



## ELABORATION OF SOME DECISION MODELS FOR THE NUTRI-SCORE LABEL

### 1 What is the Nutri-Score?

The Nutri-Score is a nutrition label that converts the nutritional value of products into a simple code consisting of 5 letters, each with its own colour. Each product is awarded a score based on a scientific algorithm. This formula takes into account the nutrients to avoid (energy value and the amount of sugars, saturated fats and salt) and the positive ones (the amount of fibre, protein, fruit, vegetables and nuts). **You can therefore see at a glance which products are recommended and which should be avoided**<sup>1</sup>.

In France, the Nutri-Score logo (see Figure 1) was elaborated by Santé publique France, a department of the Health Ministry, based on the scientific works of Professor Serge Hercberg (University Paris 13) and the experts of ANSES (Agence nationale de sécurité sanitaire de l'alimentation, de l'environnement et du travail), another department of this ministry.



Figure 1: Nutri-score logo

### This is how the Nutri-Score is calculated

The algorithm gives points for each element in the nutrition table (per 100 g or ml) - that means bad nutrients (energy, sugars, saturated fatty acids, salt) as well as good nutrients (proteins, fiber, percentage of fruit, vegetables & nuts). We then subtract the positive points from the negative ones and convert the result to the Nutri-Score table (see Figure 2).

Figure 3 below shows how to calculate the Nutri-score of an veggie food with the following information (detailed here <https://fic.colruytgroup.com/productinfo/fr/cogo/2493293> with no fruit/vegetable component):

- Energy (KJ): 485
- Sugars (g): 0.6
- Saturated fatty acids (g): 0.1
- Salt (g): 1.55

<sup>1</sup><https://nutriscore.colruytgroup.com/colruytgroup/en/about-nutri-score>

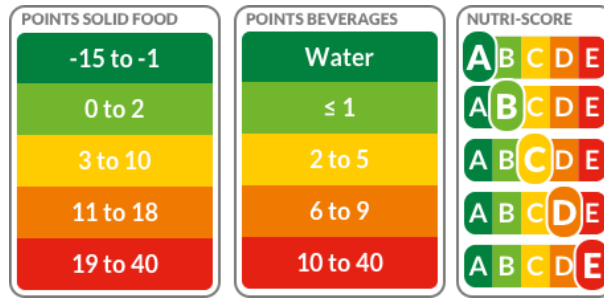


Figure 2: Assignment of foods to the classes

- Proteins (g): 22.4
- Fiber (g): 6
- Fruit/vegetable (%): 0

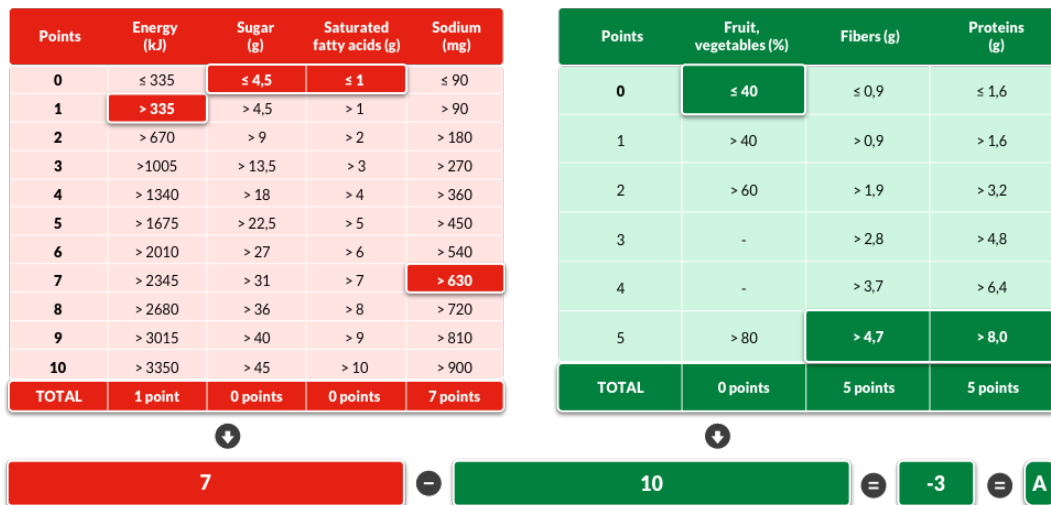


Figure 3: An example of the calculation of the Nutri-Score

## 2 The Nutri-Score viewed as a Mutlti-Criteria Decision Analysis problem

In this project, we will consider the calculation of the Nutri-Score of a food as a Multi-Criteria Decision Analysis problem where:

