Discovery of Characteristics of Patients with Increased Level of Inflammation

Ljiljana Trtica Majnarić1*, Pinar Yildirim2 and Andreas Holzinger2

1Department of Family Medicine, School of Medicine, University of Osijek, Croatia
2Department of Family Medicine, School of Medicine, University of Osijek, Croatia

Abstract

This paper is a study on knowledge discovery for the prediction of characteristics of older patients with increased level of inflammation. The etiology of inflammation is thought to be multifactorial and associated with the development of chronic aging diseases. Chronic low grade inflammation, expressed by slightly elevated serum concentrations of the inflammatory marker C-reactive protein (CRP), has been showed to be associated with increased frailty and overall and specific cardiovascular and noncardiovascular mortality. However, it has not been discovered before which conditions, taken all at once, are associated with elevated levels of inflammation and health status. To answer this question, we used the group of older primary health care attenders, burdened with multiple chronic conditions and described their health status by many aspects. The dataset was composed of 61 low-cost health parameters, many of which were data from patient health records. To predict the characteristics of patients with increased level of inflammation, we used a Linear Regression model and compared the results with some classification algorithms. In this way, we selected 11 relevant predictors of inflammation and explained their meaning according to the existing knowledge. We could realise that many of them represent the components of the highly conserved functional network in which inflammation is the intermediate mechanism, linking these components together. These components include the metabolic, the neuroendocrine and the immune system, proposed as influencing each other during the development of age-related chronic diseases. We have also identified some new components, represented by the parameters indicating inflammation-mediated locomotor system disorders and the pituitary hormone prolactin serum concentration variations. This model, resulted from the knowledge discovery procedure, can be used to provide guidelines for further research on chronic low grade inflammation and for more practical purposes, to help physicians recognizing older persons who are at increased risk for frailty and death.

Keywords: Chronic low-grade inflammation; C-reactive protein (CRP); Aging diseases; Knowledge discovery; Biomedical data mining; Linear regression

Introduction

Inflammation is a well conserved set of responses by the tissues and the body to injury or infection, aimed at returning physiological homeostasis and tissue repair. However, when uncontrolled or chronically maintained, it can also be detrimental and contribute to disease development [1]. During inflammation, the inflammatory cytokines: tumor necrosis factor-a (TNF-a), interleukin-1 (IL-1) and interleukin-6 (IL-6), are secreted to enhance inflammation and help control its course. In many inflammation-related activities, these cytokines show redundancy in action. Specifically, stimulation of acute-phase proteins production in the liver and activation of the hypothalamic-pituitary-adrenal stress axis, are two functions which are almost completely controlled by secretion of the cytokine IL-6. This cytokine is also interesting for its marked pleiotropy and involvement in the regulation of a variety of physiological processes, including hematologic, immune, neurologic, endocrine and metabolic functions [2].

Chronic low grade inflammation, expressed by slightly elevated serum concentrations of inflammatory markers, interleukin-6 (IL-6) and C-reactive protein (CRP), both known as acute phase-reactants, has for a long time been recognised as an independent cardiovascular (CV) risk factor. Especially CRP has been characterised with good prognostic properties [3]. These findings have been showed constant across various populations, including initially healthy individuals, those with marked classical CV risk factors, elderly people and those with clinically manifest CV disease [4,5]. In this regard, positive associations have been found between CRP and body mass index (BMI), fasting glucose and insulin serum concentrations, systolic blood pressure and serum lipids, indicating the involvement of inflammation in the pathogenesis of the insulin resistance (the metabolic) syndrome [6]. This is the complex trait which involves obesity of abdominal (visceral) type and is known to substantially increase the susceptibility for both, diabetes and CV disease. Prevalence of this syndrome increases with age [7]. More generally (Figure 1), evidence suggests that during aging, the tight regulation of inflammatory genes becomes less effective. As the consequence, soluble markers of inflammation, notably CRP, are measurable even when there is no apparent evidence of infection, trauma, or other stressful conditions [8,9]. The etiology is thought to be multifactorial. Some evidence indicates the particular importance of the cumulative antigenic stimulation, along with the loss of immunoregulatory functions in older age. Some other evidence indicates age-related decline in sex hormones, estrogen in women and testosterone in men [8,10]. Recent studies also highlight the importance of the genetic variation in IL-6 gene expression, for variation in serum concentrations of IL-6 and CRP, observed among individuals in the population [11]. The state of chronic low grade inflammation has been considered responsible for the initiation/progression of the main aging chronic diseases, including CV disease, Alzheimer’s and other dementia, osteoporosis, autoimmune and lymphoproliferative diseases [2,8,10]. In addition, elevated levels of inflammation have been...
proposed to play a major role in the development, in older people, of the frailty syndrome, found to be associated with their increased vulnerability for disease and death. The syndrome is characterised with certain phenotypes typically seen in people of advanced age, including decreased lean body mass, osteopenia, anaemia and low cholesterol serum concentrations [10]. Finally, in more general means, associations have been found between elevated IL-6 and CRP serum concentrations and increased overall and specific cardiovascular and noncardiovascular mortality rates. However, it has not been clearly stated, yet, whether these inflammation markers are merely reflective of established chronic diseases, or whether they also stimulate their development [5].

