Visualizing correlations among Parkinson biomedical data through information retrieval and machine learning techniques

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Received: 30 August 2020 / Revised: 3 December 2020 / Accepted: 5 January 2021 / Published online: 27 February 2021 © The Author(s) 2021

Abstract
In the last few years, the integration of researches in Computer Science and medical fields has made available to the scientific community an enormous amount of data, stored in databases. In this paper, we analyze the data available in the Parkinson’s Progression Markers Initiative (PPMI), a comprehensive observational, multi-center study designed to identify progression biomarkers important for better treatments for Parkinson’s disease. The data of PPMI participants are collected through a comprehensive battery of tests and assessments including Magnetic Resonance Imaging and DATscan imaging, collection of blood, cerebral spinal fluid, and urine samples, as well as cognitive and motor evaluations. To this aim, we propose a technique to identify a correlation between the biomedical data in the PPMI dataset for verifying the consistency of medical reports formulated during the visits and allow to correctly categorize the various patients. To correlate the information of each patient’s medical report, Information Retrieval and Machine Learning techniques have been adopted, including the Latent Semantic Analysis, Text2Vec and Doc2Vec techniques. Then, patients are grouped and classified into affected or not by using clustering algorithms according to the similarity of medical reports. Finally, we have adopted a visualization system based on the D3 framework to visualize correlations among medical reports with an interactive chart, and to support the doctor in analyzing the chronological sequence of visits in order to diagnose Parkinson’s disease early.

Keywords Biomedical data analysis · Health information visualization · Information retrieval · Machine learning

1 Introduction
In the last few years, the application of information technology to the medical field has led to the generation of large data sets that can be analyzed to identify the risk of disease and...
support the doctor in the diagnosis, therapy of the patient, and prevention of the disease itself.

The dataset Parkinson’s Progression Markers Initiative (PPMI) is a comprehensive observational, multi-center study designed to identify Parkinson’s Disease (PD) progression biomarkers important in the search for treatments of this type of illness [24]. PPMI is a “open source” reference study started in 2010 which aims at finding some new biomarkers, i.e., indicators of disease that represent missing links, important in the research of better treatments for PD. The PPMI dataset is the result of the cooperation of numerous researchers; it is constituted by a large quantity of data and samples collected and acquired by the participants, all volunteers, some of whom suffer from the disease. It is updated every 8 months and available online. The search for these biomarkers is carried out by:

- Decide on standardized protocols for acquisition, transfer and analysis of clinical, imaging and biologic data that can be used by the PD research community.
- Develop a comprehensive dataset of clinical data and image and biological samples which is uniformly acquired. It can be adopted to estimate the mean rates of change and the variability around the mean of the collected data in early PD, patients prodromal PD subjects, and PD subjects with a mutation of the \( LRKK_2 \), \( GBA \) or \( SNCA \) genes.

In [26], the authors proposed a technique to identify a correlation between the biomedical data belonging in the PPMI dataset for verifying the consistency of medical reports formulated during the visits and allow to correctly categorize the various patients. To correlate the information of each patient’s medical report, an Information Retrieval technique (named Latent Semantic Analysis) has been adopted. This technique constructs a concept space on selected patient information. This information is exploited to group and classify patients into affected or not by using clustering algorithms. Results revealed that the proposed technique reached 95% of effectiveness in the classification of patients.

However, the aforementioned paper had some limitations including:

- the tables selected for the analysis included only the patient’s diagnostic reports, but there was no information regarding the patients’ habits and lifestyle;
- only a lexical technique named Latent Semantic Analysis technique is used;
- it lacks a visualization system in order to make the doctor’s understanding of the patient’s status more immediate and simple and therefore facilitate the patient’s diagnosis.

This paper is an extension of [26]. In particular, the following improvements have been added:

- Further improvement of the analysis using the PPMI dataset updated in February 2020, which includes additional information such as information regarding personal habits and lifestyle for each patient;
- The use and comparison with machine learning techniques such as Text2Vec and Doc2Vec;
- The adoption of a visualization system to support the doctor in diagnosing the chronological sequence of visits in order to diagnose Parkinson’s disease early.

1https://www.ppmi-info.org
Information Retrieval (IR) and Machine Learning (ML) techniques have been adopted to correlate the information of each patient’s medical report. In particular, i) Latent Semantic Analysis (LSA) technique is used to construct a concept space on patient information; ii) A technique based on the concept of Word Embedding, a methodology of natural language processing composed of a set of tools, language models, and learning techniques that allow the representation of documents through the use of vectors with real components as Tex2Vec, which through a neural network and a term-document matrix creates a numerical representation of the documents and Doc2Vec a semantic machine learning technique, can be used in different methods depending on the field of application, in particular it can be used with pre-trained models. iii) clustering algorithms are adopted to group patients according to the descriptions of their medical reports; iv) a data analysis phase is performed to classify the patient groups and finally visualize the correlations to perform the diagnosis.

