Chemical Evolution of Solutions from Beans Soaking and Cooking Processes: Case Study of Phaseolus vulgaris L

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Authors’ contributions

This work was carried out in collaboration between both authors. Author JCFT designed the study, performed the statistical analysis, managed the literature searches, wrote the protocol and wrote the first draft of the manuscript. Author INT managed the analyses of the study. Both authors read and approved the final manuscript.

Article Information

DOI: 10.9734/EJNFS/2021/v13i230373

ABSTRACT

The present paper aims to highlight the physicochemical evolution of solutions from soaking and beans cooking processes. For that purpose, solutions from soaking were produced by putting in contact 2kg of sorted and quickly washed beans seed with 8kg of water with known physicochemical characteristics. Concerning the solutions from cooked beans, they were produced by putting on fire the pot containing the water mixture from soaking and bean seeds. The beans were a variety of Phaseolus vulgaris L., known as “Meringue”. The cooking process was done without salts. The samples of solutions were collected as follows: 30 and 60 minutes respectively after the start of the soaking, 30, 60, and 90 minutes after the beginning of the cooking process of the beans soaked for 60 minutes. After each sampling, the cooking solution was brought back to the gauge line using the water prepared for the cooking process. Solutions obtained from beans soaking and cooking gradually enriched in mineral salts, particularly major macro-elements (N and
Keywords: waste water; sanitation; recycling; fertilization; agriculture; nutrition; temperature.

1. INTRODUCTION

From the onset of the nineteenth century, like swarms of bees, springing from fountain pens of thinkers, disturbing facts with increasing intensities aggregated. These are facts, as different as day and night and numerous like grains of sand on a beach, which reflect the occurrence of the multifaceted links that exist between the food consumed and the state of health of the consumer. In this sense, the better the quality and composition of the food, the better the health of the consumer, and vice versa [1]. The quality of food is commonly seen as a consequence of a more or less adequate state of conservation of the given material [2]. Regarding its composition, it is linked either to the nature of the food [3] or to the modification that the food has undergone during handling [4]. Therefore, the limit between the causes of malnutrition and good nutrition becomes a little more problematic to define because we are in the presence of a multifactorial scourge [5]. Contributing to better nourishment among people then comes down to carefully scrutinizing all the practices linked to this biological function. Talking about nutrition is about referring to the set of processes by which a living being transforms food to ensure its survival [6]. Regarding humans, the countless variety of food resources has logically led to highly varied diets [7]. Thus, while some people will adopt simple diets, others will adopt rather complex diets [8]. Regarding the type of food consumed by humans, we will distinguish some that are eaten raw and others that necessarily require the cooking phase [9]. From an essentially dietary perspective, the specialists in the field increasingly recommend the consuming of at least one fruit per day. It is indeed the undisputed source of vitamins for the body, the vast majority of these first-degree functional nutrients being heat-labile [10]. Such a prescription, carried out daily with insistence and authority, is very quickly found on the front line as generating many questions for informed man. One might, therefore be tempted to ask the question of whether the vitamin concentration in cooked foods is the only one to change during the cooking process. It is undoubtedly in this sense that [11] undertook to test, fortunately for scientific research, with success, the water at the end of cooking beans on strongly desaturated and very strongly acid soils as potential way towards soil fertility recovery. Observations from these trials subsequently led [12] to successfully test the mixture of end-of-cooking solutions from beans and human urine on plant growth. If, in the first case, there were significant improvements in the chemical fertility parameters of the soils involved, the characterization of the end-of-cooking solution of the beans remained only embryonic, thus helping to keep a heavy veil on the bottomless knowledge on this fluid. In the context of this study, it is, therefore, a question of returning to the shortcomings of the work of [11] and [12] to try to shed light on the real composition of solutions at the end of cooking and soaking the beans, pledge of a hard-wearing advance in Science in the management of what one might be tempted to call “kitchen waste.” Such investigation will be done about the chemical changes undergone by the food during the processes mentioned. It will detail a question of specifying and quantifying the elements lost when food products enter the cooking stage on the one hand and the soaking phase, on the other hand. By the principles of the inferential approach as recommended by [13], these losses will be assessed by the monitoring of the composition of the cooking water and the soaking, respectively. This work takes place in a context where to cook beans, many peoples think that because of the duration of the cooking process and the expenditure of the associated energy source, it would be preferable to soak the bean during at least five hours before. Moreover,
it has been observed that, after the soaking stage, most of the people discard the soaking water, wash the bean thoroughly, boil it, and then wash it again before entering the rest of the cooking process. This is therefore a behavior that an approach such as this should help to reconsider. The beans, because of their compulsory passage through the cooking step and optional through the soaking was retained. The choice of this plant species is linked to the fact that it is one of the foods not only abundantly produced but also appreciated in Cameroon [14]. From this large group, "Meringue," one of the central representatives of the Phaseolus vulgaris L. species, was chosen.

2. MATERIALS AND METHODS

2.1 Materials

1.1.1 Bean seeds

Most of the African population is engaged in agriculture [15-17]. Among the main cultivars, we have the bean [18]. This food is very popular throughout Cameroon because of its flavor and its dietary potential [19-21]. For 100g of this food, there are organic compounds such as proteins (9.06g), carbohydrates (27.91g), lipids (0.49g), fibers (5.3g); water (61.2g); varieties of vitamins including vitamins B1 or thiamine (0.257mg), B2 or riboflavin (0.063mg), B3 / PP or niacin (0.57mg), B5 or pantothenic acid (0.299mg), B6 (0.175mg), E or tocopherol (0.98mg), K (3.7μg); total folate (168μg); many mineral elements including potassium (508mg), sulfur (225mg), phosphorus (165mg), magnesium (65mg), calcium (52mg), iron (2.3mg), sodium (2mg), zinc (0.96mg), manganese (0.548mg), copper (0.271mg), selenium (1.4μg) [22-25]. The total nitrogen content (1450mg) was deduced from the protein content by applying the Jones factor [26] according to which protein content = total nitrogen * 6.25.

About 85% of dry beans are consumed in some of the countries where they are grown. The remaining 15% are marketed [27]. In the case of Cameroon, importation is almost absent. Large quantities are produced, but the bulk is for export [28].

The flatulent effect of beans is universally known and has, undoubtedly, been a source of discomfort throughout history [29].

2.1.2 Solution from cooked beans

The solution from cooked beans is a heterogeneous mixture, and particularly a proteic globular suspension, with considerable amounts of carbohydrates within. It has a pH value of 6.4. At rest, that fluid divides itself into two superimposed domains: a flaky superficial domain and a liquid lower domain. The flaky domain is the organic part and the liquid domain is the water and the mineral salts provider. The density of the flaky domain is 0.964 and that of the liquid domain is 1.011. The average speed of the growth of Aspergillus at the surface of the water from cooked beans is 3.17 cm²/H; they cover in five days a surface of 379.74 cm². The physic-chemical characteristics of the flaky domain floating on the liquid domain make it an adequate area for the development of molds (Aspergillus L.). Solution from cooked beans Seems then to contain all the nutrients required for an optimal development of those beings (Aspergillus L.), and in the same way for an optimal fertilization of soils; this include water, organic matters and mineral salts among which nitrogen, potassium, phosphorus, sulfur and calcium can be named. It is then a complete liquid organic fertilizer. That solution positions itself also as a high grade activator for soils micro flora. It represents finally a way for a sustainable improvement of agriculture in developing countries and a way for a sustainable development of soils micro flora, required for the reach of the food self-sufficiency. Solution from cooked beans is finally not only a source of nutrients for plants, but it also positions itself as a high grade activator for soils micro flora [13].

2.2 Methods

To achieve the fixed objectives, the work was carried out in the field and in the laboratory.

