An application of machine learning with feature selection to improve diagnosis and classification of neurodegenerative disorders

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Abstract

Background: The analysis of health and medical data is crucial for improving the diagnosis precision, treatments and prevention. In this field, machine learning techniques play a key role. However, the amount of health data acquired from digital machines has high dimensionality and not all data acquired from digital machines are relevant for a particular disease. Primary Progressive Aphasia (PPA) is a neurodegenerative syndrome including several specific diseases, and it is a good model to implement machine learning analyses. In this work, we applied five feature selection algorithms to identify the set of relevant features from 18F-fluorodeoxyglucose positron emission tomography images of the main areas affected by PPA from patient records. On the other hand, we carried out classification and clustering algorithms before and after the feature selection process to contrast both results with those obtained in a previous work. We aimed to find the best classifier and the more relevant features from the WEKA tool to propose further a framework for automatic help on diagnosis. Dataset contains data from 150 FDG-PET imaging studies of 91 patients with a clinic prognosis of PPA, which were examined twice, and 28 controls. Our method comprises six different stages: (i) feature extraction, (ii) expert knowledge supervision (iii) classification process, (iv) comparing classification results for feature selection, (v) clustering process after feature selection, and (vi) comparing clustering results with those obtained in a previous work.

Results: Experimental tests confirmed clustering results from a previous work. Although classification results for some algorithms are not decisive for reducing features precisely, Principal Components Analysys (PCA) results exhibited similar or even better performances when compared to those obtained with all features.

Conclusions: Although reducing the dimensionality does not mean a general improvement, the set of features is almost halved and results are better or quite similar. Finally, it is interesting how these results expose a finer grain classification of patients according to the neuroanatomy of their disease.

Keywords: Machine learning primary progressive aphasia, Supervised algorithm, Unsupervised algorithm, Clustering Analysis

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Background
Learning from data is one of the most successful fields applicable to many heterogeneous areas and disciplines like statistics, artificial intelligence, engineering, health, etc. Digital machines such as Magnetic Resonance Imaging (MRI), Mass Spectrometry (MS) or Positron Emission Tomography (PET), among others, and new generation sensors, have found their way in biomedical systems. In the recent decades, a plethora of advances in biotechnology have increased the diagnosis precision and the efficiency with tailored treatments. Furthermore, these machines generate wealth of data, which must be analyzed to extract the valuable knowledge that they contain. Data mining and machine learning techniques are crucial to explore, extract and interpret the generated data. Bioinformatics as a multidisciplinary area includes the application of computational tools to biological, medical or health data acquired. During the last decade, the amount of data has exponential growth, and the need to extract new knowledge for tackling diseases places bioinformatics as a priority research area. In this regard, machine learning and big data methods have been applied to better understand and fight many diseases.

The results of data analysis allow to identify patterns and develop predictive models to determine future behaviors in a particular context. In a biomedical scenario, machine learning techniques will provide an informed decision making, which will also have positive consequences for the health of patients. It is possible to implement a predictive model from a set of heterogeneous data acquired from diverse patients, which can be used to predict accurately the outcome for a new set of instances.

A major concern from a health and economic point of view is the impact of the aging population in the healthcare system, which is being studied and documented in all countries. Neurodegenerative diseases affect a large percentage of the old population and, to date, there is no cure for these diseases. Researchers focus on discovering new treatments for slowing down their progress and improve the quality of life of the patients. However, the success of clinical trials is hampered by a wide heterogeneity in clinical, genetic and pathological characteristics of neurodegenerative disorders.

Machine learning and data mining techniques can help to know the disease evolution and personalize treatments according to the needs and outcome of each patient. Personalized treatments rely on a proper diagnosis, where a specific subtype of the disease can be determined by subtle details in the imaging tools, and rule the prognosis. In this area, identifying the best classifier and clustering method will be the first step. However, taking a decision about the best classifier or clustering method to be used for reaching the most accurate results is a complex task. On the other hand, not all features are equally relevant in order to identify a disease or the stage of its evolution.

In this paper, we focus on Primary Progressive Aphasia (PPA). PPA is a clinical syndrome characterized by the neurodegeneration of language brain networks. Generally, PPA is the clinical onset of specific and frequent neurodegenerative diseases such as Alzheimer’s disease, tauopathies, or TDP-43 proteinopathies, which represent the main pathological subtypes of neurodegenerative disorders. Intriguingly, three clinical variants have been described in PPA (ie. nonfluent, semantic, and logopenic PPA), and each variant impairs certain brain regions and is more or less suggestive of a pathological subtype. In this regard, non-fluent or agrammatic variant is associated to left frontal lobe damage and is suggestive of tauopathies or less frequently TDP-43 proteinopathies; semantic variant is characterized by anterior temporal lobe impairment and is highly suggestive of TDP-43 type C pathology; and in logopenic variant, left parieto-temporal lobe is involved and it often precedes Alzheimer’s disease. This heterogeneity regarding the underlying pathology and different clinical courses in patients with a similar clinical presentation (ie. aphasia) makes PPA a good model among neurodegenerative disorders to apply the machine learning techniques. Indeed, we recently have introduced the possibility of the existence of some additional subtypes within non-fluent and logopenic variants, with a better prediction of outcome using clustering analysis.

Diagnosis of neurodegenerative disorders and, in particular PPA, has been improved with the use of advanced neuroimaging techniques. Among them, PET with 18F-fluorodeoxyglucose (FDG-PET) is a marker of synaptic dysfunction and reflects the brain topography of neurodegeneration with the measurement of brain metabolism. As a consequence, it is considered a useful, early and reliable tool in the diagnostic assessment of cognitive disorders and PPA, because each neurodegenerative disease trends to impair specific brain regions. However, the assessment of brain metabolism requires experience. Inter-rater agreement for visual analysis of PET imaging is usually moderate, but it could be improved with the use of some semiautomatic and statistical techniques.