The paper is structured as follows: Section 2 discusses related work, Section 3 describes the PD and the PPMI dataset, while Section 4 presents the analysis process; Section 5 reports the data analysis we performed on the considered dataset and a discussion of the results; Section 6 shows the visualization of our dataset. Finally, Section 7 concludes the paper.

2 Related work

The choice to adopt IR techniques in the medical field is increasingly common and could be a common practice in the future to support medical diagnosis. Recently, these techniques have been applied to biomedical data [9, 16, 22, 23]. In particular, Gefen and Miller [16] use the LSA on medical records related to congestive heart failure to identify patterns of associations between terms of interest. Similarly, Li and Wu [22] propose the KIP software tool for the identification of topical concepts from medical documents. An advantage of these studies is that the knowledge of the diagnosis and the treatment to be applied could be kept up to date, while one of the disadvantages could be that medical documents are analyzed without considering the real meaning of the words and this could cause errors and confusion.

Mao and Chu [23] propose a phrase-based vector space model for indexing medical documents, whilst Chou and Chang [9] have developed an IR system for doctors and patients to retrieve similar medical case records or related documents from various databases, deriving the similarity between the concepts using their relationship based on knowledge and the similarity between two sentences was measured using their root overlaps and the similarity between the concepts.

While as regards ML techniques, documents can be classified in three ways: unsupervised, supervised and semi-supervised methods, these techniques are widely used for the extraction of knowledge in biomedical data [2, 5–7, 19, 35]. In particular Beam et al. [3] presents a new reference methodology based on statistical power specifically designed to test incorporations of medical concepts, called cui2vec. This study, however, has limitations as most of the sources of health data are not easily shared, which limits the study to small local data sources. Finally, they provide a downloadable set of pre-trained embeds as well as an online tool for interactive exploration of the embeds.

Dynomant et al. [12] used the Doc2Vec algorithm to train models that allow you to vectorize documents on the PubMed database to analyze whether you can replace the statistical model PubMed Related Articles (pmra). This algorithm was able to link documents sharing MeSH labels in a similar way the pmra did.
Chen and Sokolova [8] they used Word2Vec and Doc2Vec unsupervised to analyze sentiment summary reports. They aimed to detect whether there is any latent prejudice towards or against a particular disease. They used SentiWordNet to establish a golden sentiment standard for data sets and evaluate the performance of the Word2Vec and Doc2Vec methods.

The visualization of the data is very important, especially in the medical field, it is used to synthesize all the information relating to a patient but in particular, the visualization of the data is used to support decisions and diagnosis [28, 33].

Especially Lesselroth et al. [21] underline some problems concerning the management of information at the point of care and propose strategies for a better visualization of data including multimedia displays, clinical dashboards, concept-oriented views, metaphor graphics and probability analysis.

Ropinski et al. [30] examine glyph-based visualization techniques that have been exploited when viewing spatial multivariate medical data. To classify these techniques, they derive a taxonomy of the properties of glyphs that is based on classification concepts established in the information display.

Blaas et al. [4] presented to highly interactive, coordinated view-based visualization approach that has been developed for dealing with multi-field medical data. This type of visualization is based on intuitive interaction techniques and integrates analysis techniques from pattern classification to guide the exploration process.

In our work, we use IR and ML techniques to identify eventual correlations between documents in order to recognize the different classes of patients based on the specific medical reports. Furthermore, we perform a semantic analysis, Text2Vec and Doc2Vec techniques on the medical report with the aim of highlighting the most characterizing keywords for Parkinson’s disease. Finally, a visualization system was adopted to support diagnosis for doctors.