In the group of older, community-dwelling patients, burdened with chronic medical conditions, we wanted to find out hidden relationships in their health data and to link them to elevated serum concentrations of CRP, a well confirmed marker of inflammation. For this purpose, we used a dataset in which the health status of subjects was determined systematically, by many aspects. The model resulted from the knowledge discovery procedure and the selected predictors can be used to provide guidelines for further research and for more practical purposes, to help physicians to recognize older persons, primary health care attenders, who are at increased risk for frailty and death.

Related Work

There were several studies performed for knowledge discovery on C-reactive protein (CRP), but no one comprehensively designed. The first example. Dahlstrom et al. examined whether representation of physicians’ expert knowledge in a simple heuristic model can improve data mining methods in prognostic assessment of patients with rheumatoid arthritis (RA). They interviewed consultants and considered several prognostic indicators which they suggested. Then they carried out a clustering analysis by using a k-means algorithm and some of these suggested variables, such as the erythrocyte sedimentation rate (ESR), C-reactive protein (CRP), the number of swollen joints and the number of tender joints. They identified some prognostic subgroups and found that there is a positive correlation between ESR and CRP in this patient setting [12]. The second example. CRP, a marker of inflammation, has been identified as a risk factor for cardiovascular disease and mortality. Alley et al. used data on adults aged 20 and more from the fourth National Health and Nutrition Examination Survey, a nationally representative cross-sectional survey, and applied statistical methods, T tests and multinomial logistic regression, to examine the association between socioeconomic status and CRP in US adults. They found these associations to occur only at high levels of CRP, while there was no significant difference in the prevalence of moderate or high values of CRP according to the differences in socioeconomic status [13].

Materials and Methods

The sample

The study was conducted in a family practice in an urban area (the town of Osijek, the north-eastern part of Croatia). The region is known for high prevalence of cardiovascular and other chronic disorders, above the average of Croatia [14]. The sample consisted of 93 subjects, aged 50 years and more, who gave their consent. There were 35 males and 58 females, 50-89 years old (median 69), burdened with chronic medical conditions. The study protocol was approved by the local ethics committee.

Parameters description and the modelling process

For the purpose of this study, health data were collected systematically, to determine many aspects of the health-status of patients. Only not costly, easy available parameters were used, a large proportion of which were routinely collected parameters used from patients’ health records. There was a total number of 61 parameters (Tables 1-3). Nominal parameters indicated age and sex, diagnoses of the main chronic diseases, information on drugs use and anthropometric measures. A substantial proportion of subjects were burdened with different chronic disorders (Table 4). A large number of haematological and biochemical laboratory tests were performed, to determine the age-related pathophysiologic changes. These tests included parameters indicating: 1) inflammation (parameters: LE, NEU, EO, MO, LY, CRP, ALFA1 and 2, BETA, GAMA), 2) the nutritional status (parameters: E, HB, HTC, MCV, FE, PROT, ALB, VITB12, FOLNA, HOMCIS), 3) the metabolic status (FGLU, HbA1c, Chol, TG, HDL, IN5), 4) chronic renal impairment (parameters: Clear), 5) latent chronic infections (parameters: CMV, EBV, HBG, HPA), 6) impaired humoral immunity (parameters: RF, IGE, ANA) and 7) the neuroendocrine status (parameters: CORTIS, TSH, FT3, FT4, PRL) (Tables 2 and 3).

