# Università degli studi di Milano-Bicocca

# ADVANCED MACHINE LEARNING FINAL PROJECT

# Deep Learning-Enabled Decoding of Raman Spectroscopy

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#### Abstract

Raman spectroscopy is commonly used in chemistry to provide a structural fingerprint by which molecules can be identified. Machine Learning (ML) methods are used to decode the resulting spectra. The purpose of this study is to identify patients affected by Amyotrophic Lateral Sclerosis (ALS) using ML methods. The main problem for an efficient classification is the data scarsity. Indeed, Raman spectra are subject to medical confidentiality. Regarding the limited dataset, a model pre-trained on a larger one, composed of bacterial spectra, is a good solution to obtain better performances. This report presents a Transfer Learning (TL) approach combined with data augmentation techniques.

#### 1 Introduction

Raman spectroscopy allied with ML methods promises a new, rapid and non-invasive technique to diagnose patients. This ability has led researchers to highlight that Convolutional Neural Networks (CNNs) outperform other ML methods. However, CNNs are "data-hungry", they require a massive amount of data in order to extract the right features. Usually, a CNN can be well-trained on a few thousand samples.

Unfortunately, the dataset is composed of less than 600 spectra, which is not really sufficient to properly train a model. In addition, a problem lies in the repartition of the number of samples per patient and per group. Transfer Learning aims at improving the performance on a target domain by transferring the knowledge contained in different but related source domains.

In this work, the effort is focused on trying different TL experiments. The first is fine-tuning which consists of unfreezing a pre-trained models and re-training it on the new data. The second experiment is the most common incarnation of transfer learning: features extraction. After taking the layers from a pre-trained model, the layers are freezed and some new trainable layers are added on top of the frozen ones. These new layers will learn to turn the old features into predictions on a new dataset.

Finally some data augmentation methods will be applied. It is a well-known technique for improving the training of neural networks. The idea is to expand the number of training samples by adding some noise and small variations resulting in a more robust training. For spectral data, random offsets, random multiplications and Gaussian noises are commonly used.

#### 2 Datasets

The dataset provided is composed of 591 spectra, belonging to 30 patients, divided into two classes: patients affected by ALS and healthy ones (CTRL). 393 spectra are provided for the ALS patients and 198 for the CTRL ones as it is shown in table 1. Thus, it is quite unbalanced. Furthermore, notice that the spectra of a same patient are correlated, so it is necessary to treat them together.

Thanks to Dario Bertazioli who provided the data, they are filtered and pre-processed. The only addition is the negative values removal from the spectra. In fact, these points are meaningless, having negative value would literally mean that energy is produced by scattering.

The bacteria-ID dataset, on which the models used for TL has been pretrained, consists of 30 bacterial and yeast isolates. 2000 spectra are associated for each of the 30 reference isolates.

# 3 The Methodological Approach

The first core idea of this project was the comprehension of the dataset and its limits.

At the beginning, 4 classical ML classifiers have been implemented:

- Decision Tree Classifer
- Logistic Regression
- SVM
- Random Forest Classifier

If the correlation of spectral data among the same patient is ignored, the models are overfitted which is caused by a data leakage. Therefore, to properly have an unbiased diagnostic method, there is a need to carefully split the training and test data. It is mandatory that spectra belonging to the same patients don't show up both in the training set and in the test set. To maximally exploit the available training data, a Leave-One-Patient-Out Cross Validation and a 8 GroupKFold on patients have been implemented.

It is worth mentionning that some feature preprocessing has also been done with Principal Component Analysis (PCA) which seems to increase the

Table 1: This table represents the repartition of patients and spectra belonging to the ALS and CTRL groups

Repartition of spectra and patients in dataset					
Groups	Patient ID	Samples range	Samples count	Number of samples per group	
ALS	ALS01	1-60	60		
	ALS02	61-78	18		
	ALS05	79-114	36		
	ALS07	115-150	36		
	ALS08	151 -194	44		
	ALS09	195-210	16		
	ALS10	211-225	15	393	
	ALS11	226-242	16*		
	ALS12	243-256	14		
	ALS13	257-281	25		
	ALS14	282-300	19		
	ALS15	301-314	14		
	ALS16	315-324	10		
	ALS17	325-334	10		
	ALS18	335-344	10		
	ALS20	345-354	10		
	ALS22	355-364	10		
	ALS23	365-374	10		
	ALS24	375-384	10		
	ALS25	385-394	10		
CTRL	CTRL01	1-33	33		
	CTRL02	34-76	43		
	CTRL03	77-91	15		
	CTRL04	92-138	47		
	CTRL05	139-149	11	198	
	CTRL06	150-158	9	130	
	CTRL07	159-168	10		
	CTRL08	169-178	10		
	CTRL09	179-188	10		
	CTRL10	189-198	10		

 $<sup>^{*}</sup>$  The 227th sample is missing. It has probably been removed while processing because it was "too bad".

performance of the models (except SVM because the kernelized machine is already doing a similar work).