3 Dataset

PD is a neurodegenerative disease, belonging to the “Movement Disorders” category. It is originates from the degeneration of neurons in the brain that produce the neurotransmitter “dopamine”. In the early stages of the disease, the most obvious symptoms are related to movement, and include tremors, stiffness, bradykinesia, postural instability, slowness in movement and difficulty walking. Afterwards, cognitive and behavioral problems may arise, such as dementia, depression, psychotic features, autonomic dysfunction, oculomotor abnormalities [17]. In particular, in PD the production of dopamine in the brain decreases consistently and the reduced levels of dopamine are due to the degeneration of neurons in an area called substantia nigra (cell loss is over 60% at the onset of symptoms). Moreover, from the marrow to the brain accumulations of a protein called alpha-synuclein begin to appear. This insoluble protein accumulates within neurons forming inclusions, called Lewy bodies [10]. The causes of PD are not yet known but, it seems that there are multiple elements that contribute to its development. These factors are mainly:

- Genetic mutations: among these, the mutation of the LRRK 2, named also PARK8, is the most relevant. The heterozygous mutation, 2877510 G → A, of this gene is the most commonly described, representing the majority of familial cases of cases of idiopathic PD [10].
- Toxic factors and work exposure such as some insecticides or herbicides.
The diagnosis of PD remains a clinical diagnosis because there are neither objective tests nor specific biochemical and neuroradiological markers. However, in the last decade one of the objectives of the research has been to improve the specificity of the classical diagnostic criteria.

The diagnosis of Parkinson’s disease remains a clinical diagnosis since there is no objective test or specific biochemical and neuroradiological markers. In the last decade, however, one of the research objectives has been to improve the specificity of classical diagnostic criteria: in fact, the “United Kingdom Parkinson’s disease Society Brain Bank” has proposed clinical criteria that are still widely used in clinical practice and research protocols. These diagnostic criteria establish that the sign necessary to diagnose Parkinson’s disease is bradykinesia or akinesia, associated with at least one of the other so-called major signs mentioned above, i.e. muscle stiffness, tremor at rest and postural instability. These diagnostic criteria underline how clinical diagnosis is based on the combination of some “cardinal” motor signs and on the exclusion of symptoms considered “atypical” [17]. In conclusion, the symptoms of Parkinson’s disease manifest themselves differently in different patients, who may experience some symptoms and not others, and also the rate at which the disease progresses varies from individual to individual. For this, the misdiagnosis rate can be relatively high.

The activity conducted by PPMI is an “open source” study, the data and samples collected and acquired by volunteer participants, affected and not by the disease, will allow the development of a database and a complete biorepository, which is currently available online and updated every eight months. Being data collected from patients from various continents one of the main tasks of PPMI is to coordinate the management of the various data, defining a protocol for the collection and coding of data. The elaborated repository can be downloaded by accessing the portal of the PPMI site to allow the scientific community to conduct complete and exhaustive research.

The PPMI dataset is the result a clinical study based solely on observations aiming at fully evaluating significant cohorts of interest by using advanced imaging, biological sampling, clinical and behavioral assessments to identify biomarkers related to the progression of PD. The collected data may be helpful in the research of therapies to slow down or stop this progression.

The complete dataset consists of 145 files in CSV format, containing information about six macro-areas listed in the following:

- **Biospecimen**: collection of data related to clinical tests, such as blood collection, DNA and lobar puncture.
- **Imaging**: use of imaging techniques, such as Magnetic Resonance, PET and DatScan through which it is possible to observe non-visible areas of the organism.
- **Medical History**: clinical history of patients from the first symptoms of the disease to the latest health conditions. The collection includes possible side effects of the medicines taken, results of neurological examinations, physical and so on.
- **Motor MDS-UPDRS**: collection of motor disturbance data through the use of the MDS-UPDRS scale to evaluate the stage of Parkinson’s disease.
- **Non Motor Assessments**: collection of data related to cognitive and emotional-behavioral disorders.
- **Study Enrollment**: collection of conclusive data on particular studies conducted on patients.
- **Found**: collection on personal habits and lifestyles data.
Figure 1 shows the data model of the PPMI dataset, in which there are five entities:

- **Patient**: represents the set of patients participating in the study.
- **Event**: represents the set of tables that refer to the visits and analyzes to which patients are subjected.
- **Biospecimen Analysis Result**: represents the set of tables in which the analysis of the results for the controls to which the patients have undergone are present.
- **Family History**: a set of tables that describe the patient’s family histories.
- **Medication**: a set of tables in which the medicines taken by patients are cataloged.

The entities described above are connected through relationships:

- **R**: represents the relationship that exists between the entities, Event, Patient, and Biospecimen Analysis Result, through the PATH, that is the unique attribute that identifies the patients;
- **HAS**: is the relationship that represents the connection between patients and their family history;
- **ASSUMES**: is a relationship that associates to each patient the medicines he takes.

### 4 The analysis process

In this section we present the process that allows us to find the correlation between the information on the visits and the patient’s disease status, which may be: sick (PD and GENPD), healthy (HC, GENUN and SWEDD) and healthy with typical symptoms of the disease (PRODROMAL, i.e., subjects suffering from insomnia and have mutations of the \( LRRK2 \) gene). The version of the PPMI data used is updated in February 2020.