| No | Attribute Description | Type |
|----|-----------------------|------|
| 1  | Age Age (years)       | Numeric |
| 2  | Sex (M=Male, F=Female) | Nominal |
| 3  | Hyper Hypertension (yes, no) | Nominal |
| 4  | DM Diabetes mellitus (yes, IGT=Impaired glucose tolerance, No) | Nominal |
| 5  | F Glu Fasting blood glucose (mmol/L) | Numeric |
| 6  | HbA1c Glycosilated Haemoglobin (%) (showing average blood glucose during last three months) | Numeric |
| 7  | Chol Total Cholesterol (mmol/L) | Numeric |
| 8  | TG Triglycerides (mmol/L) | Numeric |
| 9  | HDL HDL-cholesterol (mmol/L) | Numeric |
| 10 | Statins Therapy with statins (yes, no) | Nominal |
| 11 | Anticoag Therapy with anticoagulant/antiaggregant drugs (yes, no) | Nominal |
| 12 | CVD Cardiovascular diseases (yes, no) (myocardial infarction, angina, history of revascularisation, stroke, transient ischaemic cerebral event, peripheral vascular disease) | Nominal |
| 13 | BMI Body Mass Index (kg/m²) | Numeric |
| 14 | WtH Waisthip ratio | Numeric |
| 15 | Arm cir Mid arm circumference (mm) | Nominal |
| 16 | Skinf Triceps skinfold thickness (mm) | integer |
| 17 | Gastro Gastrudodenal disorders (yes, no) (gastritis, ulcer) | Nominal |
| 18 | Uro Chronic urinary tract disorders (yes, no) (recurrent cystitis in women, symptoms of prostatism in men) | Nominal |
| 19 | COPD Chronic obstructive pulmonary disease (yes, no) | Nominal |
| 20 | Aller d Allergy (Rhinitis and/or Asthma) (yes, no) | Nominal |

Table 1: The attributes used in the dataset (1-20).
The same dataset, although with minimal corrections in variables selection, we have already used for solving other complex research tasks [15-17]. We wanted to show that by collecting health data systematically and by using data mining techniques for their analysis, it is possible to elucidate important clinical contexts of some complex age-related medical conditions, usually presented with multi-morbidity (the simultaneous presence of several disorders at the same individual).

The modeling procedure can reveal many important elements of the networked pathophysiology reaction. The results can be used for prediction purposes, to support medical doctors in their decision making, or to accelerate research, by mapping the main components and relationships within the network. In this terms, this procedure may be recommended as a new integrative approach, with the potential to add value to much more complex and time and cost consuming systems 

### Linear Regression

Regression analysis is an important statistical method for the analysis of medical data. It enables the identification and characterization of relationships among multiple factors [19]. Linear regression is a well known method of mathematical modeling of the relationship between a dependent variable and one or more independent variables. Regression uses existing (or known) values to forecast the required parameters [20]. The goal of this model, is to predict the response to n data points($x_1,y_1),(x_2,y_2),...,(x_n,y_n)$ by a regression model given by

$$y = a_0 + a_1x$$

Where $a_0$ and $a_1$ are the constants of the regression model. A measure of goodness of fit, that is, how well $a_0 + a_1x$ predicts the response variable $y$ is the magnitude of the residual $\epsilon_i$ at each of the n data points.

$$\epsilon_i = y_i - (a_0 + a_1x)$$

Table 2: The attributes used in the dataset (21-40).

| No | Attribute | Description | Type |
|----|-----------|-------------|------|
| 21 | Dr aller  | Drugs allergy (yes, no) | Nominal |
| 22 | Analg     | Therapy with analgetics/non-steroidal anti-inflammatory drugs (yes, no) | Nominal |
| 23 | Derm      | Chronic skin disorders (yes, no) (chronic dermatitis, dermatomyositis) | Nominal |
| 24 | Neo       | Malignancy, including skin malignancy (yes, no) | Nominal |
| 25 | OSP       | Osteoporosis (yes, no) | Nominal |
| 26 | Psy       | Neuropsychiatric disorders (yes, no) (anxiety/depression, Parkinson’s disease, cognitive impairments) | Nominal |
| 27 | MMS       | Mini Mental Score – test for screening on cognitive dysfunction Max Score>30 Score <24 indicates cognitive impairment | integer |
| 28 | CMV       | Cytomegalovirus specific IgG antibodies (IU/ml) | Numeric |
| 29 | EBV       | Epstein-Barr virus specific IgG (IU/ml) | Numeric |
| 30 | HBG       | Helicobacter pylori specific IgG (IU/ml) | Numeric |
| 31 | HPA       | Helicobacter pylori specific IgA (IU/ml) | Numeric |
| 32 | LE        | Leukocytes Number × 10^9/L | Numeric |
| 33 | NEU       | Neutrophils % in White Blood Cell differential | Numeric |
| 34 | EO        | Eosinophils % in White Blood Cell differential | Numeric |
| 35 | MO        | Monocytes % in White Blood Cell differential | Numeric |
| 36 | LY        | Lymphocytes % in White Blood Cell differential | Numeric |
| 37 | E         | Erythrocytes number × 10^12/L | Numeric |
| 38 | HB        | Haemoglobin (g/L) | integer |
| 39 | HTC       | Haematocrit (erythrocyte volume blood fraction) | Numeric |
| 40 | MCV       | Mean cell Volume (fL) | Numeric |