2.2.1 In the field

In the field, it was a question of obtaining the beans necessary for the production of the end-of-cooking solution for this food. The variety purchased is "Meringue" (photograph 1), one of the representatives of the species Phaseolus vulgaris L. It was also a question of getting closer to the families to question them in relation to the actions they take during the preparation of the bean-based meal. In total, between
September 2019 and December 2019, 500 families were interviewed, i.e. 50 families in each of the 10 regions of Cameroon.

2.2.2 In the laboratory

2.2.2.1 Handling of bean samples and preparation of water samples

In the laboratory, the bean was sorted in order to remove the stones. Then followed the weighing phase at the end of which 2,000 g of beans were taken for testing. The sample was then quickly washed to remove dust and all forms of impurities, then soaked for 1 hour in 8 liters of water, or 1 mass of bean for 4 mass of water. In detail, we started on the basis that 1 liter of water weighs 1 kg according to [30]. During the soaking phase, two samples of the same amount of water were taken. The first setting was made thirty minutes after the start of the soak and the second setting 60 minutes after; after each intake, the amounts of water removed were replaced by water drawn for cooking. Just before placing the pot containing the soaked water-bean mixture on the fire, a gauge line was marked on the rim of the pot in order to keep the water level intact during cooking by regular addition of water provided for cooking. Three water samples of the same quantities were taken during the cooking phase: the first 30 minutes after the start of cooking, the second 60 minutes after the start of cooking, and the third 90 minutes after the start of cooking. After each setting, the cooking water was brought back to the mark by adding water provided for cooking. The collected waters were cooled, bottled and the bottles labeled. These samples were supplemented by a sample of water used for cooking as control. Has followed the analytical phase. It focused on pH, electrical conductivity (EC), Ca²⁺, Mg²⁺, K⁺, Na⁺, HPO₄²⁻, SO₄²⁻, NH₄⁺, Cl⁻, and HCO₃⁻.

The IonPac CS12A and IonPac AS12A analytical columns were respectively used for the separation of ammonium (NH₄⁺), calcium (Ca²⁺), magnesium (Mg²⁺), sodium (Na⁺), and potassium (K⁺) as far as cations were concerned, and chloride (Cl⁻) and sulfate (SO₄²⁻) in the case of anions. The flow rate was set to 1.0 mL/min for both cations and anions optimization. The different cations were isocratically separated with a 20mM Methane sulfonic acid solution within 15 minutes. Concerning the ions studied here, isocratic elution with 4mM and 20mM Sodium hydroxide solution were employed for their separation, respectively for chloride and sulfate, during 15 minutes. All eluents were degassed and pressurized under high purity nitrogen to prevent dissolution of carbon dioxide and subsequent production of carbonate.

Concerning bicarbonate ion (HCO₃⁻), its amount was measured by using titrimetric method. The pH was determined with a pH-meter having a glass electrode. Electric conductivity for its own was measured using a conductivity meter fitted with a calibrated measuring cell; the results were expressed in mS/cm.

2.2.2.2 Determination of bean seeds mass loss

In order to determine the mass loss by bean seeds during soaking and cooking processes, 1kg of beans was taken. From there, six batches of 100g beans were made. The first batch served as a control (mto control mass = 100g). The other five batches were respectively washed quickly. The first batch of the five was soaked for 30 minutes, then dried (mht30cr = raw bean mass soaked for 30 minutes). The second batch was soaked for 60 minutes then dried (mht60cr = raw bean mass soaked for 60 minutes). The third batch was soaked for 60 minutes then cooked for 30 minutes and dried (mht60cu30 = bean mass soaked for 60 minutes and cooked for 30 minutes). The fourth batch was soaked for 60 minutes then cooked for 60 minutes and dried (mht60cu60 = bean mass soaked for 60 minutes and cooked for 60 minutes). The fifth batch was soaked for 60 minutes then cooked for 90 minutes and dried (mht60cu90 = bean mass soaked for 60 minutes and cooked for 90 minutes). In all five samples, drying was carried out in the open air until complete desiccation. To evaluate the mass losses (pm), we set respectively pm = mt - (mhtc)i. The percentage of material lost at each stage (pmpce) was obtained by setting pmpce = (pm / mt) * 100.

Photograph. 1. « Meringue »
3. RESULTS AND DISCUSSION

3.1 Results

3.1.1 Field facts

In Cameroon, beans are consumed in more than one form: boiled then fried, boiled then pounded, in the form of donuts, in the form of a cake, in the form of stew, boiled and then fried in mixture with boiled corn. Etc. By far, the most popular are the forms of cooking that integrate boiling in their cooking phases. The volume of water from cooking this cultivar is large, and increases with the amount of beans boiled.

To cook the beans, the populations, in general, sort, float, wash, soak, and boil. Regarding the trends in the use of end-of-cooking solution for beans (Fig. 1), 465 (93% of all) families (F1) throw end-of-cooking solution far from the habitat due to the subsequent odor induced. 15 (3% of all) families (F2) continue cooking with this water. 20 (4% of the whole) families (F3) cool this water and give it to their pigs to drink ...).

Pig farmers who feed their animals with spent grain mixed with bean end-cooking solution justify their actions by the extraordinary results obtained. The families who continue to cook with this solution justify this act by an inheritance from their parents, who have regularly claimed that in doing so, we are better nourished. They do it as part of the preparation of the culinary dish called "pounded". For this, they use the solution from the boiling of the beans to cook what they will pound the beans with. The families who throw out the bean after cooking water justify their act by the fact that it is dirt. Indeed, they are based on the fact that when one comes to pour the water at the end of cooking around the house, shortly after, one notices the emanation of a strong odor which can only disappear after two or even three days depending on the amount of bean solution poured. It shows that the majority of the population at the end of the cooking process rejects as waste water the water from the end of cooking of the bean (Fig. 1).

3.1.2 Analytical facts

All the parameters monitored within the framework of this study are on the rise despite local variations.

3.1.2.1 Cations

Among the cations, the decreasing order of concentration of the elements in the different solutions is as follows: $\text{NH}_4^+ >>> \text{K} >>> \text{Mg} \geq \text{Ca} >>> \text{Na}$.

Earth alkali cations

There is a strong similarity in the behavior of calcium and magnesium when the seeds are either soaked or boiled. In fact, the calcium contents (Fig. 2A and Table 1) are as follows: 0.0045g / l in the water used for cooking the seeds (E1), 0.046g / l after 30 minutes of soaking (E2), 0.103 after 60 minutes of soaking (E3), 0.183g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 0.321g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.403g / l after 60 minutes of soaking and 90 minutes of cooking (E6). Regarding magnesium (Fig. 2B and Table 1), its contents change as follows: 0.0034g / l in the water used for cooking the seeds (E1), 0.062g / l after 30 minutes of soaking (E2), 0.138 after 60 minutes of soaking (E3), 0.244g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 0.429g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.538g / l after 60 minutes of soaking and 90 minutes of cooking (E6). With the involvement of the temperature, the calcium and magnesium contents gradually and significantly increase during cooking to reach their maximum values at the end of the phenomenon. It can also be noted that between the thirtieth minutes and the sixtieth minutes of cooking, the respective content of these two elements tend to double before gradually increasing until the end of the cooking process.

Alkali cations

The potassium contents (Fig. 3A and Table 1) behave as follows: 0.0095g / l in the water used for cooking the seeds (E1), 1.255g / l after 30 minutes of soaking (E2), 1.294g / l after 60 minutes of soaking (E3), 2.358g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 2.519g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 2.68g / l after 60 minutes of soaking and 90 minutes of cooking (E6). Concerning sodium (Fig. 3B and Table 1), its contents change as follows: 0.0154g / l in the water used for cooking the seeds (E1), 0.008g / l after 30 minutes of soaking (E2), 0.019 after 60 minutes of soaking (E3), 0.034g / l after 60 minutes of soaking and 30 minutes of cooking.
(E4), 0.0596g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.07487g / l after 60 minutes of soaking and 90 minutes of cooking (E6). The intervention of temperature contributes to greatly increase the potassium and sodium contents in solutions even if the sodium concentrations remain low. Unlike potassium, the increase in sodium content occurs in conjunction with the duration of cooking. On the other hand, for potassium, the sudden increase is felt from the start of cooking; thereafter and until the end of the process, it increases rather slightly.