#### 4.1 Dataset pre-processing

The preprocessing of a text is one of the key components for classifying the text, it is carried out by cleaning from widely used terms, conjunctions, adverbs, and in general the so-called “empty” words, but also the removal of additional spaces [32].
After an initial analysis of the dataset, a skimming of the tables was carried out by selecting only those of interest for the text analysis. This selection was made only on the tables in which symptoms strongly correlated with the disease appear, excluding all those containing diagnostic information. Moreover, we evaluated only the screening visits (SC), the basic line visits (BL), the visits 1 – 15 (V01-V015), the symptomatic therapy visits (ST) and the adverse events visits (LOG). The process examines them in chronological order.

Following the selection and recognition phase, a modification of the tables is made, transforming in textual form only the columns related to the relevant symptoms, in which the answers of the patients to the various questionnaires or of the doctors were present in numerical form. Subsequently, the columns that did not contain any type of relevant information regarding symptoms such as numeric and text fields containing abbreviations, which may vary from table to table have been completely eliminated. Finally, a further skimming was carried out by eliminating all the tables that gave reliable information on the diagnostic status of the disease.

The process of correlating information was made sequentially as the visits progressed, adding the records of the next visit to previous ones. For each collection of documents taken into consideration, information is extracted for each individual patient. All the extracted information is kept in a new collection (Corpus). Subsequently, an initial cleaning of the text is performed. At first we made the tokenization of the text. Tokenizing a text means dividing the sequences of characters into minimal units of analysis called “tokens”, after which we have reported all the text in lower case and we have removed the non-textual tokens (i.e., operators, special symbols and numbers). We have also removed white space, terms with a length of less than three characters and stopwords. In particular, we performed stopword removal by also excluding words specific of the disease, such as “parkinsonian”, “parkinsonism” and “parkinson”, because they could be discriminatory terms for the classification.

4.2 The correlating information process

We used a stemming algorithm to transform the words in their root form, called “theme”. Then, we created the $n \times m$ document-terms matrix $A$, where a generic entry $A_{i,j}$ denotes the number of times that the $i_{th}$ term in the $j_{th}$ document appears. For the weight associated to each pair (term, document) we used the term frequency-inverse document frequency, also known as $tf - idf$ [11]:

$$tf_{idf}(A) = \log(tf(A)) \times idf(A).$$

In particular, for every term $t_i$ and document $d_j$ in $A$, $tf_{idf}$ is computed as follows:

$$tf_{idf}(A[t_i, d_j]) = \log(A[t_i, d_j] + 1) \times \ln \left( \frac{|d|}{\sum_{i,j} A[t_i, d_j] > 0} \right)$$

The $tf_{idf}$ is a function used in information retrieval to measure how important a word or a document is in a corpus [27]. The $tf$ value proportionally increases the number of times a word or document appears in the corpus. In our previous work [26] we adopted LSA for correlating information. In this paper we also explore the use of two Natural languages Processing techniques: Tex2Vec and Doc2Vec. The techniques used for correlating information are:

- **Tex2Vec**: Tex2Vec [31] is technique of text analysis and Natural Language Processing (NLP) by building machine learning algorithms based on text data which main goal is to provide an efficient framework with concise APIs for text analysis. It is built around
streaming APIs and iterators, which allows the construction of the corpus from iterable objects. This analysis allows us to build a matrix of document terms (DTM) and to elaborate the text by creating a map from words in a vector space. This technique is based on the concept of Word Embedding, a methodology of natural language processing to map words or phrases present in vocabulary, in a corresponding vector of real numbers, used to discover semantic correlations between them. To identify similar documents, we use cosine similarity identically. Also, in this case, a corpus of documents is built by selecting only those that coincide with the context in which the phenomena of interest reside, since even the inclusion of a large collection of high-quality documents could fail if the context of these documents does not align with the phenomenon of interest. Moreover, it is necessary to have a very large corpus to create a representative sample and to increase the chances of a word appearing in it. In this case, the stemming and the SVD are not applied. To represent documents in a vector space, we need to map terms with identifiers. In such a way as to represent a set of documents as a sparse matrix, where each row corresponds to a document and each column to a term. In our case, we have created a Document Term Matrix based on vocabulary. Doing nothing but cataloging the unique terms and assigning a unique ID. After, the similarity matrix is calculated using the DTM applying the cosine similarity. The comparison between documents with the cosine similarity also takes place. These are used to find which vectors are most similar to each other and which documents have a similarity greater than a specified threshold.