The purpose of our work is to propose a machine-learning based framework suitable for the analysis of FDG-PET images of PPA patients, in order to solve two critical questions: i) to help on the automatic diagnosis of this disease; ii) to identify subtypes of the illness that have correlations with the anatomy of the damaged brain.

In [18], the authors applied Hierarchical clustering according to the clinical language deficit to classify patients. As a result, two groups were identified corresponding to nonfluent aphasia and semantic dementia. [19] studied the association between the PPA severity
and the disease duration, and disease duration and temporal atrophy on patients diagnosed of logopenic variant of primary progressive aphasia (lvPPA). They identified three variants of lvPPA using Hierarchical clustering on language, and neurocognitive test scores.

To perform this work, we used two data sets from 150 FDG-PET brain images, which comprised 116 brain attributes belonging to the normalized metabolism of brain regions and three more attributes, which are sex, age and clinical prognosis. Characteristics of participants and data acquisition methodology are explained as follows.

Participants
This study involved 150 FDG-PET imaging studies belonging to 91 patients with PPA (31 of them were examined twice in two different moments of the disease) and 28 healthy controls.

PPA is a rare neurodegenerative syndrome and thus, 91 patients in PPA is considered a large sample, even more considering that all patients were studied with a comprehensive protocol including FDG-PET. Regarding this, in the recent joint effort of the European Association of Nuclear Medicine and the European Academy of Neurology [20] to address the utility of FDG-PET in the assessment of primary progressive aphasia, four articles with PPA and FDG-PET were selected. The sample size of these works was in all cases < 55. Although PPA is a rare disease, we chose to study this disorder because it is a good model of brain-behaviour relationships, and it may be the onset of different clinical-pathological neurodegenerative entities.

PET imaging was performed in single-center between November 2011 and May 2017. All PPA patients met the current consensus criteria [21] and were classified into the three clinical variants taking into account the diagnostic criteria and follow-up. Further details about clinical and neuropsychological assessments are shown in a previous work [13].

The Institutional Research Ethics Committee from Hospital Clinico San Carlos approved the research protocol.

FDG-PET imaging acquisition and preprocessing
PET images were acquired following European guidelines [22]. All images were obtained in the same scanner, which is a Siemens Biograph True Point PET-CT that integrates a 6-detector CT with a late-generation PET using lutetium oxyorthosilicate crystals. A mean dose of 185 MBq was administered 30 minutes before the acquisition of the images and after at least 6 hours of fasting. Patients remained at sensory rest prior to the acquisition. CT scan parameters were: kVp/effective mAs/rotation: 130/40/1; slice thickness: 3 mm; reconstruction interval: 1.5 mm; and pitch: 0.75. Acquisition time was 10 min.

Statistical Parametric Mapping software was used for imaging preprocessing (http://www.filion.ucl.ac.uk). Images were realigned and normalized to the Montreal Neurological Institute standard space using a specifically developed brain FDG-PET template for dementia [23]. Global mean normalization was performed individually for intensity scaling. Marsbar software was used to perform a region of interest analysis, calculating the mean uptake value of each participant for the 116 brain areas of the Automatic Anatomical Labeling atlas belonging to the whole brain.

The rest of this paper is organized as follows. Experimental results are analyzed in “Results” section. Results discussion is presented in “Discussion” section and conclusions are drawn bellow. “Methods” section describes the methodology applied in this work.

Results
The software toolkit WEKA was used in our experimental approach. WEKA is available in http://www.cs.waikato.ac.nz/ml/weka as a complete industrial-strength software for machine learning. Analyzed data come from FDG-PET images of each obtained cluster and were compared to an additional control group of 32 healthy subjects. Previously to be entered in the statistical analysis, images had been spatially normalized and smoothed at 12 mm full-width at half maximum. Statistical Parametric Mapping version 8 was used for preprocessing and analysis. A two-sample T test was conducted to compare between groups, using age and gender as covariates. Statistical significance was set at $p < 0.05$ using family-wise error correction at cluster level.

For exploring the dataset, K-Fold Cross Validation was used as testing methodology. K-fold cross validation avoids overlapping by splitting data into $k$ subsets and makes $K$ iterations. For each iteration, a different subset was chosen for testing and the remainder for training. We picked $k = 10$ because this value is considered appropriate to obtain an accurate estimation.

Our first experimental work, which corresponds to the phase 1 in the flow chart in Fig. 1, compared a set of supervised learning algorithms in order to find the best classifier. In this step, we considered the fully set of brain attributes. Figure 2a shows the classification results obtained for each evaluated classifier. Figure 2b shows the True Positive rate and the Receiver Operating Characteristics metric (ROC) for each algorithm. ROC is an independent measure to evaluate the performance of a prediction model, that relates the True Positive Rate (TPR) against to the False Positive Rate (FPR). The optimal prediction model will have a ROC value of 1. Data were sorted by the number of instances correctly and incorrectly classified and the True Positive rate (TP rate), respectively. According
to these results, *SMO* algorithm presents the best performance with 127 instances correctly classified and 23 incorrectly classified (with respect to the clinical diagnosis criteria). *SMO* is followed by *IBK* algorithm with 114 instances correctly classified and 36 instances incorrectly classified.

Looking at *PART* and *Bayesian Naives* classifiers, *PART* obtained similar results to *IBK* with 113 and 37 instances correctly and incorrectly classified, respectively. *Naives* is the third best classifier, with 111 and 39 instances correctly and incorrectly classified, respectively.

Regarding the ROC metric, *SMO*, *BayesNet* and *Naives* presented the best performance with a ROC value of 0.9, followed closely by *Kstar* with 0.89, the closest values to 1, the optimal value. A good performance was obtained by *IBK* and *PART* with 0.83.