- The set of alternatives  $X$  corresponds to the foods analyzed.
- The seven criteria (the set  $N$ ) to take into account are:

1. *en*: **Energy** (KJ) (criterion to be minimized)
2. *su*: **Sugars** (g) (criterion to be minimized)
3. *fa*: **Saturated fatty acids** (g) (criterion to be minimized)
4. *sa*: **Salt** (g) (criterion to be minimized)
5. *pr*: **Proteins** (g) (criterion to be maximized)
6. *fi*: **Fiber** (g) (criterion to be maximized)
7. *fr*: **Fruit/vegetable** (%) (criterion to be maximized)

Let us consider a food  $x$ . We set  $x_i$  the number of points obtained by  $x$  on each component (criterion)  $i$ ,  $i \in \{en, su, fa, sa, pr, fi, fr\}$ , by using the real Nutri-score scale. Hence the food  $x$  can be denoted by:

$$x = (x_{en}, x_{su}, x_{fa}, x_{sa}, x_{pr}, x_{fi}, x_{fr}). \quad (1)$$

For each criterion  $i$ ,  $i \in \{en, su, fa, sa, pr, fi, fr\}$ , let us define the following new scale transformation:

1.  $x'_i = 10 - x_i$  for the criteria to be minimized  $i \in \{en, su, fa, sa\}$ ;
2.  $x'_i = 2 \times x_i$  for the criteria to be maximized  $i \in \{pr, fi, fr\}$ ;

This new scale leads to this another representation of the same food  $x$ :

$$x = (x'_{en}, x'_{su}, x'_{fa}, x'_{sa}, x'_{pr}, x'_{fi}, x'_{fr}). \quad (2)$$

From this new representation of  $x$  (see Equation 2), let us define the function  $F$  as follows:

$$F(x'_{en}, x'_{su}, x'_{fa}, x'_{sa}, x'_{pr}, x'_{fi}, x'_{fr}) = x'_{en} + x'_{su} + x'_{fa} + x'_{sa} + \frac{1}{2}(x'_{pr} + x'_{fi} + x'_{fr}) \quad (3)$$

If we denote by  $\text{Nutri-Score}(x)$  (respectively  $F(x)$ ) the score of the food  $x$  obtained by applying the real Nutri-score (respectively the function  $F$  of the Equation 3), then we have the following result:

$$\text{Nutri-Score}(x) = 40 - F(x) \quad (4)$$

Equation 4 means that the algorithm of the Nutri-score is equivalent to the weighted sum, implying that these 7 criteria satisfy the mutual independence preference property.

But there is a correlation between some criteria since the following Equation (of the computation of Energy) shows that the criterion “Energy” depends on “Fat”, “Sugars”, “Proteins” and “Fibers”:

$$\text{Energy} = (9 \times \text{fat}) + (7 \times \text{alcohol}) + (4 \times \text{protein}) + (4 \times \text{sugar}) + (2.4 \times \text{organic acids}) + (2.4 \times \text{polyols}) + (2 \times \text{fibers}) \quad (5)$$

### 3 Ecoscore

The Ecoscore is an experimental indicator that represents the environmental impact of food products. It classifies products from A (low) to E (high) according to their impact on the environment. Eco-Score aims to inform consumers on the environmental impact of the food they choose so as to guide them towards more responsible consumption. This indicator helps producers and consumers to decide and to be responsible.

The computation of the Ecoscore is based on the following principles:

- Public data : quantitative data on product Life Cycle Assessment (LCA) from the Agribalyse database <https://doc.agribalyse.fr/documentation-en>, drawn up by experts and implemented in Agribalyse. Impacts on environment through production, transport, fabrication, and packaging are taken into account, giving a score out of 100.
- Data not included in LCA but which do take into account the positive or negative impact on the environment: data on the product label or given by the producer, as well as additional quality criteria : recyclability of packages, labels (Bio, quality etc.), where the ingredients come from, seasonality of food used (for recipes and ready meals). All these data will give a bonus/malus which will influence the score.
- The total mark out of 100 gives a score from A to E.