After setting up these baselines, the second and central step was the discovery and the application of the methods used in the bacteria-ID paper [1]. In this respect, some basic predictions have been firstly computed on the raw dataset using the three pre-trained models (CPKT extension):

- pretrained model
- finetuned model
- clinical pretrained model

There are saved parameters for the pre-trained CNN which is based on the ResNet architecture and dynamically modified to fit the input and output dimensions.

The fine-tuning experiments resulted from the observation that none of the model obtain good performances. It is due to the difference between the source and target domains of the training set.

Given the custom splitting method presented in the bacteria-ID paper[1] the patients are divided into sub-groups of 5 patients each. The samples of the first three are used for the training set, the fourth for the validation set and the last one for the test set. Each of the samples selected are then shuffled to make sure that no implicit link may be found by the models. Some of these spectra are selected for finetuning the models and the remaining ones for predicting on the finetuned models. However as there is a clear lack of data it is not possible to apply this method (some of the patients have only 9 samples!). Therefore, a more convenient method has been developped. First, the data is divided into:

- a "finetunable" dataset for finetuning the pretrained models. It is composed of 20 patients.
- a "full test" set which is used for computing the performances of the finetuned models. It is composed of 10 patients.

As mentionned above, the repartition among the patients have to be wisely selected due to data scarsity. Each of the groups need to be well split into the 2 datasets. A raisonable ratio, in terms of patient, would be 12/8 for ALS (i.e. 8/2 for CTRL) or 14/6 for ALS (i.e. 6/4 for CTRL).

It appears that this method allow to have an increase in terms of accuracy performances and more stability arises (the standard deviation is lowered). But, as discovered after multiple iterations, the choice of the patient repartition (and implicitly the number of samples of one group over another) have a great influence on the final result. A finetuned model which has been trained on a bigger amount of ALS samples will have a big problem to deal with CTRL when predicting. Therefore the results are biased. However, the case where the finetuned model is trained on a bigger amount of CTRL never happens!

An other TL experiment is, then, made, using the technique of "features extraction". The pre-trained models are used, not to predict classes, but to extract features from data.

To this end, each of the three provided models are "cut" at different levels:

- After the last block: in this case, only the Linear layer is removed. This kind of model extracts 37 features for each spectrum.
- Before the last block: here, the last entire block is removed. It results in 74 features for each spectrum.
- Before the two last blocks: the entire two last blocks are removed from the pre-trained model. It is producing 147 features for each spectrum.

The "block" term is used in reference at the "Residual block" class defined by the bacteria-ID paper[?]. This class is the main component of each pretrained model.

From this technique, 9 features extractors are obtained, taking the data as input ang returning features as output. To obtain predictions, two different models are proposed:

- Classical ML model: Decision Tree Classifier
- Deep ML model described in table 2.

Table 2: This table describes the deep ML model created

Layer type	Input	Output
InputLayer	Number of features extracted	Number of features extracted
DenseLayer	Number of features extracted	128
DenseLayer	128	64
DenseLayer	64	32
DenseLayer	32	16
Dropout	16	16
DenseLayer	16	1

On classical models, the performance are increased. On deep models, the same problem of data repartition (as mentionned in the finetuning part) is raised. Indeed, when data are randomly "well" distribued, this method can achieve really good performances. But when data are not, very bad performances are noticied.

Nevertheless, the repartition among patients is not the core problem. The number of spectral samples per patient influence drastically the final result. The logical conclusion to raise is to develop some techniques of data augmentation in order to have the intersection of:

- 1. a better repartition of samples per group (ALS/CTRL)
- 2. a better repartition of samples per patient (on average)

This led us to the final step of our project which is on the data augmentation techniques. 3 basic ones have been tested:

- Random Offset  $(\pm 0.1 \text{ x standard deviation of training set})$
- Random Mulitplication (1  $\pm$  0.1 x standard deviation of training set)
- Gaussian noise

Although change is low, the performances increased in the finetuning case as shown in the following section.

The data augmentation techniques previously described are also applied with features extraction. For runtime reasons, only one features extractor is used, which is the one obtained from the second pre-trained model with a cut before the last block. As expected, the dataset obtained is more balanced, which allows more constant results.

### 4 Results and Evaluation

For each experiment, the results are variable. So, the best results obtained are presented below.