According to these results, we selected *SMO* as the best among those evaluated and *IBK* as the second one. However, as aforementioned, this approach used all the attributes for classification, while not all of them have the same weight neither provide relevant information for PPA diagnosis. Therefore, it is interesting to analyze which features are most important and provide more information for an accurate diagnosis. For that purpose, we developed a feature selection process by evaluating the algorithms presented in next section.

**Feature selection results**

The feature selection process, identified as the phase 2 in the flow chart, was launched and for each feature selection algorithm a new dataset was obtained that comprises only the features considered as most relevant by every algorithm. In the particular case of PCA, we tested different number of PCs in order to achieve accurate results. *ChiSquaredAttributeEval* and *ClassifierAttributeEval* generate a ranked dataset, where each feature is ranked by their merit value. We picked those with a merit greater than 0 for generating the new subset. The two other algorithms return directly a dataset with the subset of most relevant features according to the specific implementation.

Algorithms *CfsSubsetEval* and *WrapperSubsetEval* apply *Best First* as the search method and return the number of subsets evaluated and an internal metric used by both algorithms to choose the best subset. 3314 and 1258 subset were evaluated, until the best one was found, in the case of the *CfsSubsetEval* and *WrapperSubsetEval*, respectively.

At this step, we incorporate the expert knowledge and the subsets of features selected by the algorithms are presented to an experienced neurologist who is required to apply his criteria. This step is represented in the flow chart as the phase 3. The neurologist is able to evaluate the coherency of the features returned by each algorithm with respect to the neuroanatomy of the disease. Although this step can be easily automatized by a set of rules that categorize the brain regions that can be damaged by APP variants, we decided to present a clear a meaningful organization of selected features to the neurologist in order to acquire their knowledge without further noise.
In our case, the neurologist examined the results obtained for each algorithm. For the CfsSubsetEval algorithm, he considered the use of right angular as justifiable, however the algorithm immediately selected the cerebellum cruss and posterior cingulum, which is directly associated to logopenic variant. Moreover, cuneus is too posterior for three standard variants and next selected features get into the frontals. Hence he ruled out CfsSubsetEval as the most promising for the next phase.

With respect to WrapperSubsetEval, he considered that starting with the left calcarine is not a good option. This area is too posterior and it is not associated to language regions. Then, it follows in a right way, but it also uses the occipital area, which is also found very posterior and more linked to visual function. The expert dismissed the dataset resulting for this algorithm.

ChiSquaredAttributeEval and ClassifierAttributeEval were presented as two really reasonable proposals and the datasets could be used for the next phase. They have lots of similarities. However, ChiSquaredAttributeEval included many temporal areas before other regions were included. Moreover, the occipital regions were included before the frontal regions. Hence, it was finally dismissed.

The neurologist considered the features selected by the ClassifierAttributeEval algorithm, shown in Table 1, as the most suitable option. The main reason was the high ratio of attributes selected between the different lobes. Thus, the dataset resulting from this algorithm will be used in the following classification and clustering phases.

Principal Components (PC) as a result of PCA is a different matter. Each PC is an uncorrelated variable consisting on a number of probably correlated variables and it covers a part of the data variance. Table 2 shows the PCs with an eigenvalue higher than 1, placed in the second column, the third column corresponds to the proportion in the total, the cumulative value is displayed in the fourth column, and finally the features included. Features chosen belong to different PCs in order to launch the classification process again and analyse its results. Repeated features through different PCs were selected only once.

These results are analysed by the expert neurologist, which presented as the most relevant the principal components ranging between 1 to 8, and 10 to 15. These are chosen for the next phase.

### Classification after feature selection

Data subset obtained in the feature selection process and chosen by the neurologist has been as input to the classification process under the same conditions as the first experimental work. Figure 3a shows for the feature selection algorithm chosen in the previous phase, and classifiers addressed, the number of instances correctly and incorrectly classified. Figure 3b represents the evolution of the True Positive rate and the ROC metric for each feature selection algorithm and classifiers.

We analyzed these new results and compared them to those with all features. Regarding results after feature selection, the SMO algorithm is presented as the best classifier with 126 instances correctly classified, only one less than with all features. Naives ranks second behind SMO with 116 instances correctly classified, and Bayesnet, Kstar and IBK classified correctly 115 instances.

Comparing classification before and after feature selection, 6 out of 13 analyzed classifiers improved their results (IBK, Naives, BayesNet, Kstar, DecisionTable, LWL), 2 obtained identical results (DecisionStump and OneR), while results for 5 algorithms got worse (SMO, Part, J48 (prunned), J48 (unprunned) and RepTree). From the performance perspective of the classification model using the ROC metric, BayesNet improved with a value of 0.92, the best one, followed closely by SMO and Kstar with 0.9 and 0.89, respectively. PART and IBK presented good performance with 0.84 and 0.83, improved and sustained their

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| Feature | % Feature | % Feature | % Feature | % Feature | % Feature |
|---------|-----------|-----------|-----------|-----------|-----------|
| 0.144 TemporalPole Mid L | 0.087 Parahippocamp L | 0.055 Occipital Mid R | 0.031 Supramarginal R | 0.015 Paracentrallobule LM |
| 0.137 SMA L | 0.080 Occipital Mid L | 0.051 Frontal Sup Medial L | 0.031 Precuneus L | 0.015 Precentral R |
| 0.135 TemporalInf L | 0.077 CingulumPost L | 0.050 Cerebellum 8 L | 0.030 Cerebellum 9 L | 0.010 Rolandic Oper R |
| 0.133 Fusiform L | 0.077 TemporalSup L | 0.049 Frontal Sup R | 0.029 Insula R | 0.010 Cerebellum 8 R |
| 0.123 SMA R | 0.075 Cerebellum 6 R | 0.049 CingulumAnt R | 0.027 ParietalInf L | 0.010 Olfactory R |
| 0.117 Occipital Inf L | 0.072 Cuneus L | 0.047 Frontal Inf Tri L | 0.024 Insula L | 0.010 ParietalSup L |
| 0.110 CingulumAnt L | 0.067 Frontal Sup L | 0.041 TemporalMid R | 0.023 Occipital Sup L | 0.007 Paracentrallobule LM |
| 0.100 Angular R | 0.064 Vermis B | 0.037 Supramarginal L | 0.021 Frontal Mid R | 0.003 Poscentral L |
| 0.097 TemporalMid L | 0.059 Frontal Inf Oper L | 0.037 Hippocampus L | 0.019 Thalamus L | 0.001 Frontal Inf Orb L |
| 0.093 Angular L | 0.057 Precentral L | 0.037 TemporalInf R | 0.019 Heschl L | 0.000 Sexo |
Table 2 Principal components of PCA with eigenvalue higher than 1