See <https://docs.score-environnemental.com/en> for more details about the Ecoscore.

## 4 Elaboration of your own database of foods as the set of alternatives

1. Each group have to elaborate his own set of alternatives, in an Excel file, from one or some categories of foods (e.g. breakfast cereals, cookies, aperitif, industrial ready-made dishes based on seafood, industrial ready-made dishes based on vegetables, industrial ready-made dishes based on meat,...). This database will contain at least 200 foods.
2. The Ecoscore label (from A to E) and its global score, associated to each food, should be integrated in your database.
3. The set of alternatives to be evaluated will be the set of foods contained in this database.
4. The databases of any two different groups should not contain, in common, more than 30% of foods. Each group will implement a python function comparing two databases and allowing to satisfy this requirement.
5. The information about many foods is available by using the following this link: <https://fr-en.openfoodfacts.org/>. In this website, you can choose, for instance, the country of the products you want to evaluate. For instance, the information about foods of Spain are available in <https://es.openfoodfacts.org/>. The computation of the nutri-score label is given for all the foods.

## 5 Elaboration of a Nutri-Score model based on a simple sorting (ordered classification) model

Build the functions `PessimisticmajoritySorting` and `OptimisticmajoritySorting`, respectively based on the Pessimistic and Optimistic version of ELECTRE-Tri, which returns an Excel file containing the classified foods in the previous five Nutri-Score labels. The input will be a database contained in an Excel file.

- In your program, the limiting profiles could be set directly or manually (for instance, you could play yourself the role of the Decision Maker). You can also determine them by using an appropriate approach, like a data segmentation approach (e.g. a quintile approach), that you will detail.
- You will first test your program by using the following set of weights, given in a Table 1 below, associated to the seven criteria.
- Provide also, and justify, another set of weights and test also your programs with that set.
- You should use these three majority thresholds,  $\lambda = 0.5$  (or  $\lambda = 50\%$ ),  $\lambda = 0.6$  (or  $\lambda = 60\%$ ) and  $\lambda = 0.7$  ( $\lambda = 70\%$ ), in your tests. Of course, you can also test your programs with any other majority thresholds.

	1- Energy	2-Sugars	3-Satu. fat.	4- Salt	5- Proteins	6- Fiber	7-Fruits & vegetables
Weights $w_i$	4	3	3	3	2	2	1

Table 1: An example of the weights associated to the criteria

- Compute all the assignments of foods obtained from your database and compare your results with the real Nutri-Score assignments (by using, for instance, a confusion matrix). **Our objective is not to minimize the errors between these two classifications approaches.**
- In fact, the ordered classification models you computed above could be considered as your own Nutri-Score classification models. Provide an analysis of your results. For instance, the original Nutri-Score could be explained (or be approximated) by one of your classification models ?

## 6 Elaboration of two MCDA models combining nutri-score and Eco-score

Elaborate two decision models combining two criteria: the original nutri-score and the Eco-score. One of this model will be a weighted sum and the other one a sorting model or a model based on simple decision rules.

## 7 Elaboration of Nutri-Score model based on a machine learning classification model

Build a python function `AnotherMethodNutriScore` returning an assignment of each food to a predefined Nutri-score class, by using a machine learning algorithm (e.g. a decision tree, random forest, ...). Test and compare the obtained results with your previous results and the real Nutri-score assignments.

## 8 A comparison with another group

Apply one of your MCDA model to a database used by another group. Compare the obtained results with their results.

## 9 Minimal requirements

1. For this project, each group will be constituted by **two or three students**.
2. Each group will present (by using a power-point slides for instance), on December 9th, their preliminary results during 15 minutes.
3. Each group will write a report (document in .doc or .pdf) explaining and justifying their results, the parameters chosen, the interpretation of the obtained results, etc.
4. Which model you seem comfortable with (among all the models developed in this project)? Which model is suitable to calculate your own Nutri-Score of a food ? Among your models, which could be suitable to explain to a consumer the computation of a nutritional score ? Justify all your answers.
5. Send your report and source files by the **9th January 2025, 23h59 (Paris hour)**.