Using the second pre-trained model (finetuned on bacterial spectra) to predict on raw dataset, an average accuracy score of 66% with a standard deviation of 1.11 is obtained. (Figure 1)

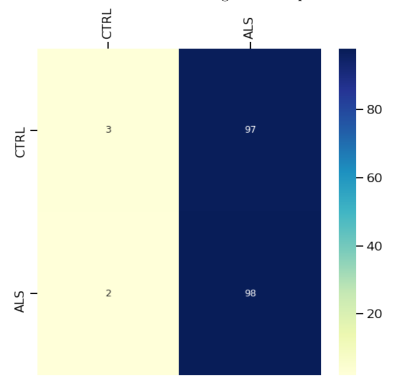
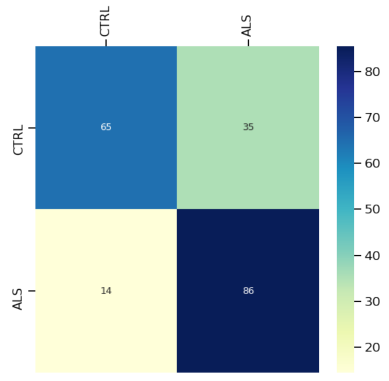


Figure 1: Confusion-matrix resulting from the pre-trained model

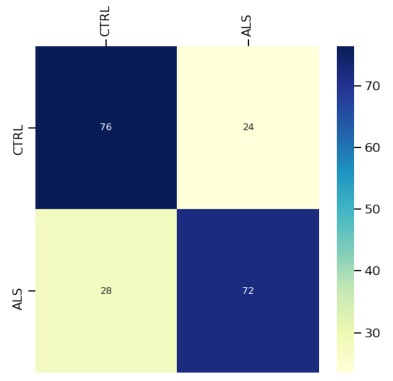
Adding fine-tuning method described in the previous section to the same model resulted in an accuracy of  $72.8 \pm 0.04$ . These results are illustrated by the confusion matrix below. (Figure 2)

Figure 2: Confusion-matrix resulting from the pre-trained model, with fine-tuning method  $\,$ 



Whith the data augmentation, the accuracy obtained is sligthly better: 74.4  $\pm$  0.01. (Figure 3)

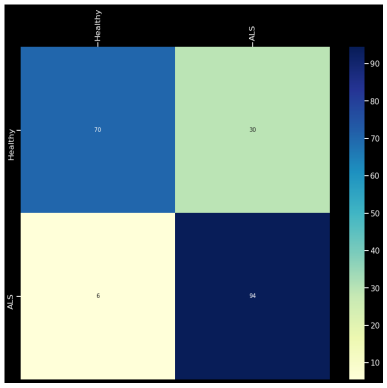
Figure 3: Confusion-matrix resulting from the pre-trained model, with finetuning and data augmentation



About the features extraction, the best performances are computed using the same pre-trained model and cuting its last block. The accuracy obtained is 72.6% which is slightly lower than the one with the fine-tuning process. This result is attained using a Deep ML model as predictor. (Figure 4) Applying the classical model the accuracy is better, but overfitting occurs.

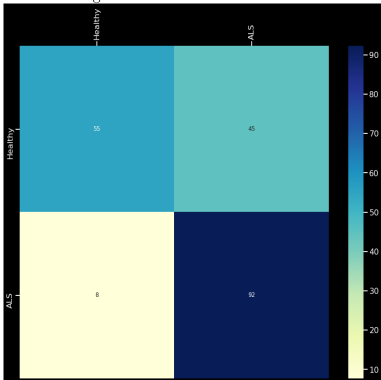
Figure 4: Confusion-matrix resulting from the pre-trained model as features

extractor



When data augmentation is applied to the previous extractor model, the results slightly improved. The accuracy obtained is about 77%. (Figure 5)

Figure 5: Confusion-matrix resulting from the pre-trained model as features extractor, with data augmentation



## 5 Discussion

As mentionned in the previous section the results chosen are the best obtained ones after multiple runs and are quite variable. Depending on the distribution of the patients (and the samples) in the finetuning and features extractor, performances can drop. It is quite problematic as it is a medical concern. In fact, if a healthy patient is diagnosed and treated for an illness it can be dangerous.

Our approach seems to be promising. However, some improvements could be done :

- Better data augmentation methods
- Different splitting method

• Try different deep and ML models (and better tune them!)

### 6 Conclusions

The Transfer Learning approach is a good approach for the analysis of the Raman spectra, especially in case of data scarsity. In addition, data augmentation techniques are helpful to make more robust models. A future work could be to find some better way to augment spectral data like Generative Adversarial Networks (GANs). The generative models could demonstrate an improvement in the accuracy of ML classification and also a solution to the problem of requiring a large amount of training data.

# 7 Data and code availability

All the data and the code are available on GitHub

### References

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