**Ammonium**

Ammonium is the most concentrated cation. Its contents (Fig. 4 and Table 1) change as follows: 0.009g / l in the water used for cooking the seeds (E1), 0.31g / l after 30 minutes of soaking (E2), 0.32g / l after 60 minutes of soaking (E3), 4.851g / l after 60 minutes of soaking and 90 minutes of cooking (E4), 5.183g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 5.41g / l after 60 minutes of soaking and 90 minutes of cooking (E6). We can then note that for this element, the intervention of the temperature causes a strong increase in its concentration in the solutions.

**3.1.2.2 Anions**

Among the anions, the decreasing order of concentration of the elements in the different solutions is as follows: $S > HCO_3 > P > Cl$.

The sulfate contents (Fig. 5A and Table 1) change as follows: 0.0085g / l in the water used for cooking the seeds (E1), 0.1236g / l after 30 minutes of soaking (E2), 0.1237g / l after 60 minutes of soaking (E3), 0.134g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 0.2633g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.29g / l after 60 minutes of soaking and 90 minutes of cooking (E6). Concerning bicarbonate (Figure 5B and Table 1), its contents change as follows: 0.0013g / l in the water used for cooking the seeds (E1), 0.0012g / l after 30 minutes of soaking (E2), 0.0011 after 60 minutes of soaking (E3), 0.0026g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 0.0182g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.095g / l after 60 minutes of soaking and 90 minutes of cooking (E6). These two most concentrated anions are also influenced by temperature. This physical magnitude allows them respectively to reach their respective highest content at the end of cooking. However, in the particular case of sulfates, its content increases suddenly and then remains relatively constant in the solutions from the start of soaking until the end of the thirtieth minute of cooking. At the end of the 60th minute of cooking, this content doubles before gradually increasing until the end of cooking. Regarding bicarbonates, their contents remain very low until the end of the thirtieth minute of cooking before being multiplied by 6 at the end of the sixteenth hour of cooking; at the end of the 90th hour of cooking, there is a more than sudden increase in these quantities, or they are multiplied by about 8.

Regarding the levels of phosphates (Fig. 6A and Table 1), they are 0.00146g / l in the water used for cooking the seeds (E1), 0.001826g / l after 30 minutes of soaking (E2), 0.00246g / l after 60 minutes of soaking (E3), 0.0298g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 0.0319g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.0339g / l after 60 minutes of soaking and 90 minutes of cooking (E6). For the chloride contents (Fig. 6B and Table 1), we have 0.0016g / l in the water used for cooking the seeds (E1), 0.0034g / l after 30 minutes of soaking (E2), 0.0098 after 60 minutes of soaking (E3), 0.00476g / l after 60 minutes of soaking and 30 minutes of cooking (E4), 0.0171g / l after 60 minutes of soaking and 60 minutes of cooking (E5), and 0.029g / l after 60 minutes soaking and 90 minutes of cooking (E6). Despite the low levels of these two anions, they respond favorably to seed contact with water, and especially with the involvement of temperature. In the particular case of phosphates, its contents first increase very slightly until the sixtieth minute of soaking. From the start of cooking, this content is multiplied by 10 at the end of the thirtieth minute before continuing to increase gradually until the end of cooking. For chlorides, despite an unexplained drop at the end of the thirtieth minute of cooking, the tendency is upward, with the end of cooking as the time of reaching their maximum concentration.

**3.1.2.3 EC and pH**

According to Fig. 7A and Table 1, the values of electrical conductivity remain low. However, they increase gradually from the end of the thirtieth minute of soaking until the end of the cooking where the value is maximum. In detail, we have in turn 0.029 in the water used for cooking the seeds (E1), 0.071 after 30 minutes of soaking (E2), 0.169 after 60 minutes of soaking (E3), 0.303 after 60 minutes of soaking and 30 minutes of cooking (E4), 0.532 after 60 minutes
of soaking and 60 minutes of cooking (E5), and 0.668 after 60 minutes of cooking and 90 minutes of cooking (E6).

As for the pH, its value increases by about one unit at the end of the thirtieth minute of soaking (from 5.1 to 6.02). From this point on, there is a slight but constant increase until the end of cooking. We then have respectively 6.02 in E2, 6.1 in E3, 6.3 in E4, 6.41 in E5, and 6.45 in E6 (Fig. 7B and Table 1).

From the above analysis, it appears that in the different solutions, ammonium followed by potassium are the two most concentrated main elements. They are followed from afar by magnesium, calcium, and sulfates, then very far by bicarbonates, sodium, phosphates and chlorides respectively (Fig. 8).

3.1.3 Correlation studies

The correlation is positive and perfect linear between magnesium and calcium; the cloud seems to regroup in three sets (the two red points for the first set, the three blue points for the second set, and the yellow point almost halfway for the third set); the correlation coefficient here is 1 (Figs. 9 and 63). That between calcium and potassium is just as positive with nonlinear positive monotonicity; the correlation coefficient is 0.8299 (Figs. 10 and 63). Dots seem to constitute three groups: the three red dots for the first group, the middle yellow dot for the second group, and the two blue dots for the third group.

The correlation between sodium and calcium is positive with 0.9813 as correlation coefficient. The major trend is linear. However, we note a distribution of the points in three groups: the two yellow points for the first group, the almost median red point for the second group, the three blue points for the third group (Figs. 11 and 63). The bond between bicarbonate and calcium appears to be positive, with a correlation coefficient of 0.7956. But, the grouping of points is very telling. Indeed, the first four points in blue, forming the first group, are arranged horizontally; red (second group) and yellow (third group) dots deviate from the first four respectively (Figs. 12 and 63).

Fig. 13 shows a positive nonlinear monotonic correlation between sulfate and calcium, with Pearson’s coefficient 0.9537 (Fig. 63). But in detail, we note the grouping of points in three sets: the yellow point for the first set, the three blue median points for the second set, and the two red points for the third set. Regarding Figure 14, it reveals equally a monotonic positive nonlinear correlation between phosphate and calcium, with Pearson’s index 0.9051 (Figure 63). The clustering of points is just as notable here. Thus, we have the first group represented by the three blue dots, the second group represented by the red dot, and the third group represented by the two upper yellow dots.

Fig. 1. Managing the bean end-of-cooking water in Cameroon
Table 1. Characterization of fluids during the process of soaking and beans cooking (mg/l)

|                                | Masse totale (g) | pH  | Ca2⁺   | Mg2⁺   | K⁺    | Na⁺   | CO₃²⁻ | SO₄²⁻ | HPO₄²⁻ | NH₄⁺ | Cl⁻ | E.C. (mS/cm) |
|--------------------------------|-----------------|-----|--------|--------|-------|-------|-------|-------|-------|------|-----|--------------|
| Eau simple (E1)                | 88.9            | 5.1 | 0.0045 | 0.0034 | 0.0095| 0.0154| 0.0013| 0.0085| 0.00146| 0.009| 0.0016| 0.0290       |
| Eau de trempage durant 30 minutes (E2) | 91.3            | 6.45| 0.046  | 0.062  | 1.255 | 0.098 | 0.0012| 0.1236| 0.00182| 0.310| 0.0034| 0.071        |
| Eau de trempage durant 60 minutes (E3) | 94.2            | 0.103| 0.138  | 1.294  | 0.019 | 0.0011| 0.1237| 0.00246| 0.320  | 0.0038| 0.169       |
| Eau de cuisson durant 30 minutes après 60 minutes de trempage (E4) | 98.3            | 6.46 | 0.183  | 0.244  | 2.358 | 0.034 | 0.0026| 0.134  | 0.0298 | 4.851 | 0.00476 | 0.303         |
| Eau de cuisson durant 60 minutes après 60 minutes de trempage (E5) | 105             | 6.30 | 0.321  | 0.429  | 2.519 | 0.0596| 0.0182| 0.2633 | 0.0319 | 5.183 | 0.0171 | 0.532         |
| Eau de cuisson durant 90 minutes après 60 minutes de trempage (E6) | 109.6           | 6.31 | 0.403  | 0.538  | 2.63  | 0.07487| 0.095 | 0.29   | 0.0333 | 5.41  | 0.029  | 0.668        |
Fig. 2A. Evolution of the amounts of Ca