## A ELECTRE TRI methods

### A.1 Elaboration of the outranking relation $\mathcal{S}_\lambda$

Let  $A$  be a set of alternatives evaluated on  $n$  real-valued criteria  $g_i : A \rightarrow \mathbb{R}$ ,  $i \in N = \{1, \dots, n\}$ . We denote by  $g_i(a)$  the performance of the alternative  $a$  on criterion  $i$ . A nonnegative weight  $w_i$  is also assigned to each criterion  $i$  (w.l.o.g. we suppose  $\sum_{i=1}^n w_i = 1$ ).

We associate with each criterion  $i \in N$ , a nonnegative preference threshold  $p_i \geq 0$ . If the value  $g_i(a) - g_i(b)$  is positive but less than  $p_i$ , it is supposed that this difference is not significant, given the way  $g_i$  has been built. Hence, on this criterion, the two alternatives should be considered indifferent.

Using this information, we define on each criterion  $i \in N$  the partial concordance index  $c_i : A \times A \rightarrow [0, 1]$  as follows:

$$c_i(a, b) = \begin{cases} 1 & \text{if } g_i(b) - g_i(a) \leq p_i \\ 0 & \text{if } g_i(b) - g_i(a) > p_i \end{cases} \quad (6)$$

The valued relations  $c_i$  are aggregated to a single concordance index  $c : A \times A \rightarrow \mathbb{R}$  by using the following Equation:

$$c(a, b) = \sum_{i=1}^n w_i c_i(a, b) \quad (7)$$

The binary relation on  $A$  called outranking relation is defined by:

$$a \mathcal{S}_\lambda b \text{ iff } c(a, b) \geq \lambda \quad (8)$$

where  $\lambda \in [0, 1]$  is a cutting level (usually called a threshold and taken above  $\frac{1}{2}$ ).

**Interpretation:** An alternative  $a \in A$  outranks an alternative  $b \in A$  if it can be considered at “least as good” as the latter (i.e.,  $a$  is not worse than  $b$ ), given the values (performances) of  $a$  and  $b$  at the  $n$  criteria. If  $a$  is not worse than  $b$  in every criterion, then it is obvious that  $a \mathcal{S}_\lambda b$ . However, if there are some criteria where  $a$  is worse than  $b$ , then  $a$  may outrank  $b$  or not, depending on the relative importance of those criteria and the differences in the evaluations (small differences might be ignored).

From  $\mathcal{S}_\lambda$  we derive the following three binary relations:

☞ “Strictly better than” relation:

$$a \mathcal{P}_\lambda b \text{ iff } [a \mathcal{S}_\lambda b \text{ and not}(b \mathcal{S}_\lambda a)] \quad (9)$$

☞ “Indifferent to” relation:

$$a \mathcal{P}_\lambda b \text{ iff } [a \mathcal{S}_\lambda b \text{ and } (b \mathcal{S}_\lambda a)] \quad (10)$$

☞ “Incomparable to” relation:

$$a \mathcal{P}_\lambda b \text{ iff } [\text{not}(a \mathcal{S}_\lambda b) \text{ and not}(b \mathcal{S}_\lambda a)] \quad (11)$$

### A.2 ELECTRE TRI (also called ELECTRE TRI B)

Let us consider  $r$  ordered categories  $C^1, C^2, \dots, C^r$ ,  $C^1$  is the worst one and  $C^r$  is the best one. The category  $C^k$  is modeled by using limiting profiles. The lower limiting profile of  $C^k$  is  $\pi^k$ . The upper limiting profile of  $C^k$  is  $\pi^{k+1}$ . We suppose that the limiting profiles are such that  $\pi^{k+1}$  strictly dominates  $\pi^k$ <sup>2</sup>. The profile  $\pi^1$  (respectively  $\pi^{r+1}$ ) is taken low (respectively

<sup>2</sup>An alternative  $a$  dominates an alternative  $b$ , we note  $a \Delta b$  iff [for all  $i \in N$ ,  $g_i(a) - g_i(b) \geq 0$ ].  $a$  strictly dominates  $b$  if  $[a \Delta b \text{ and not}(b \Delta a)]$

high). It will be convenient to suppose that  $\pi^k \in A$ , for each  $k = 2, 3, \dots, r$ , while  $\pi^1, \pi^{r+1} \notin A$ . With this convention we have

$$\text{For all } a \in A, a \mathcal{P}_\lambda \pi^1 \text{ and } \pi^{r+1} \mathcal{P}_\lambda a. \quad (12)$$