| Principal components analyses | PC | EigenValue | Proportion | Cumulative | F1 | F2 | F3 | F4 | F5 |
|------------------------------|----|------------|------------|------------|----|----|----|----|----|
| 1                            | 29.385 | 0.249 | 0.249 | Frontal Mid Orb L | Frontal Inf Orb L | CingulumAnt L | Frontal Inf Tri L | Frontal Sup Orb L |
| 2                            | 19.402 | 0.164 | 0.413 | Occipital Mid L | Frontal Inf Orb R | TemporalMid L | Fusiform L | Occipital Inf L |
| 3                            | 13.378 | 0.113 | 0.527 | Precentral R | Frontal Mid R | ParietalInf R | Poscentral L | Poscentral R |
| 4                            | 7.585 | 0.064 | 0.591 | TemporalPole Sup R | Parahippocamp R | Hippocampus R | TemporalPole Mid R | Precentral L |
| 5                            | 6.411 | 0.054 | 0.645 | Thalamus R | Heschl L | Calcarine L | Heschl R | Cerebellum Crus1 L |
| 6                            | 4.807 | 0.041 | 0.686 | TemporalInf R | TemporalMid R | EdadPET | Rectus R | Frontal Sup Orb R |
| 7                            | 3.946 | 0.033 | 0.720 | Rolandic Oper L | Thalamus R | CingulumPost R | Poscentral L | TemporalPole Mid L |
| 8                            | 3.318 | 0.028 | 0.748 | Pallidum L | Thalamus L | Vermis 10 | Pallidum R | Putamen L |
| 9                            | 2.845 | 0.024 | 0.772 | Sexo | Rolandic Oper L | Rolandic Oper R | Heschl R | Paracentrallobule R |
| 10                           | 2.314 | 0.020 | 0.791 | CingulumPost R | CingulumPost L | Vermis 10 | Caudate R | Cerebellum 9 R |
| 11                           | 2.279 | 0.019 | 0.811 | Vermis 10 | SMA L | Cerebellum Crus1 R | Rectus R | Pallidum L |
| 12                           | 1.779 | 0.015 | 0.826 | Amygdala R | TemporalPole Mid R | Amygdala L | Sexo=2 | Vermis 9 |
| 13                           | 1.674 | 0.014 | 0.840 | Vermis 12 | ParietalSup L | Calcarine L | ParietalInf R | Cerebellum 3 L |
| 14                           | 1.460 | 0.012 | 0.852 | Cerebellum 10 R | Cerebellum 10 L | Amygdala L | Cerebellum 3 R | Amygdala R |
| 15                           | 1.156 | 0.010 | 0.862 | Olfactory R | Olfactory L | TemporalPole Sup R | TemporalPole Sup L | Sexo |
| 16                           | 1.122 | 0.010 | 0.872 | Putamen R | Putamen L | Cerebellum Crus2 R | Parahippocamp R | Frontal Inf Oper R |
| 17                           | 1.066 | 0.009 | 0.881 | Amygdala R | Cerebellum 10 R | Amygdala L | Cerebellum 10 L | TemporalInf R |

previous performance values. However, Naives obtained a poor value 0.68 compared to the previous one 0.9.

After this analysis, we cannot conclude that feature selection improves the number of instances correctly classified because the improvement percentage is not meaningful, although results are promising regarding the performance reached by some classifiers.

Regarding PCA results obtained in the classification process and shown in Fig. 4, it is difficult to generalize that PCA improves classification. However, experimental results demonstrated that some classification algorithms are able to improve or at least equal the number of instances correctly classified with a less number of features. Therefore, BayesNet obtains an improvement of 6 instances. Naives and Part equal previous results. Kstar classifies correctly 5 more than original tests, while LWL, DecisionStump and J48 improves previous results in 3 instances. This is an interesting result as we can achieve similar classification performance from a reduced number of features, hence simplifying the selection criteria by a set of rules that relate brain anatomy and PPA.

Finally, SMO, IBK, DecisionTable, OneR and RepTree obtain worse results. Moreover, these results were reached with less than half of the features 55 out 119. However, SMO obtained the best result with 120 instances correctly classified. Moreover, SMO, BayesNet, Naives obtained ROC values close to the optimal 1, 0.89, 0.9, 0.9, 0.9.

Fig. 2 Classification with all features. a represents the number of instances correctly and incorrectly classified for each evaluated algorithm. b graphs the True Positive rate and the ROC metric for each evaluated algorithm.
Fig. 3 Classification after feature selection except PCA. a represents the number of instances correctly and incorrectly classified for each classification algorithm and the set of features selected above. b graphs the True Positive Rate and the ROC metric for each classification algorithm and the set of features selected above.

results obtained after the feature selection process. Different colors in bars represent the three subtypes currently recommended in consensus PPA criteria. Dark blue is non-fluent/agrammatic, light blue represents semantic, green corresponds to logopenic and controls are represented by the yellow color. Particularly, instances assigned for each cluster are shown in Table 3.

After the feature selection phase, the expert neurologist evaluated four feature selection algorithms and he selected the ClassifierAttributeEval algorithm as the most suitable, hence the subset of features provided by this algorithm are considered as the most relevant. Therefore, we launched a new set of experiments to compare these results with those from the previous study.