Fig. 2. Evolution of the amounts of calcium and magnesium in response to the contact of bean seeds with water

Fig. 2B. Evolution of the amounts of Mg

Fig. 3A. Evolution of the amounts of K

Fig. 3. Evolution of the amounts of potassium and sodium in response to the contact of bean seeds with water

Fig. 3B. Evolution of the amounts of Na

Fig. 4. Evolution of the amounts of NH$_4^+$ in response to the contact of bean seeds with water
Fig. 5A. Evolution of the amounts of $\text{SO}_4^{2-}$

Fig. 5B. Evolution of the amounts of $\text{HCO}_3^-$

Fig. 5. Evolution of the amounts of sulfates and bicarbonates in response to the contact of bean seeds with water

Fig. 6A. Evolution of the amounts of $\text{HPO}_4^{2-}$

Fig. 6B. Evolution of the amounts of chlorides

Fig. 6. Evolution of the amounts of phosphates and chlorides in response to the contact of bean seeds with water

Fig. 7A. Evolution of EC values

Fig. 7B. Evolution of the pH values

Fig. 7. Evolution of the EC and pH in response to the contact of bean seeds with water
Fig. 8. Global distribution of ions in solutions from the contact of bean seeds with water

Fig. 9. Ca$^{2+}$ and Mg$^{2+}$ correlation

Fig. 10. Ca$^{2+}$ and K$^+$ correlation

Fig. 11. Ca$^{2+}$ and Na$^+$ correlation

Fig. 12. HCO$_3^-$ and Ca$^{2+}$ correlation

For Fig. 15, it reveals an equally monotonic positive nonlinear correlation between ammonium and calcium, with Pearson's coefficient 0.9061 (Fig. 63). The regrouping of the points here brings out a first group (the three blue points), a second group (the yellow point), and a third group (the two red points). Fig. 16 for its one shows a globally monotonic positive nonlinear binding between chlorides and calcium, with 0.923 as correlation coefficient (Fig.
The distribution of points around the middle line shows a grouping into five groups: the two blue points (group 1), the red point (group 2), the yellow point (group 3), the green point (group 4) and the purple point (group 5).

The correlation is almost linearly positive between electrical conductivity (EC) and calcium; the cloud seems to be grouped into three sets (the two blue points for the first set, the median red point for the second set, and the three yellow points for the third set); the correlation coefficient is 0.9994 (Figs 17 and 63). That between calcium and pH is non-linear monotonic positive, with the correlation coefficient 0.9081 (Figs 18 and 63); the points, located very high, seem to constitute three groups (the three blue points for the first group, the median red point for the second group, and the two yellow points for the third group). The steepest slope along the trendline is delimited between the first two blue points (from left to right); the rest of the points, almost perfectly aligned, generate a slight slope according to the trendline.

Fig.19 shows a globally monotonic positive nonlinear bond between potassium and magnesium, with the correlation coefficient 0.8942 (Fig. 63). The distribution of points around the middle line shows a grouping into four groups: the two brown points (group 1), the red point (group 2), the two blue points (group 3), and the yellow point (group 4). The correlation between sodium and magnesium is positive with the correlation coefficient 0.16601. The major trend is linear. However, we note a distribution of the points in three groups: the two blue points for the first group, the almost median yellow point for the second group, the three red points for the third group (Fig. 20 and 63).

The bond between bicarbonate and magnesium seems positive, with correlation coefficient 0.802. But, the grouping of points is very telling. Indeed, the first four points in blue, forming the first group, are arranged horizontally; the red (second group) and yellow (third group) dots respectively deviate from the first four (Figs. 21 and 63). The correlation between sulfate and magnesium is monotonic positive nonlinear, with the correlation coefficient 0.9515. We note a distribution of the points in three groups: the two yellow points for the first group, the three blue points for the second group, and the red point for the third group (Figs. 22 and 63).
Figs. 23 and 24 show a positive nonlinear monotonic correlation between phosphate and magnesium and between ammonium and magnesium, respectively. The correlation coefficient is 0.9044 in the first case and 0.9055 in the second case (Fig. 63). The regrouping of the points here brings out a first group (the three blue points), a second group (the yellow point), and a third group (the two red points).

The correlation is globally positive nonlinear monotonic between chlorides and magnesium (Fig. 25); their correlation coefficient is 0.9228 (Fig. 63). The distribution of points around the middle line shows a grouping into five sets: the two blue points (group 1), the red point (group 2), the yellow point (group 3), the green point (group 4), the brown point (group 5). The correlation is quite positive linear between electrical conductivity (EC) and magnesium; the cloud seems to be grouped into three sets (the three blue points for the first set, the median red point for the second set, and the two brown points for the third set); the correlation coefficient here is 0.9989 (Figs. 26 and 63).
The bond between magnesium and pH is monotonic positive nonlinear, with coefficient of correlation 0.9236 (Figs. 27 and 63); the points, located very high, seem to constitute three groups (the three red points for the first group, the median yellow point for the second group, and the two blue points for the third group). The steepest slope along the trendline is delimited between the first two red dots (from left to right); the rest of the points, almost perfectly aligned, generate a slight slope according to the trendline. Regarding the correlation between sodium and potassium, it seems positive with regard to the trend curve (Fig. 28), with the correlation coefficient 0.6175 (Fig. 63). The scatter plot here reveals four groups: the blue point for the first group, the two red points for the second group, the yellow point for the third group, and the two purple points for the fourth group.

According to the trend curve in Fig. 29, the correlation is positive between bicarbonate and potassium, with the correlation coefficient 0.8577 (Fig. 63). But, the distribution of points is ambiguous, suggesting four groups: the red point for group 1, the two blue points for group 2, the two yellow points for group three, and the purple point for group 4. Also, we see that the points of group 1 and group 2 are completely aligned horizontally. From the analysis of Fig. 30, there is also a positive correlation between sulfate and potassium, with Pearson's coefficient 0.8577 (Fig. 63). As before, the distribution of the points suggests four groups: the red point for the first group, the two blue points for the second group, the yellow point for the third group, and the two purple points for the fourth group. However, the points of the first and second group are well aligned along the trendline.

Figs 31 and 32 respectively highlight the positive correlation that exists between phosphate and potassium on the one hand, and between ammonium and potassium on the other. The correlation coefficients are 0.8995 and 0.9101 respectively (Fig. 63). In both cases, the cloud generated brings out three groups: the red point for the first group, the two blue points for the second group, the three yellow for the third group.

The correlation between chlorides and potassium is positive with regard to the trend curve (Fig. 33); the correlation coefficient is 0.7109 (Fig. 63). But, we note a strong dispersion between the points of the cloud, bringing out five groups in total: the red point for group 1, the two blue points for group 2, the yellow point for group 3, the green point for group four, and the purple point for group 5. Concerning Fig. 34, it shows for its part a positive correlation between electrical conductivity (EC) and potassium, with the correlation coefficient 0.8797 (Fig. 63). The dispersion of the dots brings out four groups: the red point for group 1, the two blue dots for group 2, the yellow point for group 3, and the two purple dots for group 4.