– **Doc2Vec**: Doc2Vec [20] is a technique that allows you to transform textual documents into vectorial representations that protect their semantics, trying to keep all possible information expressed in the text within the vectors, for example managing to interpret the information of similarity or thematic diversity between various text blocks. In reality, the Doc2Vec is an evolution of the Word2Vec technique, which consists of a group of models, used to do word embedding, whose purpose is to translate words or sentences into vectors of real numbers, or, in a form easily computable by compilers and which it manages to represent are not the word intended as a “sequence of characters” but also the meaning it assumes, thus managing to create a coding that allows, for example, to summarize concepts of similarity or opposition about other terms. These models are nothing more than two-level neural networks trained, through an unsupervised approach, to reconstruct the linguistic contexts of words; Word2Vec takes as input a large fragment of text and builds a vector space in which each word is uniquely assigned to a corresponding vector in space. The goal of Doc2Vec is to create a numeric representation of an entire document regardless of its length.

The purpose of Doc2Vec for its similarity to Word2Vec is to create a vectorial representation of an entire document regardless of its length, therefore, the vectors obtained will summarize the main theme or the global meaning of the entire document. It makes use of the Word2Vec model and in input, another vector is added, called DocumentID. So after training the neural network, you will have not only the word-vector (the vector representation of the words) but also a document vector (vector representation of the document). The purpose is simple, taken as input the DocumentID, the model uses the similarities between the words learned during the training (the word-vector) to build a vector that will include the words contained in it. By comparing these vectors, for example using the cosine similitude, we can then compare multiple documents with each other to verify their similarities. Doc2Vec, according to the Word2Vec approach used as
a base, is divided into two methodologies; in particular, we have the “Distributed Memory version of Paragraph Vector” (PV-DM) deriving from CBOW and the “Distributed Bag Of Words memory version of Paragraph Vector” (PV-DBOW) deriving from Skip-Gram. In particular, we based ourselves on the Word2Vec Skip-Gram, where the task of the neural network is to calculate, given an input word, the probability for each word of the vocabulary (together with all the words obtained from the training documents) to be close (juxtaposed within the text) to it. In reality, the concept of “closeness” between words is described through the definition of a measure, called windows size, which describes the number of terms to be analyzed, preceding or following the word given as input.

The network, therefore, for each input word, will have to find the probability that that specific word forms a pair with another word of the vocabulary; therefore, the net will be trained according to the number of times each pair is used. To allow training of the neural network it is necessary to provide a numerical representation of the words, as these, in the form of strings, cannot be easily used; for this reason, a vocabulary of the words obtained from the training documents is built and one-hot vectors will be used as input to the neural network; vectors of size equal to the size of the vocabulary, consisting of all negative bits except a positive one in correspondence with the reciprocal term in the vocabulary. The output of the neural network will also be a vector of the same size, as it will also use the indices to refer to the terms of the vocabulary, but, it will contain the probabilities of the various terms of being “close” to the word given as input.

4.3 The analysis of correlating information

After running the LSA, Text2Vec or Doc2Vec algorithms and obtaining the similarity matrix between the various documents, a clustering technique can be applied. In this work we use two types of techniques to compare categorization on the type of patients: the k-means [1] and the Fuzzy c-means clustering [15]. The main difference lies in the way in which the classification of the elements takes place:

- In the k-means technique the elements can belong only in mutual exclusion to a cluster and once assigned to a given cluster they can no longer be moved. The k-means algorithm is part of the “hard clustering” techniques and we exploited the kmeans function in the r package “cluster”.\(^2\) It is a partition clustering algorithm that allows to subdivide a set of objects in K groups based on their attributes, by partitioning the data set into unique homogeneous clusters whose observations are similar but different from other clusters. The k-means iteratively improves the initial centroids by minimizing the total intracluster variance, i.e., maximizing the similarity between the documents. The resulting clusters remain mutually exclusive.

