic, not-  
144 naturally-occurring vitamer, and is thus not present in apple and carrots.

145 In order to verify the homogeneity of fortification preparations, the apple purée was supplemented  
146 with folic acid. Recovery rates accounted to  $107 \pm 5 \%$  and  $109 \pm 1 \%$ , which were determined by  
147 analysing two aliquots of two independently prepared fortification mixtures. Extraction was thus  
148 quantitative and homogeneity was acceptable, indicated by low standard deviations.

### 149 **3.2 Stability in buffer solution and apple purée**

150 The stability of the folic acid and 5-methyltetrahydrofolate was first evaluated in a formate buffer (10  
151 mL/L, pH 3.5) heated at 80 °C. The daily folate amount recommended by the European Food Safety  
152 Authority (EFSA) depends on the origin of the folate, as natural folates are less bioavailable than folic  
153 acid (EFSA, 2014). A dietary folate equivalent (DFE) is usually applied to convert folic acid amounts  
154 to the concentration that would be ingested if natural folates were absorbed. One  $\mu\text{g}$  of DFE is defined  
155 as 1  $\mu\text{g}$  food folate, which in turn corresponds to 0.6  $\mu\text{g}$  of folic acid from fortified food. This  
156 conversion factor was considered when preparations were carried out and it results in differing initial  
157 values depending on the vitamer used (Figure 2). Consistent with literature results (Delchier et al.,  
158 2016; O'Broin, Temperley, Brown, & Scott, 1975; Paine-Wilson & Chen, 1979), folic acid was more  
159 stable than 5-methyltetrahydrofolate (Figure 2A). Folic acid dissolved in a formate buffer was  
160 completely stable at 80 °C for the duration of 180 minutes while in turn 5-methyltetrahydrofolate  
161 degraded entirely within 15 minutes. The stability of folates depends on ions (Indrawati et al., 2004;  
162 Paine-Wilson & Chen, 1979); hence direct comparison of degradation rates with literature results is  
163 not possible as other buffer solutions were used. However, the degradation of 5-methyltetrahydrofolate  
164 occurred within the same time range as previously reported, registering a complete loss within some  
165 minutes (O'Broin et al., 1975; Paine-Wilson & Chen, 1979).

166 According to literature, the stability of 5-methyltetrahydrofolate for 15 minutes up to 170 °C can be  
167 obtained by adding ascorbic acid (Oey et al., 2006). In the present study, the addition of 2840  $\mu\text{mol/kg}$   
168 of ascorbic acid induced complete stability of 5-methyltetrahydrofolate in the formate buffer for three  
169 hours at 80 °C (Figure 2A). The apple purée contains fructose, which is known to accelerate the  
170 degradation of folates (Verlinde et al., 2010), but also polyphenols that may potentially exhibit a  
171 protective effect as a result of their anti-oxidative properties. Therefore, the stability of folic acid and



172 5-methyltetrahydrofolate was also assessed in apple purée (Figure 2B). As in the formate buffer, folic  
173 acid was completely stable over the investigated time range and 5-methyltetrahydrofolate degraded  
174 rapidly (Figure 2B). Mnkeni and Beveridge (1983) found that 5-methyltetrahydrofolate is also fairly  
175 unstable in apple juice. When 2840  $\mu\text{mol/kg}$  of ascorbic acid were added, the complete stability of 5-  
176 methyltetrahydrofolate was also attained in the apple purée. Naturally present polyphenols in the apple  
177 purée did not lead to the same stability as the addition of 2840  $\mu\text{mol/kg}$  of ascorbic acid. Ascorbic acid  
178 completely protected the 5-methyltetrahydrofolate in the apple purée even in the presence of a high  
179 amount of fructose. This is coherent with the result of Verlinde et al. (2010), who studied the stability  
180 of 5-methyltetrahydrofolate (0.04 mmol/L) in the presence of fructose (0.04 mmol/L) and ascorbic  
181 acid (1.13 mmol/L) in a model solution which was heated for 45 minutes at 100 °C.