The correlation is positive between pH and potassium (Fig. 35), with Pearson's coefficient 0.9511 (Fig. 63). The cloud of points, located very high, is organized into three groups: the red point for group 1, the two blue points for group 2, and the three yellow points for group 3. Overall, despite the dispersion points, they are all very close to the trendline. The correlation is also positive between sodium and bicarbonate by referring to the trend curve (Fig. 36); their correlation coefficient is 0.1405 (Fig. 63). The cloud inducted individualizes four groups: the three blue points for the first group, the red point for the second group, the yellow point for the third group, and the purple point for the fourth group. But, the points of the first and second group line up horizontally along the abscissa axis.

Fig. 37 shows a positive correlation between sulfate and sodium, with the correlation coefficient 0.1503 (Fig. 63). The distribution of points individualizes three groups: the red point for the first, the three blue points for the second, the two yellow points for the third. Figure 38 for its part also shows a positive correlation between phosphate and sodium with regard to the trend curve. The correlation coefficient is 0.1515 (Fig. 63). The points of the cloud are individualized into three groups: the three blue points for the first, the red point for the second, and the two yellow points for the third.

The trend line in Fig. 39 shows a positive correlation between chlorides and sodium. The correlation coefficient is 0.1549 (Fig. 63). The dispersion of the points of the cloud individualizes four groups: the three blue points for the first, the red point for the second, the yellow point for the third, and the purple point for the fourth. Regarding Fig. 40, the distribution of points highlights a positive correlation between electrical conductivity (EC) and sodium, with the Pearson's index of 0.1672 (Fig. 63). The distribution of the points of the cloud reveals three groups: the three blue points for the first,
the red point for the second, and the two yellow points for the third. The vast majority of points are almost carried by the trendline.

The correlation is positive between pH and sodium (Fig. 41), with 0.1312 as the correlation coefficient (Fig. 63). Located high, the scatter plot for the most part clusters very close to the trendline. Also, this cloud individualizes three groups: the three blue points for the first, the red for the second, and the two yellow points for the third. The trend curve in Fig. 42 shows a positive correlation between sulfate and bicarbonate, with the correlation coefficient 0.7332 (Fig. 63). The distribution of points reveals four groups: the red point for the first, the two blue points for the second, the yellow point for the third, the purple point for the fourth. We note in detail that the fourth group is very eccentric with respect to the other three groups.
In Figs. 43 and 44, there is a positive non-linear monotonic correlation between phosphate and sulfate on the one hand and between ammonium and bicarbonate on the other hand. The correlation coefficients are respectively 0.5972 and 0.5949 (Fig. 63). The distribution of points shows four groups in the first case and three groups in the second case. In the first case we have: the two blue points for the first group, the red point for the second group, the two yellow points for the third group, and the purple point for the fourth group. In the second case, we have: the three blue points for the first group, the two yellow points for the second group, and the red point for the third group. In both cases, if the first groups tend to approach, the last group is completely eccentric.

Figs. 45 and 46 show a monotonic positive non-linear correlation between chlorides and bicarbonate on the one hand and between electrical conductivity (EC) and bicarbonate on the other. The correlation coefficients are respectively 0.9138 and 0.8088 (Fig. 63). The distribution of points highlights three groups in both cases: the three blue points for the first group, the two yellow points for the second group, and the red point for the third group. In both cases, if the first two groups tend to come closer, the last group is completely eccentric.

Regarding the correlation between chlorides and sulfates (Fig. 50), the correlation between electrical conductivity (EC) and sulfate (Fig. 51), and the correlation between pH and sulfate (Fig. 52) respectively, the trend curve shows that it is globally linear positive, with the respective correlation coefficient 0.8987, 0.3662, 0.8696 (Fig. 63). The point cloud of the pH-sulfate correlation has the particularity of being very high. Overall, this cloud makes it possible to visualize three groups of points, relatively distant from each other: the red point for the first group, the three blue points for the second group, and the two yellow points for the third group.

The correlation is positive nonlinear monotonous between the pH and the bicarbonate (Fig. 47), with the correlation coefficient 0.4522 (Fig. 63). The point cloud is high. Most of these points are concentrated towards the left end of the trendline. This cloud individualizes three groups: the four blue points for the first, the red point for the second group, and the yellow point for the third group. The correlation between phosphate and sulfate (Fig. 48) on the one hand and between ammonium and sulfate (Fig. 49) on the other hand is monotonic positive nonlinear, with correlation coefficient 0.8003 and 0.8060 respectively (Fig. 63). The point cloud in both cases is organized into four groups: the brown point for the first, the two blue points for the second, the yellow point for the third, the two red points for the fourth. We note in detail that these groups are highly distant from each other.

The trendline of the correlation between ammonium and phosphate (Fig. 53), pH and ammonium (Fig. 54), and pH and phosphate (Fig. 55) shows a positive relationship. The correlation coefficient is 0.7964 in the first case, 0.7395 in the second case, and 0.7275 in the third case (Fig. 63). The point cloud initiated by the correlation pH-NH₄⁺ and pH-HPO₄²⁻ is positioned higher (Figs. 54 and 55). But, in all three cases, this cloud individualizes two main groups, distant from each other: the three blue dots for the first group and the three red dots for the second group.

Regarding the correlation between electrical conductivity (EC) on the one hand (Fig. 56) and the correlation between electrical conductivity
and phosphate on the other hand (Fig. 57), the trend line shows a positive correlation. The respective correlation coefficients are 0.9065 and 0.9064 (Fig. 63). The cloud of points individualizes three groups: the three blue points for the first group, the red point for the second group, and the two yellow points for the third group.

In the context of the correlation between chlorides and ammonium (Fig. 58) on the one hand and between phosphates and chlorides (Fig. 59) on the other hand, the trend curve shows a positive relationship. The correlation coefficients are 0.6826 and 0.6823 respectively (Fig. 63). The scatter plot shows four groups: the three blue points for the first group, the red point for the second group, the purple point for the third group, and the yellow point for the fourth group.

The trend curve for the correlation between pH and chlorides (Fig. 60) on the one hand and between pH and electrical conductivity (EC) on the other hand (Fig. 61) shows a positive relationship in both cases, with for respective correlation coefficient 0.6456 and 0.7576 (Fig. 63). The distribution of points here brings out three groups for each of these cases. Regarding the correlation between pH and chlorides, the groups are: the four blue points for the first group, the red point for the second group, and the yellow point for the third group. Regarding the second case, the groups are: the three blue dots for the first group, the red dot for the second group, and the two yellow dots for the third group. In both cases, the third group is distant from the first two groups.

![Fig. 31. HPO$_4^{2-}$ and K$^+$ correlation](image1)

![Fig. 32. NH$_4^+$ and K$^+$ correlation](image2)

![Fig. 33. Cl$^-$ and K$^+$ correlation](image3)

![Fig. 34. EC and K$^+$ correlation](image4)
3.1.4 Mass variations of the beans during soaking and cooking processes

During soaking and cooking the beans, one can denote the mass loss at the different stages by bean seeds (Figure 64 and Table 2). However, the mass loss is lower during the soaking of the beans compared to the situation observed during the cooking process. In detail, from 100g at the beginning (mt), he mass reaches 97.6g after 30 minutes of soaking (mht30cr), 94.7g after 60 minutes of soaking (mht60), 90.6g after 60 minutes of soaking and 30 minutes of cooking (mht60cu30), 83.9g after 60 minutes of soaking and 60 minutes of cooking (mht60cu60), and 79.3g after 60 minutes of soaking and 90 minutes of cooking (mht60cu90). Then, compared to the initial situation (mt), the loss is
as follow: 2.4g, 5.3g, 9.4, 16.1g, 20.7g respectively in mht30cr, mht60cr, mht60cu30, mht60cu60, mht60cu90 stages (Fig. 63).