We analyzed the Linkage clustering algorithm, and compared results before and after feature selection. Clustering results after the feature selection process for the Linkage clustering algorithm are outlined in Fig. 5b and d, where X-axis represents the number of clusters evaluated and Y-axis corresponds to the number of patients assigned to each cluster. The clinical PPA diagnosis for each patient is represented by a different color within the cluster. We provide a more detailed information of the distribution of patients per cluster in Table 3 b and d.

respectively. Kstar obtained good performance with 0.88, PART improved with 0.84 and IBK got a little worse from 0.83 to 0.82.

Nevertheless and once the feature selection algorithm was chosen and the best set of features were selected from PCA, we explored in the following section unsupervised learning process to compare with previous experiments carried out in [13].

Clustering results after feature selection
In a previous work [13], we applied Agglomerative Hierarchical Clustering (AHC) as unsupervised learning algorithm [24], particularly the Ward Linkage [25] algorithm, to evaluate how patients were split into subgroups taking into account clusters from 4 to 8. According to the experimental results, the distribution of patients in 8 clusters led 6 different subtypes of PPA, what reinforces the latest clinical studies [26–28]. These experiments were carried out considering all attributes for classification. Figure 5 shows the distribution of patients within each cluster for 4 and 8 cluster, before and after the feature selection process. The X-axis is the number of clusters and Y-axis the number of patients belonged to each cluster. Figure 5a and c represent results obtained in the previous work, without feature selection. Figure 5b and d correspond to
We compared the classification results for 4 and 8 clusters after feature selection with those obtained in our previous study. Regarding 4 clusters, the group k1 contains in both cases all no-fluent and semantic PPA patients, either after the classification performed with all attributes, and after feature selection. However, a logopenic patient is moved to k3 group, and two patients previously belonging to the healthy control are moved to k4 and k2, respectively.

With respect to 8 clusters, the first group k1 remains unchanged and it contains the same patients than k1 group with all attributes. It comprises 16 patients with non-fluent and 1 patient with logopenic variants. K2 group includes same patients as the former k2 group

| Table 3 | Distribution of patients per cluster and clinical PPA diagnosis for Linkage and 4 and 8 cluster |
|---------|------------------------------------------------------------------------------------------------|
| 4 Clusters |                                                                                                   |
|          | Before Feature Selection | After Feature Selection |
| PPA      | K1  | K2  | K3  | K4  | K1  | K2  | K3  | K4  |
| 1        | 46  | 0   | 0   | 0   | 46  | 0   | 0   | 0   |
| 2        | 15  | 0   | 0   | 0   | 15  | 0   | 0   | 0   |
| 3        | 2   | 37  | 22  | 0   | 1   | 37  | 23  | 0   |
| HC       | 2   | 0   | 0   | 26  | 0   | 1   | 0   | 27  |
|          | a)  |                                              | b)                                  |
| 8 Clusters |                                                                                                   |
|          | Before Feature selection | After Feature selection |
| PPA      | K1  | K2  | K3  | K4  | K5  | K6  | K7  | K8  | K1  | K2  | K3  | K4  | K5  | K6  | K7  | K8  |
| 1        | 16  | 0   | 24  | 0   | 0   | 6   | 0   | 0   | 16  | 0   | 10  | 20  | 0   | 0   | 0   | 0   |
| 2        | 0   | 0   | 0   | 15  | 0   | 0   | 0   | 0   | 0   | 0   | 0   | 15  | 0   | 0   | 0   | 0   |
| 3        | 1   | 37  | 0   | 0   | 22  | 1   | 0   | 0   | 1   | 37  | 0   | 0   | 0   | 23  | 0   | 0   |
| HC       | 0   | 0   | 2   | 0   | 0   | 14  | 12  | 0   | 0   | 1   | 0   | 0   | 0   | 14  | 13  | 0   |
|          | c)  |                                              | d)                                  |

Parts labeled as a and c correspond to results before the feature selection process, whereas b and d represent results with the set features selected above. Number of patients assigned to each cluster are shown, and their previous clinical PPA diagnosis is represented by the column labeled PPA. Values 1, 2 and 3 in the first column correspond to nonfluent/agrammatic, semantic and logopenic PPA, respectively, whereas HC represents healthy controls. Kn specifies the number of clusters, where \( n = 0, 1, \ldots, N \) and \( N \) is a value between 4 and 8.
and one healthy control. This healthy control is the only difference between both results.

The former k3 group included 24 patients with nonfluent and 2 controls. After feature selection, k3 is subdivided into two subgroups k3 (n=10) and k4 (n=20).

Patients diagnosed with semantic PPA are identified in both experiments in a single group, k4 in the former study and k5 after feature selection. The k5 group after feature selection includes all patients, which were in the k5 group in the previous work. Moreover, one patient is moved to k5 when it belonged to the former k6. Finally, healthy controls were also subdivided in two subgroups k7 and k8. K7 comprises 14 patients all of then women, and k8 13 men patients. The only difference is given in the k8 group with a new patient moved from k3 in the previous study, where patients were mostly diagnosed with logopenic variants. As we previously mentioned, the subdivision of healthy controls in two groups probably reflects gender differences in regional brain metabolism, because 100% of cases in k7 and k8 were women and men, respectively [29].

Our results expose a finer grain classification of patients according to the neuroanatomy of their disease, and give a step further in the field of PPA diagnosis. However, with respect to the previous work the clustering analysis after feature selection had not profound improvements and we addressed the Principal Component Analysis (PCA)[30].