182 Thus, 5-methyltetrahydrofolate, in the presence of 2840  $\mu\text{mol/kg}$  of ascorbic acid is equally as suitable  
183 for fortification as folic acid. Both vitamers withstood long periods of heating that can be encountered  
184 when food is reheated and kept warm for several hours. Until now, flour products have been  
185 exclusively chosen for mandatory folic acid fortification in the United States. However, folic acid, a  
186 fairly stable folate vitamer, is lost during the bread baking process (Gujska & Majewska, 2005).  
187 Neither folic acid nor 5-methyltetrahydrofolate in the presence of 2840  $\mu\text{mol/kg}$  of ascorbic acid were  
188 lost in the apple purée under the reported conditions. Thus, this study suggests that with regard to  
189 stability, apple purée exhibits an advantageous fortification matrix compared to flour that is baked into  
190 bread.

### 191 **3.3 Concentration effect of ascorbic acid on the stability of 5-methyltetrahydrofolate** 192 **tested in apple and carrot purées**

193 The amount of ascorbic acid (2840  $\mu\text{mol/kg}$ ) used for the assessment above was added in excess when  
194 compared to the content of 5-methyltetrahydrofolate (4.0  $\mu\text{mol/kg}$ ). It was also much higher than the  
195 concentration of dissolved oxygen, which was not measured but is of 258  $\mu\text{mol/kg}$  in water at 25 °C  
196 and is probably even lower under the studied conditions, given the matrices and temperatures used  
197 (Penicaud, Peyron, Gontard, & Guillard, 2012). In the next phase, lower amounts of ascorbic acid  
198 were added to the apple purée so as to verify if stability was still maintained. However, the addition of

199 neither 570  $\mu\text{mol/kg}$  nor 1420  $\mu\text{mol/kg}$  led to the absolute stability of 5-methyltetrahydrofolate within  
200 the investigated time range (Figure 3A). When adding 570  $\mu\text{mol/kg}$  of ascorbic acid, degradation  
201 started within 60 minutes and amounted to  $30 \pm 1\%$  after 180 minutes and in the case of 1420  
202  $\mu\text{mol/kg}$ , it initiated within 120 minutes and reached  $17 \pm 3\%$  after 180 minutes. However, once  
203 degradation started, the depletion pace for both ascorbic acid concentrations was still slower than  
204 when no vitamin C was added. In order to examine whether the loss of complete preservation of 5-  
205 methyltetrahydrofolate was linked to a loss of ascorbic acid, the concentration of the latter was  
206 followed over the time period analysed (Figure 3B). However, the amounts of ascorbic acid still  
207 present when the degradation of 5-methyltetrahydrofolate started were quite high.

208 Oey et al. (2006) reported that even low amounts of ascorbic acid (284  $\mu\text{mol/L}$ ) are sufficient to  
209 completely protect 5-methyltetrahydrofolate (4  $\mu\text{mol/L}$ ) at 80 °C. From these results, it can be  
210 concluded that besides the amount of ascorbic acid present in the medium, another time-dependent  
211 factor seems to crucially influence the stability of 5-methyltetrahydrofolate. It might be that dissolved  
212 oxygen is involved in this phenomenon. Dissolved oxygen was consumed within one hour in apple  
213 purée heated at 80 °C under the conditions of Herbig, Maingonnat, and Renard (2017), but this was  
214 not tested in the present work due to the difficult access of the oxygen sensor to the medium in the  
215 experimental set-up. The degradation of 5-methyltetrahydrofolate does not proceed when the medium  
216 is deprived of oxygen before the heat treatment (Delchier et al., 2014). It may also be that the  
217 generation of reactive oxygen species might as well be implicated in this phenomenon. Boatright  
218 (2016) measured up to 32.5  $\mu\text{mol/L}$  of hydrogen peroxide during the oxidation of 100  $\mu\text{mol/L}$  of  
219 ascorbic acid. The formation of hydrogen peroxide and other reactive oxygen species during the  
220 oxidation of ascorbic acid and possibly also other components in the apple purée medium, might  
221 explain the time-limited protective effect of ascorbic acid. A longer protection with increasing  
222 amounts of ascorbic acid is consistent with the assumption of reactive oxygen species being involved,  
223 as the probability rises that reactive oxygen species encounter an ascorbic acid molecule before a 5-  
224 methyltetrahydrofolate one.