3.2 DISCUSSION

When the seeds are soaked, the amounts of the various target ions in the scope of this study increase significantly. During boiling, the stage after the soaking phase, this increase is greater. Overall, this increase is due to the hydrolysis of reserves contained in the cotyledons according to [31] and [32], the main seed storage organs as proved by [33]. The products resulting from this hydrolysis then pass through the teguments by diffusion to end up in the soaking or cooking solutions. [34] explains this integument crossing by relying on the permeable or semi-permeable character of biological membranes. This postulate corroborates the observations made by [35]. The sudden increase in the quantities of ions in solution during cooking is justified by the fact that the temperature, high or low, greatly influences the lysis of compounds [36]. The addition of the amounts of Ca$^{2+}$, Mg$^{2+}$, K$^+$, Na$^+$, SO$_4^{2-}$, HPO$_4^{2-}$, Cl$^-$, NH$_4^+$, Cl$^-$, and HCO$_3^-$ in the soaking or cooking solutions is in accordance with the composition of bean seeds as highlighted by [24] have shown this. In addition, [35] completes these observations by specifying the amounts of sulfur in bean seeds. NH$_4^+$ and HCO$_3^-$ come from the hydrolysis of organic compounds present in the cotyledons, in particular proteins for the first ion, and carbohydrates and lipids for the second ion. This is in agreement with the results of [36].

Likewise, the increase in pH and electrical conductivity (EC) is noted. The increase in pH is the consequence of the increase in the amounts of exchangeable basic cations in soaking or cooking solutions. This observation is commonly made in soil physiology by pedologists like [37,35], etc. Regarding electrical conductivity (EC), this is a quantity that is related to the presence of electrolytes in solution. Its behavior is consistent with the increase in ion quantities in the solutions followed here. However, according to [38], it is weak. This is the consequence of the low sodium contents in the solutions. The main ions contained in the solutions are NH$_4^+$ and K$^+$ respectively, followed by Mg$^{2+}$, Ca$^{2+}$, SO$_4^{2-}$, for the most abundant. This is in accordance with the order of decreasing concentration of bean seeds in these different elements. However, one can also relate the enrichment of solutions resulting from cooking bean seeds in Ca$^{2+}$, Mg$^{2+}$, K$^+$, and SO$_4^{2-}$ to the fact that these elements are highly influenced by temperature. In this sense, [39] and [40] shows through his two series of crystallization of minerals in a magmatic environment that the Ferromagnesians (forsterite type olivines) crystallize parallel to the calcium feldspars (anorthites) at high temperature. Lower, that is, at a lower temperature, crystallize potassium and sodium silicates, followed by other minerals. Moreover, it has been observed that during volcanic eruptions, many gases including sulfur dioxide are released [41]. This ease of sulfur to express in the presence of heat is consistent with the increase in sulfates in bean seed cooking solutions.

![Fig. 41. pH and Na+ correlation](image1)

![Fig. 42. SO$_4^{2-}$ and HCO$_3^-$ correlation](image2)
Further on, all the correlations are positive between the parameters followed in this present study. This is in agreement with the joint increase of these different parameters. One would therefore be tempted to think that the different parameters influence each other in solutions when bean seeds are soaked or boiled. However, a detailed observation shows that the points clouds organize themselves into groups, and that the points of several of these groups are arranged either horizontally as is the case in the correlation between sodium and bicarbonate, potassium and bicarbonate, bicarbonate and magnesium, and between bicarbonate and calcium, either vertically, as is the case in the correlation between electrical conductivity and bicarbonate, chlorides and bicarbonates, phosphates and sulfates, ammonium and bicarbonate, sulfate and bicarbonate, pH and bicarbonate, electrical conductivity and sulfate, and between electrical conductivity and ammonium. This demonstrates independence in the behavior of the parameters concerned according to the work of [42]. This is the reason...
why in a magmatic environment, some minerals will form first before others depending on the environmental conditions. Also, the points of the clouds organize themselves in groups which are sometimes very distant. This is the consequence of the presence of another underlying parameter which controls the behavior of the elements involved in the correlation. This is in agreement with the observations made by [42]. On a completely different level, we can see that in the organization of point clouds, many points are completely off-center, imposing a final trend that does not correspond to reality. This is the case in the correlation between pH and bicarbonate, electrical conductivity and bicarbonate, sulfate and bicarbonate, phosphate and sulfate, ammonium and bicarbonate, sulfate and bicarbonate, bicarbonate and potassium, bicarbonate and magnesium, and between bicarbonate and calcium. This corroborates the notable independence in the behavior of certain parameters taken in pairs according to [42]. In the same vein, the trendline of the correlations between some of these elements taken into account are linear positive as is the case for the correlation between calcium and magnesium, or even non-linear monotonic positive as it is the case for the correlation between pH and chlorides, pH and electrical conductivity, pH and phosphate, pH and ammonium, pH and bicarbonate, chlorides and bicarbonate, electrical conductivity and bicarbonate, ammonium and bicarbonate, phosphate and sulfate, and between magnesium and pH.[42] explains this fact by specifying that in this case, the value of one of the elements concerned in the correlation can, depending on the environmental conditions, strongly or weakly influence that of the other. This is in accordance with the temperature variation brought into play in the present study through the simple soaking and cooking of bean seeds channel. The correlations involving sodium have the lowest coefficients. This reflects a low degree of enrichment of sodium in solutions. This can be justified by the starting composition of bean seeds, which have low concentrations of this element, in particular about 2 mg per 100 g of bean seeds according to [24].

Such a cause and effect phenomenon has also been observed in the magmatic environment, where the composition of the final rock is closely related to that of the starting magma as shown by [40]. Still in the analysis of correlations, we see that between magnesium and calcium, the correlation is positive and perfect linear, with correlation coefficient 1. By drawing a parallel with the magmatic environment once again, we will notice that in [39]'s crystallization series, forsterite-type Ferromagnesians (Magnesian olivine) and anorthite-type calcium feldspars (calcium plagioclase) are the first minerals to form in the respective series, in particular the discontinuous series (black minerals) and continuous series (white minerals).

In the various solutions resulting from the contact between bean seeds and water, and especially in those whose production has undergone the intervention of temperature, the three main nutrients are, and in order of importance, nitrogen, potassium, and magnesium. According to [43], this are two of the three major macroelements (N and K), and one of the minor macroelements (Mg). After these three elements, we have the other minor macroelements (Ca and S) analyzed within the framework of the present study, and the trace elements here represented by the chlorides. Concerning the third major macroelement, in particular phosphorus, its contents are rather low. These contents in solutions, compared with those of calcium and magnesium, do not agree with the contents in seeds as showed by [24]; in fact, the phosphorus contents of bean seeds are at least twice the calcium and magnesium contents respectively. The phosphorus contents in the solutions could therefore have been at least comparable to those of calcium and magnesium. However, [44] clarifies the mystery by declaring at the end of his work that phosphorus is rather an element sensitive to the composition of the environment. In this sense, this author shows that in the presence of calcium among other things, the phosphate reacts with it to precipitate in the form of calcium phosphate as showed by [45], thus becoming not very concentrated in the solution. This is in agreement with the observations of [11]. Also, by referring to the correlations established between phosphorus and calcium, between phosphorus and magnesium, between phosphorus and potassium, and between phosphorus and sodium, we see in the cloud of points the formation of groups of points the arrangement of which is reminiscent of staircases, that is to say sequentially linear, thus revealing not only the influence that these elements can have on each other taken two by two, but also revealing the impact of temperature on their behavior according to [42]. Further on, we will establish such a theory by referring to the magmatic medium or in the same rocks, where we will find mineral associations highlighting the presence of magnesian (forsterite), calcium (anorthite), phosphate (apatite), sodium (albite)
as shown by [41]. Apart from the nutrients observed here, notably the major macroelements, minor macroelements, and trace elements, the bean end-of-cooking solution has a very high water content as showed by [46]. In addition, [13] showed that the bean end-of-cooking solution, at rest, settles, depositing organic matter. It is therefore a complete fertilizer. [13] has showed this quality by testing this solution as a substrate for heterotrophic beings, whose food needs are summed up in water, mineral salts and organic matter according to [47]. The parallelism drawn by [46] between this solution and human urine supports the observations made by [12].
Fig. 55. pH and $\text{HPO}_4^{2-}$ correlation