Clustering after the principal components analysis

After PCs with higher coverage of the variance were chosen, we launched the clustering analyses with those that reached better results in the classification process. Moreover, we tried to cover as many possibilities as possible, thus we tested different sets of PCs to indentify those features which better fit to the problem at hand. Results were compared with those from our previous study. Figure 6 displays the distribution of patients within each cluster, where X-axis represents the number of cluster and the Y-axis the number of patients assigned to each one. Colors corresponds to the clinical PPA diagnosis for patients within the cluster (1 nonfluent/agrammatic (dark blue), 2 semantic (light blue), 3 logopenic (green)) and healthy controls (yellow). More detailed information for 4 clusters is given at Table 4. Results from the study previously mentioned, labeled as (a), are detailed on the left (highlighted in bold). Next column, labeled as (b), represents the distribution of instances for the PCs selected. Clustering results for 4 and 8 clusters are drawn in Figures (a) and (b), and (c) and (d), respectively.

Regarding the column labeled as (b), which corresponds to experimental tests 1-8, 10-15 and 1-14 No Sex, we observed that all except one instances are identified according to the standard PPA types. The group k1 contains all patients diagnosis as no-fluent and one instance with a prognosis as logopenic PPA. The remaining groups correspond to patients diagnosed with semantic and logopenic PPA, and controls, respectively. These results demonstrated that with a less number of features obtained from PCA, the most of instances are almost perfectly assigned according to the clinical diagnosis.
Table 4 Distribution of patients per cluster and clinical PPA diagnosis for Linkage and 4 and 8 clusters

|                | Previous Study (a) | 1-8,10-15 PCs(b) |
|----------------|-------------------|------------------|
|                | K1    | K2    | K3    | K4    | K1    | K2    | K3    | K4    | K1    | K2    | K3    | K4    |
| PPA 1          | 46    | 0     | 0     | 0     | 46    | 0     | 0     | 0     | 1     | 60    | 0     | 0     |
| 2              | 15    | 0     | 0     | 0     | 0     | 0     | 15    | 0     | 0     | 0     | 0     | 0     |
| 3              | 2     | 37    | 22    | 0     | 1     | 60    | 0     | 0     | 0     | 0     | 0     | 0     |
| HC             | 2     | 0     | 0     | 26    | 0     | 0     | 0     | 28    | 0     | 0     | 0     | 0     |

Parts labeled as a and c correspond to results before the feature selection process, whereas b and d represent results with the set features selected above. Number of patients assigned to each cluster are shown, and their previous clinical PPA diagnosis is represented by the column labeled PPA. Values 1, 2 and 3 in the first column correspond to nonfluent/agrammatic, semantic and logopenic PPA, respectively, whereas HC represents healthy controls. Kn specifies the number of clusters, where $n = 0, 1, \ldots, N$ and $N = 4$.

We focus on the clustering analyses for 8 clusters. Figure 6 shows the distribution of patients within clusters. We extended information in Table 4(b) and we compared results of the columns namely as (b) with the column labeled as (a).

Below is the analysis for 1-8,10-15 PCs (b). In this case the group k6 and k4 in (a) are identical to k1 and k6, respectively in (b). 6 instances from k5 and 12 from k2 in (a) are placed in the group k5 in (b). K1 and k3 in (a) distributed its patients between k3 and k4 in (b). Healthy controls were assigned to the only group k7 in (b).

Discussion

Our study addresses the improvement of the PPA diagnosis from FDG-PET images applying machine learning techniques. Using results obtained in a previous work as a reference, we extend the study to identify the best classifier algorithm and the best set of features to simplify the PPA diagnosis. According to the classification results obtained with all features in phase 1, SMO followed by IBK are the best algorithms among those evaluated. Although twelve classifiers algorithms were selected, other algorithms can be evaluated in order to obtain the most precise results. Phase 2 was in charge to analyse the performance of four feature selection algorithms and PCA to identify the best set of features. This phase also considered the expert neurologist supervision and compared the results with those obtained in the phase 1. After this analysis, we cannot conclude that feature selection improves the number of instances correctly classified because the improvement percentage is not meaningful, although results are promising regarding the performance reached by some classifiers. Analysing a wider set of feature selection algorithms should allow not only to confirm these results but also reassert that other alternatives have to be explored.

According to the classification results obtained with the features extracted with PCA, we cannot generalize that PCA improves classification. There is no doubt that some classification algorithms equal or even improve the number of instances correctly classified with a less number of features.

Once the ClassifierAttributeEval algorithm selected the best set of features, we explored unsupervised learning with Hierarchical Clustering to compare with previous experiments carried out in [13]. Considering the set of features provided by the ClassifierAttributeEval algorithm, our results adjust the classification of patients according to the neuroanatomy of their disease, as we previously mentioned. Regarding the set of features from PCA, our results reinforce our previous outcomes with a less number of features. Most of the instances are almost perfectly assigned according to the clinical diagnosis. However, other machine learning branches can help to identify the best reduced set of relevant features and make easier the automation of the diagnosis of PPA.

Conclusions

In this work, we have exploited machine learning techniques to improve the PPA diagnosis from FDG-PET images. This study confirms and reinforces the results obtained in a previous clinical work, where we explored the automatic classification of PPA patients and found out new subtypes of this disease that correlate with the clinical findings and better predict the clinical course. Here, we have proposed a machine learning approach that, on one hand, validate the clinical findings and, on the other
Hand, offers a practical and automatic tool to help on the
diagnosis of PPA variants in order to improve the manage-
ment of these patients. The data acquired from FDG-PET
images are characterized by a high dimensionality, hence
the application of the feature selection process is consid-
ered as a key proposal to reduce the dimensionality in
order to improve the automatic classification. As part of
the feature selection step, the expert knowledge has been
included in a simplified and natural way for the clinical
assessing, and could be automatized in the future by a
set of rules. As a result, the number of patients correctly
classified is increased after the most relevant features are
identified, particularly after PCA analyses. As a conclu-
sion, a fine grain classification has been obtained based
on the neuroanatomy of the disease, which improves pre-
vious results for the PPA diagnosis. Overall, our study
suggests the role of machine learning techniques applied
to neuroimaging analysis in order to improve the classifi-
cation of neurodegenerative diseases, and closes the gap
between image processing and automatic diagnosis tools
that help on the clinical practice.