225 Another explanation might also be an implication of sugar derivatives. In model solutions, ascorbic  
226 acid protects 5-methyltetrahydrofolate in the presence of fructose (100 °C, 45 minutes) by reducing 5-  
227 methylidihydrofolate, the first oxidation product (Figure 1) (Verlinde et al., 2010). In an apple purée,  
228 the protective effect might, however, be counteracted by an accumulation of sugar degradation  
229 products in the course of time and consequently explain the shifted degradation initiation of 5-  
230 methyltetrahydrofolate. In addition, since some ascorbic acid degradation products, namely  
231 dehydroascorbic acid and 2,3-diketogulonic acid, are also dicarbonyls that are even more reactive  
232 Maillard precursors than sugars, they might participate in the shifted degradation beginning (Ortwerth  
233 & Olesen, 1988; Reihl, Lederer, & Schwack, 2004; Roig, Bello, Rivera, & Kennedy, 1999; Slight,  
234 Feather, & Ortwerth, 1990).

235 In future studies, it might therefore be interesting to study the combined effect of the amount of  
236 oxygen in the headspace, dissolved oxygen, the appearance of reactive oxygen species, sugar  
237 derivatives and ascorbic acid derivatives, in order to understand the interactions more profoundly.  
238 Until these interactions are understood in detail, degradation kinetics cannot be extrapolated to other  
239 experimental set-ups.

240 As the composition of the food matrix may potentially influence the stability of 5-  
241 methyltetrahydrofolate, the latter was subsequently supplemented to carrot purée so as to verify  
242 possible stability changes. As with the apple purée, when no ascorbic acid was added, the 5-  
243 methyltetrahydrofolate was very fragile and degraded rapidly. Its stability in carrot purée in the  
244 presence of 570 µmol/kg and 2840 µmol/kg of ascorbic acid was not different from that of the apple  
245 purée when the same amount of ascorbic acid was added (Figure 3C). Stability was attained with the  
246 presence of 2840 µmol/kg of ascorbic acid but not when only 570 µmol/kg were used for fortification.  
247 Coherently, Mnkeni and Beveridge (1983), who studied the stability of fortified folic acid in apple and  
248 tomato juices (100-140 °C), did not observe any difference between degradation paces in the two  
249 studied food media despite their different compositions. In accordance with results obtained for the  
250 apple purée and with those of Indrawati et al. (2004), who worked with carrot juice (500 mg/kg of

251 ascorbic acid) heated at 120 °C for 40 minutes, the presence of ascorbic acid in carrot purée was  
252 linked to a high stability of 5-methyltetrahydrofolate.

### 253 **3.4 Temperature effect (60-80 °C)**

254 Temperature diminishes the stability of 5-methyltetrahydrofolate in a model solution according to the  
255 Arrhenius equation (Indrawati et al., 2004; Nguyen et al., 2003; Oey et al., 2006). However, the extent  
256 depends on the pH used and on the buffer ions (Indrawati et al., 2004; Paine-Wilson & Chen, 1979). It  
257 was hypothesised that stability for 3 h could be reached in apple purée with only 570 µmol/kg of  
258 ascorbic acid by decreasing temperature. However, when the temperature was decreased from 80 °C to  
259 70 °C or to 60 °C, the stability did not increase at the same time (Figure 4). Apparently, the supply of  
260 energy was not the limiting factor for the degradation pace in this temperature range. Indrawati et al.  
261 (2004) studied the impact of temperature on the stability of 5-methyltetrahydrofolate in orange juice,  
262 kiwi purée, carrot juice and asparagus. In the mentioned study, orange juice and kiwi purée contained  
263 500 mg/kg of ascorbic acid i.e. 2840 µmol/kg, and carrot juice and asparagus contained 100 mg/kg of  
264 ascorbic acid, corresponding to 570 µmol/kg. Consistent with the results of this study, they observed  
265 no marked impact between the amount that degraded at 70 °C or at 80 °C in the different food  
266 matrices.