- In the Fuzzy c-means technique the elements can belong simultaneously to both clusters, without any constraint. Fuzzy c-means clustering, also referred to as soft clustering or soft k-means, is used with the fanny function of the r package cluster, each element has a set of membership coefficients corresponding to the degree of the link with a given cluster; this value can vary from 0 to 1. The Fuzzy c-means algorithm is one of the most common fuzzy clustering algorithms, the centroid of a cluster is calculated as a weighted average of all points, based on the degree of cluster membership. The

\(^2\)https://cran.r-project.org/web/packages/cluster/cluster.pdf
clustering process is accomplished through an iterative optimization of the following function:

\[
nc \sum_{v=1}^{nc} \sum_{i=1}^{m} \sum_{j=1}^{m} u_{iv}^r u_{jv}^r d(e_i, e_j) / 2 \sum_{j=1}^{m} u_{jv}^r
\]

(2)

where \(e_i\) and \(e_j\) are pairs of entities selected in the set of all cluster entities. The size of this set is \(m\), while \(nc\) is the number of clusters to identify and \(u_{iv}\) is a non-negative value that specifies the membership of the entity \(e_i\) to the cluster \(v\). The sum of all the relevances of a given entity \(e_i\) is 1, while the exponent of membership is \(r\) and can assume values between 1 and \(\infty\). In the case \(r\) is close to 1, the behavior of the algorithm is similar to that of the k-means algorithm. The clustering process will stop when the inequality occurs [29]:

\[
max_{i,v=1...nc} |u_{iv}^{t+1} - u_{iv}^t| < \varepsilon
\]

(3)

\(t\) indicates the maximum number of iterations, while \(\varepsilon\) represents a termination criterion. The value for \(\varepsilon\) in \([0,1]\). Fuzzy c-means computes a membership matrix that is used to generate clusters. We empirically set \(r = 1.01, \varepsilon = 1e^{-20}\) and \(t = 1000\).

Both algorithms need the definition of \(K\), the number of clusters in which the information is divided. Because we classify the patient in two groups (genetically affected, genetically not affected) we set \(K = 2\). Moreover, as we have already said, in the Fuzzy c-means technique an element can belong to several clusters without restrictions and with different percentages of belonging. To overcome this problem, we have cleaned up all the spurious values, i.e., the observations that were less than \(1/K\), which were therefore removed and the results obtained are shown in Fig. 2.

5 The data analysis

To perform the comparison of the processes, as we have already said previously, the techniques were performed on the dataset divided by visits. Considering that the visits follow a precise chronological order (starting from the SC-screening visit up to the LOG-diagnostic visit) an incremental subdivision of the dataset has been chosen, therefore the subset of the dataset relating to a specific visit will contain the data of it plus those of previous visits. Furthermore, we have chosen to implement a function that subdivides the dataset into a partition in which each subset corresponds to a visit and contains only the information relating to it; this is to allow future analyzes that focus solely on the data provided by a visit or by groups of visits.

The results of the clustering algorithms are analyzed, by computing Precision, Recall and F-measure for each cluster. Precision and Recall are measures used to indicate accuracy and completeness of results, respectively, while F-measure represents a trade-off between these measures. All the measures are based on a comparison between an expected result and the result obtained. In particular, Precision measures (also called positive predictive value) the ratio between the correctly obtained instances (true positives) with respect to the total number of instances returned by the processing process (true positive and false positive). The Recall measures (also known as sensitivity) the ratio of instance correctly obtained (true
Fig. 2 Graphical representation of the clusters obtained with K-means (LSA a, Text2Vec c, Doc2Vec e and Fuzzy c-means (LSA b, Text2Vec d, Doc2Vec f) algorithms for each technique

positive) with respect to the number of expected instances (true positive and true negative) [13].

\[
Precision = \frac{|R \cap D|}{|D|}, \quad Recall = \frac{|R \cap D|}{|R|}
\] (4)

The F-measure is the harmonic mean of Precision and Recall and provides a measure of how the processing is effective.

\[
F - measure = 2 \cdot \frac{Precision \cdot Recall}{Precision + Recall}
\] (5)
where D is the list of patients returned by the classification and R is the list of patients correctly classified. The results of this analysis process are summarized in Figs. 3 and 4, where is highlighted that the best classification accuracy is reached with the fuzzy technique in each visit.

Specifically, as shown in Fig. 5, from the comparison of the results, it is shown that the techniques to which the k-means clustering is applied produce lower results than the Fuzzy clustering, except for the Doc2Vec in the SC, V02, V03 visits and the Text2Vec in the SC, BL, V01 visits where they produce better results than the other algorithms. Furthermore, despite the excellent performance of the Text2Vec k-means in the BL visit (F-Score 0.92), the technique proves to be somewhat unstable, in fact, we can see its total failure in the V04-V06 visits, where the k-means fails creating a cluster of a few dozen items. While the Text2Vec technique with Fuzzy clustering is the process that produces the best results in the series of visits from SC to V02 and from V10 to LOG. During the screening visit, both Doc2Vec (k-means) and Text2Vec (k-means and Fuzzy) produce very precise and similar classifications of patients.