Table 3: The attributes used in the dataset (41-61).

| No | Attribute | Description | Type |
|----|-----------|-------------|------|
| 41 | FE        | Iron (gL) | Numeric |
| 42 | PROT      | Total serum proteins (g/L) | Nominal |
| 43 | ALB       | Serum albumin (g/L) | Numeric |
| 44 | Clear     | Creatinine clearance (mL/min) | Nominal |
| 45 | HOMCIS    | Homocistein (µmol/L) | Numeric |
| 46 | ALFA1     | Serum protein electrophoresis (g/L) | Numeric |
| 47 | ALFA2     | Serum protein electrophoresis (g/L) | Numeric |
| 48 | BETA      | Serum protein electrophoresis (g/L) | Numeric |
| 49 | GAMA      | Serum protein electrophoresis (g/L) | Numeric |
| 50 | RF        | Rheumatoid Factor level (IU/ml) | Numeric |
| 51 | VITB12    | Vitamin B12 (pmol/L) | Numeric |
| 52 | FOLNA     | Folic acid (mM/L) | Numeric |
| 53 | INS       | Insulin (µIU/L) | Numeric |
| 54 | CORTIS    | Cortisol in the morning (µmol/L) | Numeric |
| 55 | PRL       | Prolactin in the morning (mIU/L) | Numeric |
| 56 | TSH       | Thyroid-stimulating hormone (IU/ml) | Nominal |
| 57 | FT3       | Free triiodothyronine (pmol/L) | Numeric |
| 58 | FT4       | Free thyroxine (pmol/L) | Numeric |
| 59 | ANA       | Antinuclear antibodies (autoantibodies) (µIU/ml) | Numeric |
| 60 | ICE       | IgG (MU/L) | Numeric |
| 61 | CRP       | C-reactive protein (mg/L) | Nominal |
Ideally, if all the residuals $\epsilon_i$ are zero, one may find an equation in which all the points lie on the model. Thus, minimization of the residual is an objective of obtaining regression coefficients.

The most popular method to minimize the residual is the least squares methods, where the estimates of the constants of the models are chosen such that the sum of the squared residuals is minimized, that is minimize $\sum x_i y_i^2$ [20]. Measures of association provide an initial impression of the extent of statistical dependence between variables. If the dependent and independent variables are continuous, a correlation coefficient ($r$) can be calculated as a measure of the strength of the relationship between them.

In many cases, the contribution of a single independent variable does not alone suffice to explain the dependent variable $Y$. If this is so, one can perform a multivariate linear regression to study the effect of multiple variables on the dependent variable [19].

In the multivariate regression model, the dependent variable is described as a linear function of the independent variables $x_i$, as follows:

$$ y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \ldots + \beta_n x_n $$

The model permits the computation of a regression coefficient $\beta_i$ for each independent variable. Just as in univariable regression, the coefficient of determination describes the overall relationship between the independent variables $x_i$ and the dependent variable $y$. It corresponds to the square of the multiple correlation coefficients, which is the correlation between $y$ and $b_1 x_1 + b_2 x_2 + \ldots + b_n x_n$ [19].

**Decision Table**

Decision Table summarizes the dataset with a ‘decision table’, a decision table contains the same number of attributes as the original dataset, and a new data item is assigned a category by finding the line in the decision table that matches the non-class values of the data item. This implementation employs the wrapper method to find a good subset of attributes for inclusion in the table. By eliminating attributes that contribute little or nothing to a model of the dataset, the algorithm reduces the likelihood of over-fitting and creates a smaller, more condensed decision table [21-23].