## 267 **4. Conclusion**

268 The heat susceptibility of 5-methyltetrahydrofolate was confirmed in apple and carrot purées.  
269 Polyphenols and carotenoids that could, due to their anti-oxidative properties, potentially enhance the  
270 stability of 5-methyltetrahydrofolate are far from being as effective as the addition of vitamin C.  
271 Stability can only be obtained in the presence of vitamin C. The vitamin C amount which is necessary  
272 to entirely protect 5-methyltetrahydrofolate depends on the intended duration of the heat treatment.  
273 The length of time of complete stability can be prolonged by increasing the vitamin C concentration.  
274 The molar vitamin C concentration is not directly related to the concentration of 5-  
275 methyltetrahydrofolate. Even if the amount of ascorbic acid is still in excess considering the amount of

276 5-methyltetrahydrofolate or dissolved oxygen, degradation starts in the course of time. A time-  
277 dependent factor seems to intervene. Decreasing temperature in the range of 60-80 °C does not have a  
278 marked impact on the stability of 5-methyltetrahydrofolate. Thus, by heating purées at 80 °C, the risk  
279 of microbial growth can be decreased while the same nutritional value is maintained as if heated at 60  
280 °C. Apple purée and carrot purée in the presence of an adequate amount of vitamin C are thus suitable  
281 food matrices for folate fortification.

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286

## 287 **Figure captions**

288 **Fig. 1. Degradation mechanism of 5-methyltetrahydrofolate in the presence sugars.**

289 (1: 5-methyl-5,6,7,8-tetrahydrofolate, 2: 5-methyl-7,8-dihydrofolate, 3: p-aminobenzoyl glutamic acid,  
290 4: 2-amino-8-methyl-4,9-dioxo-7-methyl-p-aminobenzoylglutamate-6,7,8,9-tetrahydro-4H-  
291 pyrazino(1,2-a)-s-triazine, 5 : unknown reaction products, 6 : N(2 $\alpha$ )-[1-(carboxyethyl)]-5-methyl-  
292 5,6,7,8-tetrahydrofolic acid.)

293 Postulated and adapted from the version from Verlinde et al. (2010).

294

295 **Fig. 2. Degradation kinetics at 80 °C in a formate buffer and apple purée.**

296 ▲ Folic acid vs. ○ 5-CH<sub>3</sub>-H<sub>4</sub>folate and ● 5-CH<sub>3</sub>-H<sub>4</sub>folate in the presence of ascorbic acid (initial  
297 concentration: 2840  $\mu$ mol/kg).

298 A: Formate buffer (10 mL/L, pH 3.5)

299 B: Apple purée

300

301 **Fig. 3. Impact of the concentration of ascorbic acid on the stability of 5-CH<sub>3</sub>-H<sub>4</sub>folate**  
302 **in apple and carrot purées heated at 80 °C.**

303 A: Stability of 5-CH<sub>3</sub>-H<sub>4</sub>folate in apple purée containing an initial ascorbic acid concentration  
304 of ▽ 570 µmol/kg, □ 1420 µmol/kg or • 2840 µmol/kg.

305 B: Ascorbic acid content in apple purée during heat treatment as a function of the initial concentration.  
306 Ascorbic acid initial concentration: ▲570 µmol/kg vs. ■ 1420 µmol/kg and ♦ 2840 µmol/kg.

307 C: Stability of 5-CH<sub>3</sub>-H<sub>4</sub>folate in carrot purée after an ascorbic acid addition of \*570 µmol/kg or ◦  
308 2840 µmol/kg.

309

310 **Fig. 4. Impact of temperature on the degradation of 5-CH<sub>3</sub>-H<sub>4</sub>folate.**

311 5-CH<sub>3</sub>-H<sub>4</sub>folate in apple purée containing 570 µmol/kg of ascorbic acid

312 during a heat treatment at ▽ 60 °C, □ 70 °C, and • 80 °C.