ELECTRE TRI ([9], chap. 6) renamed ELECTRE TRI-B by Almeida-Dias et al. [4] is a MultiCriteria Decision Aid method using limiting profiles. It has two versions called “pessimistic” and “optimistic” in [9]. In [8] the name “pseudo-conjunctive” is used for the “pessimistic” version and “pseudo-disjunctive” for the “optimistic” version. These two versions are defined as follows:

**Définition 1** (Pessimistic version: ETRI-B-pc). *Decrease  $k$  from  $r + 1$  until the first value  $k$  such that  $a \mathcal{S}_\lambda \pi^k$ . Assign alternative  $a$  to  $C^k$ .*

ETRI-B-pc assigns an alternative  $a$  to the unique category  $C^k$  such that  $a$  is at least as good as to the lower limiting profile of this category and is not at least as good as its upper limiting profile (the relation “at least as good as” being  $\mathcal{S}_\lambda$ ).

**Définition 2** (Optimistic version: ETRI-B-pd). *Increase  $k$  from 1 until the first value  $k$  such that  $\pi^k \mathcal{P}_\lambda a$ . Assign alternative  $a$  to  $C^{k-1}$ .*

ETRI-B-pd assigns an alternative  $a$  to the category  $C^k$  such that the upper limiting profile of this category is better than  $a$  and the lower limiting profile of this category is not better than  $a$  (the relation “better than” being  $\mathcal{P}_\lambda$ ).

**Remarque 1.** Roy and Bouyssou ([9],chap.6,pp.393-395) have shown that if  $a \in A$  is assigned to the category  $C^k$  by the pessimistic version and to the category  $C^l$  by the Optimistic version, then  $k \leq l$ .

### A.3 Majority Rule sorting procedure (MR-Sort)

MR-Sort is a simplified version of the ELECTRE TRI sorting model directly inspired by the work of Bouyssou and Marchant [1, 2] who provide an axiomatic characterization of non-compensatory sorting methods. The general principle of MR-Sort (without veto) is to assign alternatives by comparing their performances to those of profiles delimiting the categories. An alternative is assigned to a category “above” a profile if and only if it is at least as good as the profile on a (weighted) majority of criteria.

The condition for an alternative  $a \in A$  to be assigned to a category  $C^k$  is expressed as follows:

$$\sum_{i: g_i(a) \geq g_i(\pi^{k-1})} w_i \geq \lambda \text{ and } \sum_{i: g_i(a) \geq g_i(\pi^k)} w_i < \lambda \quad (13)$$

The MR-Sort assignment rule described above involves  $r \times n + 1$  parameters, i.e.,  $n$  weights,  $(r - 1) \times n$  profiles evaluations and 1 majority threshold.

As demonstrated in [6], the problem of learning the parameters of a MR-Sort model on the basis of assignment examples can be formulated as a mixed integer linear program (MILP) but only instances of modest size can be solved in reasonable computing times. The MILP proposed in [6] contains  $m \times (2n + 1)$  binary variables, with  $n$ , the number of criteria, and  $m$ , the number of alternatives. A problem involving 1000 alternatives, 10 criteria and 5 categories requires 21000 binary variables. For a similar program in [3], it is mentioned that problems with less than 400 binary variables can be solved within 90 minutes.

In [5] a genetic algorithm was proposed to learn the parameters of an ELECTRE TRI model. This algorithm could be transposed for learning the parameters of a MR-Sort model. However, it is well known in [7] that genetic algorithms which take the structure of the problem into account to perform crossovers and mutations give better results. It is not the case of the genetic algorithm proposed in [5] since the authors’ definitions of crossover and mutation operators are standard.

Learning only the weights and the majority threshold of an MR-Sort model on the basis of assignment examples can be done using an ordinary linear program (without binary or integer variables). On the contrary, learning profiles evaluations is not possible by linear programming without binary variables. Taking these observations into account, [10] proposes an algorithm that takes advantage of the ease of learning the weights and the majority threshold by a linear program and adjusts the profiles by means of a dedicated heuristic. This algorithm uses the following components:

1. a heuristic for initializing the profiles;
2. a linear program learning the weights and the majority threshold, given the profiles;
3. a dedicated heuristic adjusting the profiles, given weights and a majority threshold.

## References

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