**Kstar**

K is an instance-based classifier, the class of a test instance is based upon the class of those training instances similar to it, as determined by some similarity function. The underlying assumption of instance-based classifiers is such as K [23,24].

**M5Rules**

The method for generating rules from model trees, which is called M5rules, is straightforward and works as follows: a tree learner is applied to the full training dataset and a pruned tree is learned. Next, the best branch (according to some heuristic) is made into a rule and the tree is discarded. All instances covered by the rule are removed from the dataset. The process is applied recursively to the remaining instances and terminates when all instances are covered by one or more rules. This is a basic separate-and-conquer strategy for learning rules: however, instead of building a single rule, as it is usually done, we build a full model tree at each stage, and make its “best” branch into a rule. This avoids potential for over-pruning called hasty generalization. In contrast to PART (Partial Decision Trees) that employs the same strategy for categorical prediction, M5Rules builds full trees instead of partially explored trees. Building partial trees leads to a greater computational efficiency, and does not affect the size and accuracy of the resulting rules [23-25].

**Random subspace**

Random Subspace Method proposed by Ho [26]. The main idea of this technique is that, instead of using a single sub-space for classification, multiple subspaces are constructed from the original space by a random procedure. The Random Subspace technique produces a classifier using each random subspace. Then the final classification decision is achieved by combining all classification results. Although each classifier may not produce a good classification, their combination can be much better than using a single sub-space [27,28].

**Regression by discretization**

The regression by discretization approach allows the use of a classification algorithm in a regression task. It works as a pre-processing step in which the numeric target value is discretized into a set of intervals. A regression scheme that employs any classifier on a copy of the data that has the class attribute (equal-width) discretized. The predicted value is the expected value of the mean class value for each discretized interval (based on the predicted probabilities for each interval) [27-29].

**RepTree**

Quinlan [30] first introduced Reduced Error Pruning (REP) as a method to prune decision trees. REP is a simple pruning method, although it is sometimes considered to over-prune the tree. A separate pruning dataset is required, which is considered a disadvantage of this method because data is normally scarce. However, REP can be extremely powerful when it is used with either a large number of examples or in combination with boosting. The pruning method that is used is the replacement of a subtree by a leaf representing the majority of all examples reaching it in the pruning set. This replacement is done if this modification reduces the error, i.e., if the new tree would give an equal or fewer number of misclassifications [27-31].

**Results and Discussion**

**Experimental results**

We used a Linear Regression model to predict characteristics of patients with an increased level of inflammation and compared the results with some classification algorithms (Decision Table, Kstar, M5rules, Random Subspace, Regression by discretization and RepTree). A comparison of these techniques was presented. WEEKA 3.6.8 software was used for analysis. WEKA is a collection of machine learning algorithms for data mining tasks and it is open source software. The software contains tools for data pre-processing, classification, regression, clustering, association rules and visualization [32,33].

Linear regression method was applied on our dataset and following regression function of the variables was generated:

\[
\begin{align*}
\text{CRP} = & -0.0867 \times \text{Age} + 0.2304 \times \text{BMI} + 1.6537 \times \text{Alog=No} + 1.8415 \times \text{OSP=No} + 0.287 \times \text{MMS} + 0.016 \times \text{EBV} + 0.1557 \times \text{HB} \\
\end{align*}
\]
As can be seen from Table 5, Regression by discretization algorithm generated good results. This algorithm obtained a high correlation coefficient of 0.2441. Also, the algorithm has lower error rates among the other algorithms with a mean absolute error of 2.0757 and a root mean square error of 3.8528.

Discussion (Explanation of the Results in Light of the Current Knowledge)

The Linear Regression method, applied on the dataset that comprehensively describes the health status of the group of older persons with multiple morbid conditions, extracted a set of 11 variables, relevant enough to allow a significant model generation, in order to predict characteristics of persons with increased level of inflammation. A relatively large number of variables selected in the model are indicative of inflammation as a multifunctional disorder. This is not unexpected, since inflammation is known as an intermediate mechanism which links many interacting components, from the genetic and molecular to the phenotypic levels, into the common functional network [35]. The second explanation is that these results are reflective of a great inter-individual variability in clinical characteristics of older persons with increased level of inflammation. This explanation is in line with the recent studies of aging showing the aging phenotype, not as an uniform pattern, but as a heterogeneous mosaic of aging traits, which is a result of the complex interactions of environmental, genetic and stochastic factors [36]. In the same line, the studies of the frailty syndrome, the age-related phenotype characterised with weakness and motor dysfunction, showed the association of this syndrome with the physiological multisystem dysregulation, that means that the likelihood of frailty increases with increasing number of dysregulated physiological systems [37].