Finally, we calculated the frequency of the different terms present in the documents and we have discarded the 95 quantile.

We have extracted the most relevant words from the documents related to the patient in the LOG visit, than we classified them in five categories:

- **Parental**: Patern and Sibling;
- **Symptomatic**: Pain, Sleep, Thyroid, Muscoloskeletal and Urinary;
- **Related disorders**: Hypothyroidism, Hypercholesterolemia, Diabet and Reflux;
- **Therapeutic**: Amantadin, Rytary, Arilict, Risagilin, Mirapex and Levodopa;
- **General terms**: Procedure and Full.

The words that are of most interest, come mostrato in Fig. 6 are Hypothyroidism, Hypercholesterolemia and Diabet. Regarding hypothyroidism, although no evidence of a higher frequency of hypothyroidism among patients with Parkinson’s disease has been reported in the literature, there may be a concomitance between these two diseases. In fact, studies conducted on patients with PD who take levodopa / carbidopa have indicated that the reduction of TSH levels is directly related to the drug, and occurs only during the first two hours after intake. It is not related to any significant thyroid dysfunction. This effect tends to be more discernible in males and is probably related to a primary or secondary propensity to hypothalamic levels specific for patients with PD [25]. Another study, however, has shown that the thyroid gland and its enzyme thyroperoxidase participate in the nitrosylation of serum proteins and can influence parkinsonian nitrosative stress and nitrosylation of serum alpha-synuclein, a potentially pathogenic factor [14]. Concerning hypercholesterolemia, many studies have not found a close correlation between the two pathologies, only a large prospective study [18] suggests that high total cholesterol at baseline is associated with an increased risk of Parkinson’s disease. As for diabetes, a large study [34] showed that diabetes was associated with a higher future risk of PD, because the insulin receptors are expressed in the substantia nigra. The dopamine agonist bromocriptine improves glycemic control and was approved for adjunctive treatment of diabetes. Conversely, the insulin sensitizer rosiglitazone protects dopaminergic neurons in animal models of PD. It is also important to point out that both diabetes and PD are age-related chronic diseases and some pathogenic processes may underlie both conditions.
Fig. 3  Clustering results for the K-means algorithm and Precision, Recall and F-measure data.
### LSA Fuzzy Clustering

| Visit | Cluster PD | Cluster GP | Results |
|-------|------------|-----------|---------|
| SC    | 856        | 1338      | PD 198  |
| BL    | 563        | 239       | 802 198 |
| V01   | 563        | 241       | 804 198 |
| V02   | 560        | 239       | 799 198 |
| V03   | 624        | 454       | 2078 198 |
| V04   | 626        | 448       | 1074 198 |
| V05   | 628        | 443       | 1071 198 |
| V06   | 621        | 432       | 1053 198 |
| V07   | 622        | 430       | 1052 198 |
| V08   | 624        | 433       | 1057 198 |
| V09   | 628        | 428       | 1085 198 |
| V10   | 634        | 447       | 1081 198 |
| V11   | 634        | 447       | 1081 198 |
| V12   | 635        | 450       | 1085 198 |
| V13   | 778        | 640       | 1413 284 |
| V14   | 509        | 313       | 824 614 |
| V15   | 221        | 93        | 318 894 |
| ST    | 221        | 93        | 318 894 |
| LOG   | 222        | 93        | 315 894 |

### TEXT2VECS Fuzzy Clustering

| Visit | Cluster PD | Cluster GP | Results |
|-------|------------|-----------|---------|
| SC    | 764        | 205       | 969 476 |
| BL    | 776        | 26        | 802 503 |
| V01   | 756        | 70        | 826 503 |
| V02   | 724        | 405       | 1179 522 |
| V03   | 726        | 405       | 1177 522 |
| V04   | 726        | 414       | 1190 513 |
| V05   | 777        | 421       | 1198 506 |
| V06   | 778        | 419       | 1197 508 |
| V07   | 777        | 414       | 1191 513 |
| V08   | 777        | 413       | 1190 514 |
| V09   | 778        | 409       | 1284 528 |
| V10   | 781        | 94        | 843 833 |
| V11   | 761        | 63        | 807 833 |
| V12   | 761        | 55        | 795 837 |
| V13   | 789        | 50        | 789 877 |
| V14   | 786        | 43        | 779 884 |
| V15   | 786        | 43        | 779 884 |
| ST    | 785        | 43        | 778 884 |
| LOG   | 785        | 44        | 779 884 |