Methods
As aforementioned, our goal is to advance on the devel-
opment of an automatic diagnosis tool based on machine
learning approaches. For that purpose, we aim to improve
the classification process carried out in [13] that brought
the finding of new PPA variants. In the previously men-
tioned work, we used Hierarchical clustering as unsu-
upervised learning algorithm [24], particularly the Ward
Linkage [25] algorithm with all attributes present in the
dataset. We concluded that unsupervised clustering anal-
ysis of FDG-PET data favored, based on the Davies-
Bouldin index, the classification of PPA into six variants
rather than three subtypes as currently recommended in
consensus PPA criteria. However, input data are multidim-
ensional; we have 150 instances and 119 attributes for
each one and some attributes may introduce noise, which
affects the classification process.

In this light, we considered necessary to identify the
most relevant attributes, reducing the multidimensional-
ity and improving the classification process, accurately.
Figure 1 shows the flow chart proposed as methodology,
which will be detailed below.

The first phase, identified as 1 in the flow chart,
provides the dataset with all features to the classifi-
cation phase, identified as 4, where for each classifier
algorithm the process is launched. The set of selected
classifiers was BayesNet, NaiveBayes, SMO, IBK, Kstar,
LWL, DecisionTable, OneR, PART, J48, DecisionStump
and RepTree from the machine learning software WEKA
(Waikato Environment for Knowledge Analysis) [31] in
order to evaluate which classifiers perform better for
the problem of improving the diagnosis of PPA from
FDG-PET images. As a result, we obtained the classifi-
cation with all features. Phase labeled as 2 is in charge
of the feature selection phase and consists in pinpointing
those attributes considered more relevant to the prob-
lem at hand (this is to improve the classification of
patients according to more accurate subtypes of PPA).
For this phase, we selected four typical feature selec-
tion algorithms: CfsSubsetEval, ChiSquaredAttributeEval,
ClassifierAttributeEval and WrapperAttributeEval, and we
applied the Principal Components Analysis (PCA) algo-
rum [30], although they are not comparable. PCA is a
powerful mathematical algorithm for analysing data and
reducing high dimensionality, preserving its variability as
much as possible, and at the same time improving the data
interpretability. PCA explores a dataset represented by a
matrix $X$, where each row is one entity $n$ and each column
identifies a numerical variable $p$. The algorithm search
the linear combination of the columns, which maximizes
the variance. This algorithm is able to identify patterns in data
and discover their differences and similarities to find the
more relevant features. We applied PCA on the dataset
previously mentioned. When the process is completed, a
set of principal components is obtained. Next step, we
decided to ignore the principal components of lesser sig-
nificance in order to preserve as much information as
possible.

Datasets obtained from the selected algorithms contain
the most relevant features according to each algorithm.
These datasets are used as input in the following phases,
classification and clustering phases, labeled as 4 and 6,
respectively.

The next phase corresponds to the clustering phase,
identified as 6, where clustering analysis [32] is performed.
In this phase, Hierarchical clustering [24] parameterized
to Ward’s Linkage is launched. Instances are classified in
different groups. The result of this process is compared
against the clusters found in our previous work, 8 clus-
ters, and also against the clusters described in the classical
medical literature, 4 clusters.

Abbreviations
AHC: Agglomerative hierarchical clustering; FDG-PET: Fluorodeoxyglucose
positron emission tomography; FPR: False positive rate; IBK: Instance based
learner K-nearest; KStar: Search algorithm K*; LVPPA: Logopenic variant PPA;
LWL: Locally weighted learning; MRI: Magnetic resonance imaging; MS: Mass
spectrometry; PART: Partial decision trees; PCA: Principal components; PCA:
Principal components analysis; PET: Positron emission tomography; PPA:
Primary progressive aphasia; ROC: Receiver operating characteristics; SMO:
Sequential minimal optimization; TDP-43: TAR-DNA binding protein-43; TPR:
True positive rate

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Authors’ contributions
JD contributed to carry out each experimental test, compile and prepare the
algorithms results, results analyses and manuscript writing. JMG contributed
to compile and prepare data from patients, as the main supervisor of clinical
results, results analyses and manuscript writing. MNCM contributed to
compile and prepare data from patients. JLJM contributed to the supervision
of the algorithms addressed, results analyses and manuscript writing. JLA as the principal supervisor and contributed to all phases and manuscript writing. All authors read and approved the final manuscript.

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Availability of data and materials
All data are available in a systematic Data Base created by the Department of Neurology of the San Carlos Hospital, in Madrid, and accessible to clinicians and researchers participating in the project. These data are not publicly available due to data privacy laws.