According to these studies, aging is considered as the complex remodeling process, that means that there is not a simple linear decline in the physiological functions over time, but instead, compensatory reactions follow detrimental changes, while the loss of some functions is maintained together with the over activation of some others [35,36]. Successful modeling of age-related functions considers the existence of efficient defense mechanisms. In this regard, chronic low grade inflammation is thought to be a result of the imbalance between inflammatory and anti-inflammatory regulatory networks and, as such, as the main driving force for age-related diseases development [35-38]. In other words, inflammation can be considered as a reflective of unsuccessful modeling, thereby linking the aging process with increased morbidity and mortality rates. The studies also say that when this point in the course of human life has passed, the mortality curve starts to slowdown instead of to continuously rise, reflecting the positive selection of very old individuals (85 and over) who are characterised with lower level of inflammation and better coping mechanisms, than their younger counterparts loaded with chronic aging diseases [38,39].

All this discussion was needed to help us explaining the obtained results on the negative association between the variable Age and the stage of diagnosis, proposing further investigations on the interaction between age, inflammation and other comorbidities.
target variable CRP. Namely, this result might be due to the fact that the median age of the subjects in this sample was 69 years, indicating the age period when the disease burden is the largest. Obviously, the statement: the bigger the number of years of life, the higher inflammation level, does not fairly deal with our results, thereby supporting the remodeling theories of aging. The answer to the question why the variable sex (male/female) maintained unselected in the model, might not be due the lack of its relevance for the pathophysiology of inflammation, but rather the reason might be due to the mixed sample, or the insufficient sensitivity of the input variables for the differences in gender. In this regard, evidence suggests that women and men use different pathways in achieving age-related morbidity changes, or in longevity. In comparison to men, women are less dependent on genetics and more on environmental and societal factors [40].

Selection of the majority of other variables in this model also argues towards the proof of the conceptual framework of the remodeling theory of aging. According to this theory, the three main physiological systems, the metabolic, the neuroendocrine and the immune systems, are interconnected and capable of influencing each other during the age-related remodeling [35,38,41,42]. Chronic inflammation has been recognised as the common mechanism which links deteriorative changes in these systems together, accelerating the development of age-related diseases [38]. Elements of these systems and their relationships with chronic inflammation (indicated by the variable CRP) are likely to be captured by the selected variables BMI, MMS, PRL, EBV and ANA. Explanations for these variables are provided in the sections below.

In regard to the variable BMI, this is an anthropometric measure and well accepted marker of obesity. Obesity is known inflammatory condition, due to high secretory activity of the adipose tissue, that may promote insulin resistance (impaired insulin-mediated glucose metabolism), subsequently leading to glucose intolerance and diabetes, both disorders known as the major CV risk factors [43]. Among many secretory products with the origin in the adipose tissue, the main role in the pathogenesis of insulin resistance is prescribed to the inflammatory cytokines TNF-α and IL-6. Insulin resistance is considered to be the key mechanism of the insulin resistance (the metabolic) syndrome, clinically expressed by the cluster of several metabolic CV risk factors, the frequency of which increases with age, including: obesity; hypertension, impaired glucose tolerance, hyperinsulinemia and dyslipidemia [7]. Some other factors have subsequently been added to the syndrome, including impaired coagulation/fibrinolysis, increased blood viscosity and low grade chronic inflammation [44,45]. In this context we can consider the variables HB and HTC, selected in the regression model. Namely, these variables have already been identified as blood viscosity measures and, thus, as the part of the metabolic syndrome [45]. Specifically, the variable HB, indicating anaemia, can also be viewed from the perspective of the frailty syndrome, which in older people is characterised by muscle wasting and low anthropometric performances [10]. Related to this, it has already been recognised that impaired body composition, not just obesity, but also the imbalance between the fat and the muscle contents, may influence the level of inflammation. Older age is associated with changes in both, in the level of inflammation and in body composition [46,47]. It is not surprisingly then that chronic inflammation has been recognised as the main reason why the frailty syndrome, despite the weight loss, increases the risk for the development of CV disease [48]. Based on this discussion, it is possible that, in addition to the variable HB, the variable BMI also has a dual function, indicating physical weakness and weight loss, rather than (or in addition to) the state of being overweight. The general implications of these results would be that any state associated with great deviations in body compositions (including either malnourishment or overweight) can also increase the level of inflammation. This statement once again argues towards the remodeling theory of aging, indicating that there is the evolutionary necessity for the close relationships between the metabolic and the immune (inflammation-related) functions [49].