### DOC2VECS Fuzzy Clustering

| Visit | Cluster PD | Cluster GP | Results |
|-------|------------|-----------|---------|
| SC    | 518        | 455       | 973 224 |
| BL    | 754        | 456       | 1210 232 |
| V01   | 735        | 426       | 1161 501 |
| V02   | 838        | 376       | 1214 551 |
| V03   | 821        | 364       | 1175 563 |
| V04   | 938        | 456       | 1387 571 |
| V05   | 938        | 456       | 1387 571 |
| V06   | 934        | 446       | 1380 492 |
| V07   | 933        | 438       | 1371 481 |
| V08   | 925        | 428       | 1353 499 |
| V09   | 929        | 419       | 1343 508 |
| V10   | 928        | 421       | 1344 507 |
| V11   | 921        | 422       | 1343 507 |
| V12   | 924        | 424       | 1348 507 |
| V13   | 925        | 423       | 1348 507 |
| V14   | 926        | 422       | 1348 507 |
| V15   | 928        | 421       | 1351 507 |
| ST    | 930        | 421       | 1351 507 |
| LOG   | 934        | 408       | 1342 519 |

Fig. 4  Clustering results for Fuzzy c-means algorithm and Precision, Recall and F-measure data
5.1 Discussion

From what has been observed we can conclude that the processes with LSA produce the worst results, and generally, the results produced by the execution of the processes via K-means are lower than those with Fuzzy clustering, this is due to the intrinsic structure of the K-means algorithm that does not allow an element that can be positioned in both clusters or that it can be moved later from one cluster to another, forcing an incorrect classification of the information processed. This is because in our data there is the PRODROMAL class, which are subjects not affected by Parkinson Disease but who present symptoms characteristic of this disease, therefore a stringent clustering could lead to a higher error rate, as can be seen from the results. In conclusion, the techniques that currently produce the best results are Text2Vec-Fuzzy, in visits from SC to V02 and from V10 to LOG, and Doc2VecM-Fuzzy in visits from V03 to V09.
The visualization of the information is very important in each field for a personalized visualization and a better understanding of the information but especially in the medical field. For example, it can act as a decision and diagnosis support for doctors. There are many tools for viewing information and data, here we use the D3.js framework\(^3\) that is a JavaScript library to create dynamic and interactive visualizations starting from organized data, visible through a common browser. for the visual rendering of the data, to be able to create both small tables, diagrams and statistics and complex graphic representations (including animations and other possibilities of interaction). Libraries are always linked to software that uses the functions of a programming library when a specific function of the collection is requested, which is why they only work within a program and cannot be performed independently.

In our case, we offer a visualization system, based on Radar chart, useful for the doctor to place each individual patient in one of the two clusters, in order to be able to make a quicker and faster diagnosis as the visits made in chronological order follow one another.

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\(^3\)https://d3js.org

Fig. 7 A data visualization based on Radar chart for fast diagnosis

6 Data visualization
For this visualization, we used the data obtained from both Text2Vec and Doc2Vec, based on the Fuzzy clustering efficiency obtained from the processing carried out for each visit. In fact, we used the Text2Vec for the SC, BL, V01, V02, V10-V15, ST and LOG visits and Doc2Vec for all the other visits, thus obtaining the chart shown in Fig. 7, for patient 3006. The chart is composed of three areas the green one indicates the non-sick patient, while the red one the sick patient; the orange line indicates the probability of patient 3006 to be included in the PD and GP areas. In the V02-V03 visits, the values indicate that the patient in question can be affected by Parkinson’s disease. Unfortunately, this is confirmed more and more in subsequent visits.

7 Conclusion

In this paper, we proposed a techniques to identify a correlation between the biomedical data in the PPMI dataset useful to verify the consistency of medical reports formulated during the visits and, then, to correctly classify the patients into affected or not. To correlate the information of each patient medical report, Information Retrieval techniques and clustering algorithms have been adopted. Furthermore, we have created a data visualization system to support diagnosis for doctors to be able to categorize the Prodromal class of patients in particular.

In the future, the analysis of the data and the correlation between them may be further extended, in particular, it could aim to establish a precise coding standard of the dataset, so that any future developments work with the updated datasets, in which the information of the tables are described with the same dictionary. In this way, the results of the various research activities can be compared more reliably. Furthermore, in the process of textual conversion of the numerical information contained in the dataset, these should be described with medical jargon; remember that we trained a model on specific medical texts on Parkinson’s, a correlation must be maintained between the two languages. Finally, we will evaluate the compression of the graphs proposed by doctors in order to validate the proposed technique.

Funding Open Access funding provided by Università degli Studi di Salerno within the CRUI-CARE Agreement.

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