Ethics approval and consent to participate
The Institutional Research Ethics Committee from Hospital Clínico San Carlos approved the research protocol with the 1964 Helsinki declaration and its later amendments. Written informed consent was obtained from all individual participants included in the study or their caregivers.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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References
1. Alpaydin E. Introduction to Machine Learning, 2nd Edition. Cambridge, MA: MIT Press; 2010.
2. NCBN. Available at the National Center for Biotechnology Information’s Web Site. Bethesda, 2018. www.ncbi.nlm.nih.gov.
3. Khemphila A, Boonying V. Comparing performances of logistic regression, decision trees, and neural networks for classifying heart disease patients. In: 2010 International Conference on Computer Information Systems and Industrial Management Applications (CISIM). Washington DC: IEEE Computer Society Press; 2010. p. 193–8. https://doi.org/10.1109/CISIM.2010.5643666.
4. Marco J, et al. Data mining methods in the prediction of dementia: A real-data comparison of the accuracy, sensitivity and specificity of linear discriminant analysis, logistic regression, neural networks, support vector machines, classification trees and random forests. BMC Res Notes. 2011;4(112).299. https://doi.org/10.1186/1756-0500-4-299.
5. Kavakiots I, et al. Machine learning and data mining methods in diabetes research. Comput Struct Biotechnol J. 2017;15:104–16. https://doi.org/10.1016/j.csbj.2016.12.005.
6. Kouroo K, et al. Machine learning analysis in cancer prognosis and prediction. Comput Struct Biotechnol J. 2015;13:8–17. https://doi.org/10.1016/j.csbj.2014.11.003.
7. Husain M. Alzheimer’s disease: time to focus on the brain, not just molecules. Brain. 2017;140(2):251–3.
8. Mesulam MM, et al. Primary progressive aphasia and the evolving neurology of the language network. Nat Rev Neurol. 2014;10(10):534–69.
9. Matias-Guiu JA, García-Ramos R. Primary progressive aphasia: From syndrome to disease. Neurología. 2013;28(6):366–74.
10. Marshall CR, et al. Primary progressive aphasia: a clinical approach. J Neurol. 2018;256:1474–90.
11. Matias-Guiu JA, et al. Visual and statistical analysis of 18F-FDG-PET in primary progressive aphasia. Eur J Nucl Med Mol Imaging. 2015;42(6):916–27.
12. Matias-Guiu JA, et al. Clinical course of primary progressive aphasia: clinical and FDG-PET patterns. J Neurol. 2015;262(5):570–7.
13. Matias-Guiu JA, et al. Clustering analysis of FDG-PET imaging in primary progressive aphasia. Frontiers in Aging Neuroscience. 2018;10:1–22. https://doi.org/10.3389/fnagi.2018.00230.
14. Shimizu S, et al. Role of Neuromaging as a Biomarker for Neurodegenerative Diseases. Front Neurol. 2018;9:265.
15. Lafort R, et al. Molecular imaging in dementia. Past, present, and future. Alzheimers Dement. 2018;14:1522–52.
16. Matias-Guiu JA, et al. Evaluation of the new consensus criteria for the diagnosis of primary progressive aphasia using fluorodeoxyglucose positron emission tomography. Dement Geriatr Cogn Disord. 2014;38(3-4):147–52.
17. Perani D, et al. Validation of an optimized SPM procedure for FDG-PET in dementia diagnosis in a clinical setting. Neuroimage Clin. 2014;4:445–54.
18. Knibb JA, Xuereb JH, Patterson K, Hodges JR. Clinical and pathological characterization of progressive aphasia. Ann Neurol. 2006;59(1):156–65. https://doi.org/10.1002/ana.20700. https://onlinelibrary.wiley.com/doi/pdf/10.1002/ana.20700.
19. Mutchulla MM, Whitford JL, Duffy JR, Strand EA, Dean PM, Serrym ML, Jack CR, Josephs KA. Identification of an atypical variant of logopenic progressive aphasia. Brain Lang. 2013;127(2):139–44. https://doi.org/10.1016/j.bandl.2013.02.007.
20. Bouwman F, Olin S, Gandolfo F, Altmare D, Fести C, Agosta F, Arbizu J, Drzezga A, Nester P, Nobili F, Walker Z, Morbelli B, Boccardi M, for the EANM-EAN Task Force for the Prescription of FDG-PET for Dementing Neurodegenerative Disorders. Diagnostic utility of fdg-pet in the differential diagnosis between different forms of primary progressive aphasia. Eur J Nucl Med Mol Imaging. 2018;45(9):1526–33. https://doi.org/10.1007/s00259-018-4034-2.
21. Gorno-Tempini, et al. Classification of primary progressive aphasia and its variants. Neurology. 2011;76(11):1006–14. https://doi.org/10.1212/WNL.0b013e31821103e6. http://neurology.nationaljournal.com/content/76/11/1006.full.pdf.
22. Varrone A, et al. EANM procedure guidelines for PET brain imaging using [18F]FDG, version 2. Eur J Nucl Med Mol Imaging. 2009;36(12):2103–10.
23. Della Rosa PA, et al. A standardized [18F]-FDG-PET template for spatial normalization in statistical parametric mapping of dementia. Neuroinformatics. 2014;12(4):575–93.
24. Evenett BS, et al. Hierarchical Clustering, in Cluster Analysis, 5th Edition. Chichester: John Wiley & Sons, Ltd; 2011. https://doi.org/10.1002/9780470977811.ch4.
25. Jr. JHW. Hierarchical grouping to optimize an objective function. J Am Stat Assoc. 1963;58(301):236–44. https://doi.org/10.1080/01621459.1963.10500845. http://www.tandfonline.com/doi/pdf/10.1080/01621459.1963.10500845.
26. Josepka A, et al. Characterizing a neurodegenerative syndrome: primary progressive apraxia of speech. Brain. 2012;135(Pt 5):1522–36.
27. Mesulam MM, Weintraub S. Is it time to revisit the classification guidelines for primary progressive aphasia? Neurology. 2014;82(13):1108–9.
28. Botha H, et al. Classification and clinicoradiologic features of primary progressive aphasia (PPA) and apraxia of speech. Cortex. 2015;69:220–36.
29. Hu Y, et al. Gender differences of brain glucose metabolic networks revealed by FDG-PET: evidence from a large cohort of 400 young adults. PLoS ONE. 2013;8(12):83821.
30. Jolliffe IT. Principal component analysis. In: International Encyclopedia of Statistical Science. Berlin, Heidelberg: Springer Berlin Heidelberg. 2011. p. 1094–6. https://doi.org/10.1007/978-3-642-04989-8_245.
31. Frank E, et al. In: Maimon O, Rokach L, editors. Weka-AMachine Learning Workbench for Data Mining. Boston: Springer; 2010. pp. 1269–77. https://doi.org/10.1007/978-0-387-09823-4-66.
32. Hall/CRC H. Handbook of Modern Statistical Methods. 6000 Broken Sound Pkwy NW Ste 300: CRC Press; 2016.