The selected variable ALFA1 indicates the increased hepatic synthesis of acute-phase proteins and in this context can be used as a complementary one to the variable CRP, that is, as a marker of inflammation [50]. We can speculate on the reasons why some other variables that indicate increased inflammation and have been used as the model’s input, including: LE, MO, LY, NEU and ALB, have not been not selected in the model. The reasonable explanation is that the inflammatory cytokine IL-6, known to be in the background of the increased synthesis of CRP, taken as the target variable in this model, can trigger a particular part of the inflammatory response [50].

According to the large body of evidence, there are multiple connections between the immune, the endocrine and the central nervous systems (CNS) [51]. The CNS regulates the immune system through neuroendocrine pathways and the autonomic nervous system, while the immune system sends signals to the brain via humoral factors, mainly cytokines, but also other immune mediators. In this terms, many cytokines and their receptors have been identified in the brain. Reversely, receptors for hormones, neurotransmitters and other neuropeptide mediators can be found on lymphocytes and other immune cells [51,52]. Knowledge on these communications at the molecular and cellular levels helps us understanding the observed clinically sound associations between inflammation and the immune mediated diseases, on the one hand, and the endocrine system related and mood disorders related diseases, on the other hand [51]. In this context, it has been known for a long time that the main neuroendocrine pathway which regulates the immune system functions is the hypothalamus-pituitary-adrenal (HPA) stress axis [51]. Reversely, the most powerful stimuli of this axis, in humans, is likely to be the cytokine IL-6, an inductor of acute-phase proteins synthesis, including also CRP, the main clinical marker of low grade inflammation [2]. Recent evidence indicates even more complex regulatory network to exist, including the neuroendocrine stress axis, obesity, inflammation and insulin resistance, linking the neuroendocrine, immune/ inflammatory and metabolic pathways, into the common network [49]. In the same line of evidence, cognitive function decline (reflective of the brain degenerative/inflammatory processes) has been recognised as the part of the frailty syndrome (reflective of chronic inflammation and changes in body composition) [53]. Within this complex context, described above, we can find explanations for the selected variables MMS and PRL. The variable MMS indicates cognitive function decline, proposed to be the result of the brain aging and neurodegenerative brain disorders [54]. Evidence indicating that increased level of systems inflammation, together with metabolic and vascular changes, due to insulin resistance, are proposed triggering/accelerating factors in the pathogenesis of neurodegenerative disorders, can explain this observed link between the variable MMS and the target variable CRP. Another selected variable, PRL, indicating hormone prolactin serum concentration variations, might be also considered as the marker of the brain aging and degenerative brain disorders, elbeit focus shifted towards the pituitary gland, the central point of neuroendocrine control [55]. In line with the previous discussion is also the evidence indicating an association between the increased serum prolactin concentrations, insulin resistance and increased risks for CV disease development [56]. In fact, evidence is still insufficient on whether there is a pro-inflammatory or anti-inflammatory role of this hormone in the pathophysiology network of neuroendocrine and metabolic age remodelling, or whether there is increase, or decrease in this hormone.

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serum concentrations [57]. Our results also support this common notion that there is the complex remodeling process along with the course of age-related disease development, rather than simple linear decline in functions.

The selection of the variables EBV and ANA in the predictive model of inflammation can also be explained in the light of existing evidence and it is associated with the immuno-regulatory role of inflammation [2]. In this regard, one line of evidence indicates the role of cytokine IL-6 in enhancing humoral immunity, according to its described function as a stimulator of B-lymphocytes proliferation/maturaion and immunoglobulins (antibodies) production [8]. The second line of evidence emphasises the role of low grade inflammation in accelerating aging of the immune system (immunosenesence). This happens in conditions of aging and the presence of age-related diseases.