Fig. 56. EC and $\text{NH}_4^+$ correlation

Fig. 57. EC and $\text{HPO}_4^{2-}$ correlation

Fig. 58. $\text{Cl}^-$ and $\text{NH}_4^+$ correlation

Fig. 59. $\text{HPO}_4^{2-}$ and $\text{Cl}^-$ correlation

Fig. 60. pH and $\text{Cl}^-$ correlation

Fig. 61. pH and EC correlation
During the processes of soaking and cooking the beans, one can denote the differential mass loss at the different stages by bean seeds. This situation is due to the more or less extended duration of the contact between the bean seeds and the water. In that vein, [48] demonstrated the lysis capacity of water when it gets in contact with matters. According to [49], the products from the bean lysis traversed the seeds membrane and reach the solutions drowning the beans. This is consistent with the increase of the concentrations of the different ions followed up in the case of the present study. However, the mass loss is lower during the soaking of the beans compared to the situation observed during the cooking process. After [50], the involvement of temperature often increases the intensity of chemical reactions. This is in accordance with the amount of mass loss at different stages. In that point of view, compared to the initial situation, the loss of matter is 2.4g after 30 minutes of soaking. After 60 minutes of soaking, the loss is a little more than the double (5.3g) of the amount previously observed. After 60 minutes of soaking and 30 minutes of cooking the beans, the lost is quite the double (9.4g) of the former amount. After 60 minutes of soaking and 60 minutes of cooking the beans, the loss is equally quite the double (16.1g) of that observed before. After 60 minutes of cooking and 90 minutes of cooking the beans, the loss is about 1.3 time (20.7g) that of the earlier situation. The reduction of the increasing in losing the matter can be due to the progressive reach of the saturation point of the solutions as demonstrated by [51].
Table 2. Matrix of correlation

|       | Ca²⁺ | Mg²⁺ | K⁺   | Na⁺   | HCO₃⁻ | NH₄⁺ | SO₄²⁻ | HPO₄²⁻ | Cl⁻ | EC | pH |
|-------|------|------|------|-------|-------|------|-------|-------|-----|----|----|
| Ca²⁺  | 1    |      |      |       |       |      |       |       |     |    |    |
| Mg²⁺  | 1    | 1    |      |       |       |      |       |       |     |    |    |
| K⁺    | 0.8299 | 0.8942 | 1    |       |       |      |       |       |     |    |    |
| Na⁺   | 0.9813 | 0.16601 | 0.8031 | 1    |       |      |       |       |     |    |    |
| HCO₃⁻ | 0.7956 | 0.802 | 0.6175 | 0.1405 | 1    |      |       |       |     |    |    |
| NH₄⁺  | 0.9061 | 0.9055 | 0.9101 | 0.1509 | 0.5949 | 1    |       |       |     |    |    |
| SO₄²⁻ | 0.9537 | 0.9515 | 0.8577 | 0.1503 | 0.7332 | 0.8061 | 1    |       |     |    |    |
| HPO₄²⁻| 0.9051 | 0.9044 | 0.8995 | 0.1515 | 0.5972 | 0.7964 | 0.8003 | 1    |     |    |    |
| Cl⁻   | 0.923 | 0.9228 | 0.7109 | 0.1549 | 0.9138 | 0.6826 | 0.8987 | 0.6823 | 1  |    |    |
| EC    | 0.9994 | 0.9989 | 0.8797 | 0.1672 | 0.8088 | 0.9065 | 0.3662 | 0.9064 | 0.9235 | 1 |    |    |
| pH    | 0.9081 | 0.9236 | 0.9511 | 0.1312 | 0.4522 | 0.7395 | 0.8696 | 0.7275 | 0.6456 | 0.7576 | 1 |

Table 3. Mass variations of the beans during soaking and cooking processes

|                | mt   | mht30cr | mht60cr | mht60cu30 | mht60cu60 | mht60cu90 |
|----------------|------|---------|---------|-----------|-----------|-----------|
| M(g)           | 100  | 97.6    | 94.7    | 90.6      | 83.9      | 79.3      |
| Perte de masse (g) | 0   | 2.4     | 5.3     | 9.4       | 16.1      | 20.7      |
| Pourcentage de | 0   | 2.4     | 5.3     | 9.4       | 16.1      | 20.7      |
| matière perdue (%) |     |         |         |           |           |           |
5. CONCLUSION

Solutions obtained from beans soaking and beans cooking are gradually enriched in mineral salts, particularly major macro elements (N and K), minor macro elements (Ca, S, Mg), and oligoelements compared to the situation noticed in the water used for the cooking process. Concerning the third major macro element, notably the phosphorous, it is present in low amounts. The pH and the electric conductivity (EC) of the solutions increase too with the duration of the soaking and cooking processes. The increasing of the different amounts of the parameters studied here is shy and gradual during the soaking process and abrupt directly after the beginning of the cooking process. The correlations are globally positive. The study of the clouds of dots reveal the impact of the temperature as the major responsible of the behavior of some of those elements despite the positive correlations established. The Pearson index in the correlations including sodium are the lowest.

6. RECOMMENDATION

Recycling the bean end-of-cooking water can be done by reintroducing it in the rest of the bean cooking processes, either as inputs in pig feed, or moreover as fertilizer. This fluid must be gainfully stored in a cool place for gastronomic purposes, or directly in the ground if it is stored for agrarian target. The watering of soils with this solution must be directly followed by earthing up in other to avoid the loose of nitrogen and sulfur through gas emanation. It use does not require any dilution.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