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389

Figure 1 :

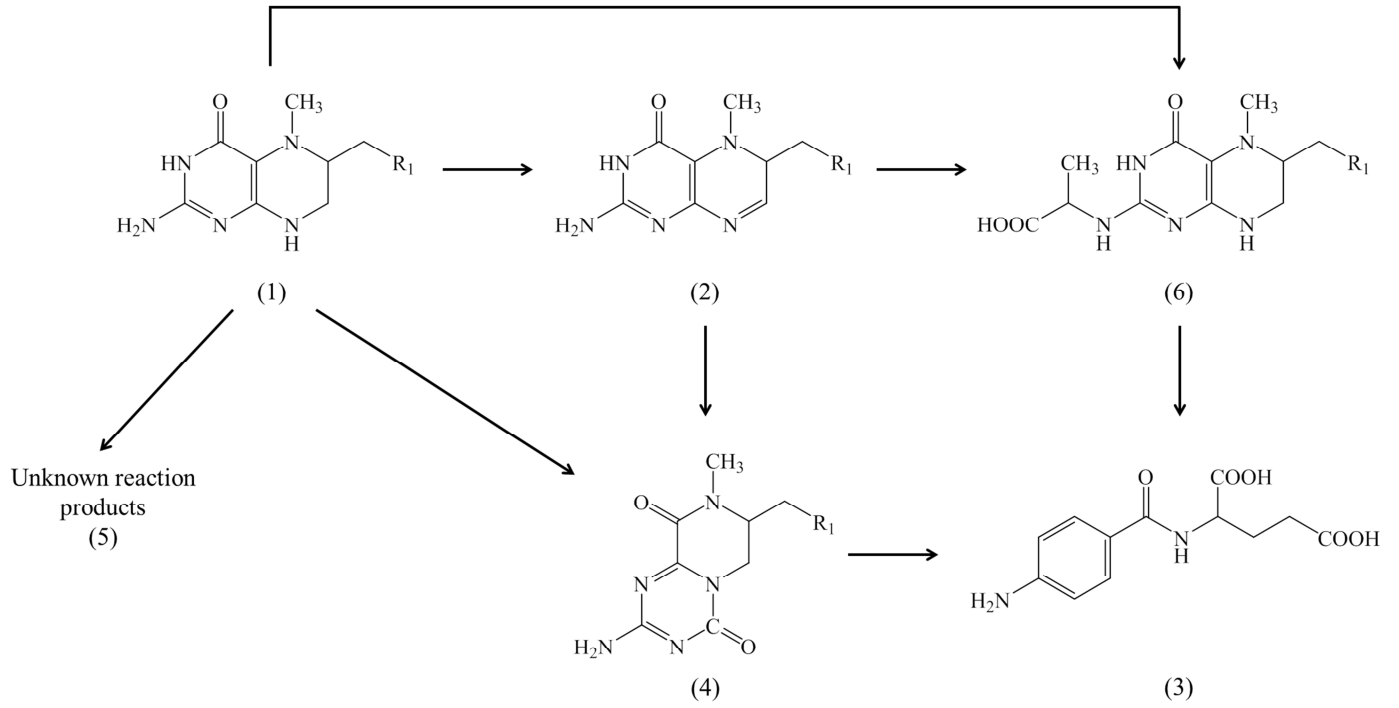




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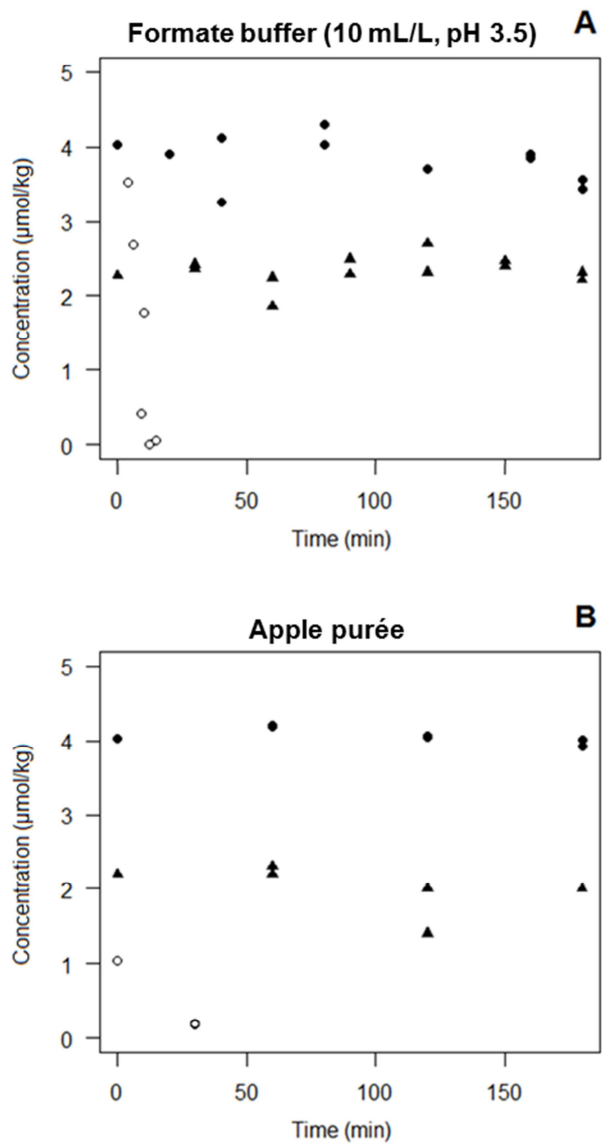