The process is characterised by the failure of specific immunity, reactivation of latent infections (in our sample indicated by the selected variable EBV) and spontaneous production of autoantibodies, even in the absence of autoimmune diseases (indicated by the selected variable ANA) [58]. In this context, ANA is a routinely used diagnostic marker in rheumatic autoimmune diseases, but its serum concentrations also increase spontaneously with age and are found to be elevated, as well in healthy elderly people, as in associations with different chronic diseases [59,60]. We can only speculate on the reasons why the variable EBV, indicating latent EBV infection, was selected in the model, but not other three variables (CMV, HBG and HPA) that indicate latent cytomegalovirus and Helicobacter pylori infections. According to the available evidence, the reason might be the synergistic action between EBV infection and chronic low grade inflammation in B lymphocytes stimulation [58].

Apart from these well known relationships involving chronic low grade inflammation, described above, our results also add new values to the knowledge of inflammation, by indicating the involvement of the locomotor system chronic inflammatory disorders, in this complex pathophysiologic network. This is indicated by the two nominal variables selected in the model, Analg=No and OSP=No. Related to these findings, osteoporosis, indicated with the variable OSP, is a condition characterised with increased bone fragility and risks for fracture, due to enhanced bone resorption and/or decreased formation. This is a well known inflammation-mediated aging disease which predominantly affects women [2,18]. In fact, multiple pathogenetic mechanisms have been proposed for this disease, including genetic, biomechanical, metabolic, hormonal and inflammatory factors [18]. We can speculate that the option "OSP=No" of this selected variable does not mean the negative selection of subjects free from this disease, but on the contrary, implicates early stages of this disease development, when the degree of inflammation, produced locally in bone tissue, is also likely to be high. These early stages are complementary to the diagnosis of osteopenia (the condition in which the bone tissue content is reduced, but at lesser degrees than in osteoporosis). In contribution to this statement is the fact that osteopenia has been recognised as the component of the frailty syndrome, known to be an inflammation-mediated condition [10]. Once when this mentioned bone disease turns into its advanced stage, the fairly large bone loss is unable to elicit any inflammatory reaction. Similar explanation might state for the second variable, "Analg=No", indicating older people who does not use analgesics and non-steroidal anti-inflammatory drugs. This statement is likely to exclude patients with severe osteo-arthritis (who does use these drugs), allowing for those ones with early stages of this disease. In this regard, recent evidence suggests that osteo-arthritis, a condition characterised with the fragility and loss of cartilage tissue in the weight-bearing joints, including prevalently the knees and the hips, has been recognised as an inflammation-mediated disorder and important source of inflammatory mediators [61]. Systemic inflammation, associated with this condition, affects the susceptibility of these patients to the development of chronic aging diseases, notably atherosclerotic CV and Alzheimer’s disease. The problem with this provided explanation for the variable "Analg=No" is the lack of the classification of osteo-arthritis disease, which makes the relationships between the disease activity and the level of inflammation difficult. Additional thing is that use of analgesics can mask the inflammation state. This all enables only general statement upon the selection of this variable, indicating analgesics use (or non use), and the existence of the conditions complementary to osteo-arthritis, without taking into account the exact position of this variable in the ordinal scale values, by means of altering the valus "yes" or "no".

Conclusions

We generated the model to predict features of older people with low grade inflammation, by using a large set of simple health data collected in a way to determine the health status of patients from many aspects, systematically. The model is likely to capture all main relationships known to link the IL-6 gene expression (a hallmark of inflammation) with the metabolic, the neuroendocrine and the immune system pathways, implicating inflammation as a part of the highly conserved pathophysiology network. Apart from these core elements, some new components of the network have been emphasised, including inflammation-mediated bone and joint disease and particular neuroendocrine pathways, with the pituitary hormone prolactin being involved in. When the existing knowledge was used to explain these variables selection, some interesting facts have emerged, leading to new hypotheses generation. One hypothesis states that there is an association between increased level of inflammation and changes in body’s composition, which might be the main driving mechanism of this network. Another hypothesis deals with the notion that there is a nonlinear remodeling of the metabolic, the immune and the neuroendocrine pathways during aging and age-related disease development. Whether the components and pathways outlined in this model are of universal importance, or whether they just reflect the characteristics of the study sample, have to be explored by repeating the same procedure on other samples. This presented approach based on using systematically collected, easily available health parameters, can be used to support decision-making when planning preventive strategies, to initially mapping components when an unexplored problem is to be solved, or to supplement some other system’s methods. Even more important, this approach could be useful to seeking for simple, low-cost markers, for making prediction in the area of chronic aging disease. According to our results, parameters indicating analgesics use (variable Analg), increased blood viscosity (variable HTC) and acute-phase response (variable ALFA1), deserve such attention.

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