1. Darche E, My diet, my best doctor! A few simple rules of food and daily hygiene for real health at any age, Embourg, Marco Pieter, 2007; 415.
2. De Heu H., Recipes and processes for the preservation of foods containing the best recognized preservation processes and more than 150 Standard Recipes, Jacobs, Brussels. 1917;46-47.
3. Pujol F. The 100 words of dietetics and nutrition, Presses Universitaires de France, 2010;74.
4. Dennekamp M, Howarth S, Dick CAJ. Ultrfine particles and nitrogen oxides generated by gas and electric cooking. Occupational and Environmental Medicine. 2001;58:511–516
5. Stratton RJ, Green CJ, Elia M. Scientific criteria for defining malnutrition. Disease related malnutrition: an evidence-based approach to treatment. Cambridge, UK: CABI Publishing. 2003;1–34
6. Tarek K, Abdel-Hamid. Modeling the dynamics of human energy regulation and its implications for obesity treatment, System Dynamics Review, Nº 4, Hiver 2002;18:431–471.
7. Marason, What Are the Components to the MIND Diet? Journal of the Academy of Nutrition and Dietetics, vol. 115, n°10, 2015;1744 (ISSN 2212-2672, PMID 26407649, DOI: 10.1016/j.jand.2015.08.002.
8. Béliveau D, Food intelligence: reprogramming the way you eat, Montreal, Les Éditions La Semaine. 2012;191.
9. Lagace J, Raw and cooked foods: Advantages and disadvantages; 2018. Available:https://maisonsaine.ca/sante-et-sante/sante/aliments-crus-et-cuits-avantages-et-inconvenients.html.
10. Tessier FJ. Effect of cooking on vitamins. Food cooking, Nutrition, and Health. Thematic file. Correspondance in Metabolisms, Hormones, Diabètes and Nutrition. 2012;XVI:5-6
11. Fopoussi Tuebue JC, Basga SD., Tematio P, Nguetnkam JP. Effect of trachyte powder, human urine and reserved water from cooked beans on andosols fertility in cameroonian western highlands. Asian Journal of Soil Science and Plants Nutrition. 2018;3(2):1-17.
12. Fopoussi Tuebue JC, Basga SD, Tematio P, Nguetnkam JP. Impact of the mixture of water from cooked bean and human urine on the growth of some common plants in Cameroon: case study of Talinum fruticosum and Ocimum gratissimum 2019a;5(1):1-15. Article no.AJSSPN.51071 ISSN: 2456-9682.
13. Fopoussi Tuebue JC, Basga SD, Tematio P, Nguetnkam JP. Water from cooked beans as substrate for some heterotrophic
organisms: Case Study of Moulds 2020a; 5(3):1-15. Article no: AJSSPN.53588 ISSN: 2456-9682.
14. Pasquet RS, Fotso M. Distribution of cowpea cultivars (Vigna Unguiculata (L.) Walp.) From Cameroon: Influence of the environment and human factors", Journald'AgricultureTraditionnelleet de Applied Botany. 1994;XXXVI(2):93-143.
15. CI. Genetic diversity in the genus Phaseolus, Study notebook, Intended to be used as a complement to the Audiotutorial Unit dealing with the same subject; Scientific content: Daniel D. Debouck; Production: Luz Maria Médina Call, Colombia, 1989:52.
16. Morin S. The fundamental dissymmetries of the West Highlands of Cameroon and their consequences on human occupation. Example of the Bambouto Mountains. The man and the tropical mountain. SEPLANIT ed., Bordeaux. 1988:49-51.
17. Shackleton CM, Pasquin MW, Drescher AW. African Indigenous Vegetables in Urban Agriculture. Earth scan Publishing for a sustainable future. London, Sterling, VA, 2009:339.
18. Aufrey P. Associated crops and cropping systems in Bamiléké country (West Cameroon). Center Univ. Dschang-CNEARC, 1st year ESAT thesis. 1985:82.
19. Lauchaux C., Dried beans with tomato; 2003. Available:http://pascal.moron.free.fr/rec
20. Andriniaina Harivolana P. Laboratory study of the behavior of some varieties of Phaseolus vulgaris L, IIS (50-1), IL5 (50-7), IL5 (56-1), IL5 (56-4), RJ2 and Lb with respect to Acanthoscides. Univ. Antananarivo, Ecole Norm. Sup., Dept of Initial Scientific Training, Sciences of Studies and Research Natural Sciences. Thes. 2006:99.
21. Marcia LP, Bykowski C, Fujiko FD, Danielle R. Reed. Excretion and perception of a characteristic odor in urine after asparagus ingestion: A psychophysical and genetic study, n° 1. Chemical Senses, 2011;36:9-17.
22. Boissaard P. Cuisinemalgache, cuisine créole. Librairie de Madagascar. 1997:144.
23. Press L. The bean book: Over Seventy Incredible Recipes. Globe Pequot Press; 2002, ISBN: 9781585744732.
24. Souci SW, Fachmann E, Kraut H. Food composition and nutrition tables. medpharm scientific publishers stuttgart; 2008.
25. Journal des Femmes. Haricot blanc ; 2019. Available:https://cuisine.journaldesfemmes .fr/encyclopédie-produits/1958658-haricots-blancs/
26. Jones DB Eds., Factors for converting percentages of nitrogen in foods and feeds into percentages of protein, US Department of Agriculture - circ. 1941:83.
27. Fanantenano J.O., Varietal improvement of white bean type white ingot by hybridization with a view to future export: 2002 - 2003 campaign in Nansana Memory of C.A.P.E.N. - ENS - University of Antananarivo. 2003:52.
28. Azefouet A, Tsague E, Medjou S, David O, Temple L, Parrot L, Quantification of cross-border flows in Cameroon. Hal Id: cirad-00948344. 2006:53.
29. Roger Fenwick R, Price KR, Johnson Ian T, Flatulence. Nutrition & Food Science. 1987;87(3):2-4. DOI: 10.1108/eb059428
30. Lide DR, Handbook of chemistry and physics, CRC, 89º éd. 2008:9-502736. ISBN: 142006679X et 978-1420066791),
31. Miege J, Reserves physiologiques-Reserves vegetales", encyclopedia Universalis; 2020. Available:http://www.universal.fr/encyclopedie/reserves-physiologiques-reserves-vegetales/
32. Grambow B, Michel N. Solubility of solids in water: surface property or solid? 6th ScientificDays of Maroucle, Subatech, Nantes. 2006 :53
33. Abderrazak Marof J.R. La botanique de A a Z, Dunod, 2007:69.
34. Rachidi W, Biological membranes: Structures and functions. health studies. Joseph Fourier University of Grenoble, First Year Medicine Course, UE2 Cellular Biology. 2012:75.
35. Fopoussi Tuebue JC. Petrological study of andosols in the West Highlands of Cameroon and fertilization test: Case of a toposequence developed on trachyte from the upper zone of the southern slope of the Bambouto Mountains (West Cameroon). Doctorate / PhD thesis, University of Ngaoundere, Faculty of Sciences, Department of earth sciences, laboratory of sciences, water and environment, 2018; 273.
36. Davis TR. The composition and origin of the gas produced during urological endoscopic resections. Br J Urol. 1983;55(3): 294-297

37. Ruellend A, Dosso M. Regard sur le sol. Université Francophone. Ed. Foucher Aupelf. 1995;192.

38. El Oumlouki K, Moussadek R, Zouahri A, Dakak H, Chati M, EL Amrani M. Study of the physico-chemical quality of water and soil in the Souss Massa region, (Case de perimeter Issen), Maroc / Study of physico-chemical quality of water and soil in the region Souss Massa (Case perimetrIssen), Morocco. J. Approx. Sci. 2014;5(S2):12-24. ISSN: 2028-2508. CODEN: JMESCN. 2365-2374

39. Bowen NL. The later stages of the evolution of the igneous rocks. Journal of Geology. 1915;23:1-89.

40. Bowen NL. The evolution of the igneous rocks. Princeton University Press; 1928.

41. Tchoua F, Contribution to the geological and petrographic study of some volcanoes of the Cameroon Line (Mont Manengouba and Mont Bambouto). Doctoral thesis State Sc., Univ. Clermont Ferrant. 1974;347.

42. Rakotomalala R., Correlation Analysis. Study of quantitative dependencies-variables. Version 1.1 UniversiteLumiere, Lyon 2, 2017.99.

43. Memento de l'agronome, French Republic, Ministry of Cooperation, reprint, 4th edition.

44. Chairat C. Experimental study of the kinetics and mechanisms of alteration of apatitic minerals. Application to the behavior of a ceramic containing minor actinides. Doctoral thesis, Univ. Toulouse, 2005;267.

45. Deferne J. At the heart of minerals, Museum of Geneva. 2014;45.

46. Fopoussi Tuebue Jc, Tchindal N, Djotsa D. Physico-chemical characterization of solutions from cooked beans and their comparison with human urine: fertilizer and food? European Journal of Nutrition & Food Safety. 2020c;12(12):9-27. Article no.EJNFS.63712 ISSN: 2347-5641

47. Djakou R, Thanon SY. Ecology. Intertropical Africa. 2nd A, C, 1st C and D. Bordas. 1986;167.

48. Duchaufour P. Pédologie: 1-pedogenesis and classification. Doc. 2nd Ed. Masson, Paris. 1983;491.

49. Rachidi W, Biological membranes: Structures and functions. Health studies. Joseph Fourrier University of Grenoble, First Year Medicine Course, UE2 Cell Biology. 2012;75.

50. David RL. CRC handbook of chemistry and physics, CRC Press, 90th Ed. 2009; 2804.

51. Lide DR. Handbook of chemistry and physics, CRC, 89 éd. 2008;9-50. ISBN: 142006679X et 978-1420066791).

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