Figure 3:

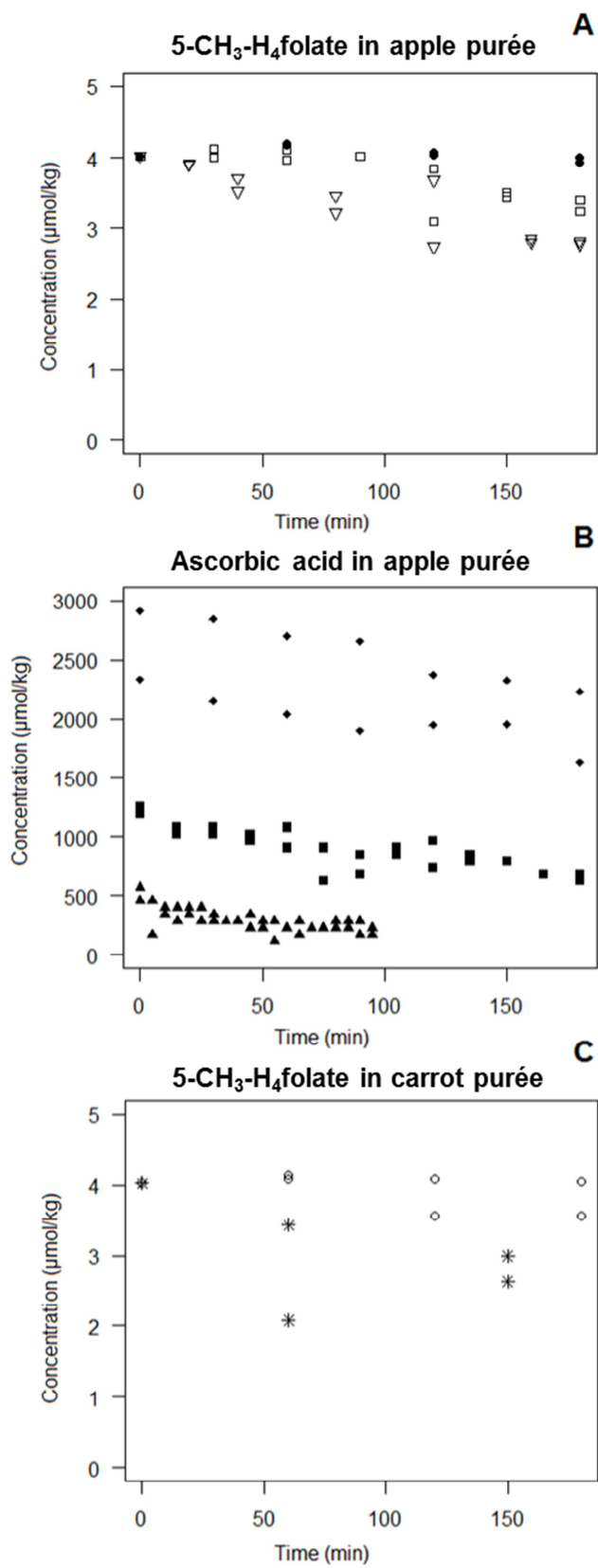


Figure 4:

