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PAPER

Deep learning of resting-state electroencephalogram signals for three-class classification of Alzheimer's disease, mild cognitive impairment and healthy ageing

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Abstract

Objective. This study aimed to produce a novel deep learning (DL) model for the classification of subjects with Alzheimer's disease (AD), mild cognitive impairment (MCI) subjects and healthy ageing (HA) subjects using resting-state scalp electroencephalogram (EEG) signals. Approach. The raw EEG data were pre-processed to remove unwanted artefacts and sources of noise. The data were then processed with the continuous wavelet transform, using the Morse mother wavelet, to create time-frequency graphs with a wavelet coefficient scale range of 0–600. The graphs were combined into tiled topographical maps governed by the 10–20 system orientation for scalp electrodes. The application of this processing pipeline was used on a data set of resting-state EEG samples from age-matched groups of 52 AD subjects (82.3 \pm 4.7 years of age), 37 MCI subjects (78.4 \pm 5.1 years of age) and 52 HA subjects (79.6 \pm 6.0 years of age). This resulted in the formation of a data set of 16197 topographical images. This image data set was then split into training, validation and test images and used as input to an AlexNet DL model. This model was comprised of five hidden convolutional layers and optimised for various parameters such as learning rate, learning rate schedule, optimiser, and batch size. Main results. The performance was assessed by a tenfold cross-validation strategy, which produced an average accuracy result of $98.9 \pm 0.4\%$ for the three-class classification of AD vs MCI vs HA. The results showed minimal overfitting and bias between classes, further indicating the strength of the model produced. Significance. These results provide significant improvement for this classification task compared to previous studies in this field and suggest that DL could contribute to the diagnosis of AD from EEG recordings.

1. Introduction

Dementia is a term that describes a collection of diseases that affects approximately 50 million people worldwide (Prince *et al* 2015). It is characterised by a measured cognitive decline in two or more domains such as memory, language, behaviour and personality, ultimately leaving the individual unable

to perform simple everyday tasks (Weller and Budson 2018). The global healthcare cost of dementia is upwards of \$818 billion, which is increasing year by year (Prince et al 2015). Alzheimer's disease (AD) contributes to approximately 60%–80% of the global dementia diagnoses and is most prevalent in adults aged 60 and above (Weller and Budson 2018). The biggest risk for AD is age, with a reported doubling

of the disease prevalence every 6.3 years after the age of 60 (Prince *et al* 2015). Other factors include health risks such as high body mass index, high fasting glucose, smoking and increased intake of sugar-sweetened beverages (GBD 2016 Dementia Collaborators 2019).

AD is related to neurofibrillary tangles and amyloid plaques developing in the cerebral cortex area of the brain, especially in the hippocampus (DeTure and Dickson 2019). An intermediary stage between healthy ageing (HA), sometimes referred to as healthy control, and AD has been widely recognised as a stage called mild cognitive impairment (MCI). It is currently unclear who amongst individuals with MCI will develop AD dementia (Kramer et al 2007). At present, the only definitive way to diagnose AD is via post-mortem examination to find the plaques or tangles within the brain (DeTure and Dickson 2019). However, with the advent of novel dementia biomarkers which can be gathered in vivo, a new biological definition of AD has been introduced (Jack Jr et al 2016) (Jack Jr et al 2018). Biomarkers are expensive, invasive, little specific, and only available in specialised centres (Parra et al 2019).

The gold standard, or current state of the art, for diagnosing AD uses mental examinations combined with costly and time-consuming neuroimaging scans such as magnetic resonance imaging and positron emission tomography (Cassani *et al* 2018). These imaging tools aim to highlight biomarkers, such as amyloid- β peptides, that indicate the formation of plaques within the brain. This method is highly dependent on trained doctors that interpret and analyse the results to determine the diagnosis. The reported diagnostic accuracy of AD by experts alone is only 77% (Sabbagh *et al* 2017).

Early diagnosis of AD, during the MCI stage, would allow healthcare professionals to avoid misdiagnosis, deliver quick and more appropriate treatment options, and provide better overall disease management. To achieve this, the analysis of electroencephalogram (EEG) signals has been suggested by researchers to find features and biomarkers that may aid in AD diagnosis (Dauwels et al 2010, Rossini et al 2020).

Conventionally, EEG analysis has been used clinically to evaluate different conditions such as epilepsy, sleep disorders and strokes (Britton et al 2016). In addition, these signals have also been analysed in research settings using visual and statistical methods for the diagnosis of AD. The drawback of this is that standalone, EEG analyses have not produced results which can compete with standard practices (Craik et al 2019). Typically, EEGs are subject to signal processing techniques that provide condensed features for classification. There are three main features of EEGs that differ between AD and HA which are: slowing of the EEG, reduced complexity of the EEG and EEG synchrony (Dauwels et al 2010). Examples of signal processing methods for the detection of slowing of the EEG include time frequency analysis

techniques such as discrete Fourier transform, power spectral density (PSD) and continuous wavelet transform (CWT). Entropy and Lempel-Ziv complexity are examples of algorithms used to explore EEG complexity and the Pearson correlation coefficient and Coherence function have been used to identify perturbations in EEG synchrony (Dauwels *et al* 2010). The problematic poor performance of EEG analysis for AD diagnosis has drawn interest from new developments in Artificial intelligence (AI), with the ability to analyse signals with increased complexity and depth.

Deep learning (DL) is a subsection of AI that has become popular in recent years due to advancements of Graphics processing units (GPUs) in computing. It aims to mimic the learning of the human brain by using complex algorithms to obtain features of data that cannot be seen using conventional statistical analysis methods. Thus, millions of learnable parameters, particularly useful in image classification problems, are built to detect perturbation features. These algorithms have proven results that exceed human performance in classification problems such as ImageNet (Dodge and Karam 2017). Many biomedical engineering applications, including EEG analysis, can produce graphical image outputs that can be used as inputs to these DL networks.

It is hypothesised that improved accuracies for the diagnosis of AD can be achieved using DL to correctly classify the EEG recordings from AD, MCI or HA subjects. This study employed a novel signal processing technique to convert complex EEG recordings into usable input images for a DL network. A pre-trained DL model was optimised for this study's focus, trained using the ground truth values associated with each image. It was then cross validated to evaluate the reported classification accuracy result.

The outline of this paper is as follows. Section 2 details the related literature and studies in this field. Section 3 describes the materials and methods used. Section 4 displays the results of the study and section 5 contains the discussion. Finally, section 6 presents the conclusions and further work.

2. Related studies

This section details seven studies that relate to the research topic of improving diagnosis of AD using DL of EEG signals. It is split into two sections detailing three-class and two-class classification.

The first reported paper to complete three-class (tertiary) classification of AD vs MCI vs HA using DL techniques was by Morabito *et al* (2016). The authors propose a two-layer deep convolutional neural network (CNN) that uses extracted features from time-frequency maps as the input to the network. The signal processing method used was CWT with the Mexican Hat mother wavelet. The reported accuracy was relatively low at 82% and used a reasonably

small data set consisting of 23 AD, 23 MCI and 23 HA subjects.

A discriminative deep probabilistic model was proposed by Bi and Wang (2019) which produced an accuracy result of 95.04% for three-class classification of AD, MCI and HA. The employed signal processing technique created spectral topography maps from raw signals, showing the relative power (RP) per frequency band across each electrode in one image. One shortfall of this paper is that, although a combined 12000 images were used across training and validation, the data originated from a very small sample size of four subjects per class. This could indicate that the results are more susceptible to error and less accurate than what is stated within the paper.

A paper published by Ieracitano et al (2019) used 2D greyscale PSD images as an input to a CNN for the three-class classification of AD, MCI and HA. The authors report an accuracy of 83,33% for the CNN, which was superior compared to shallow machine learning techniques such as support vector machine (SVM), multi-layered perceptron (MLP) and linear discriminant analysis. This result showed clear accuracy benefits for using CNNs within this field; however, the authors correctly conclude that without future work, this method would not be sufficient alone for clinical diagnosis of AD. In addition to the large data set of 63 HA, 63, MCI and 63 AD subjects, a clear explanation of the importance of avoiding bias by balancing and age-matching the data set was included. Despite the large number of subjects, there were only 2340 images used within the model due to the sampling size which is below the general guidelines of 1200 images per class (Stanford Vision Lab 2010).

The most recent published paper by Ieracitano et al (2020) used a 'multi-modal machine learning' approach to classify EEG recordings in dementia. The authors make use of a novel combination of features extracted from Bispectrum analysis and CWT time-frequency analysis to classify AD vs MCI vs HA. The results indicate a maximum of 89.22% accuracy for this classification, which is a significant increase in using either signal processing method individually. The results published are supported with statistical evidence and used the largest number of features seen within this field equalling 207900. However, the authors do not innovate within machine learning, using a simple MLP as the classifier for this problem. It successfully identifies areas of future research, such as linking this work with a feasibility study using a novel spiking neural network (NN) architecture called NeuCube (Capecci et al 2014).

Three of the seven papers analysed only reported two-class classification which severely reduced the complexity requirement of the models proposed.

Zhao and He (2015) used time-domain EEG signals as the input to a restricted Boltzmann machine (RBM) network to obtain features from the data,

which are classified using an SVM. The unique feature of this paper is that the authors conducted a detailed optimisation experiment on the number of hidden layers and number of nodes within each layer for this network. They conclude that a low number of layers (L=3) and high number of nodes (n=2000) was preferential, resulting in a two-class (binary) classification accuracy of 92%. The authors reduced computing time by only using 10 out of the 30 available subjects (15 AD, 15 HA) but showed that the sample size was small in comparison to a relatively large 19200 input images.

Kim and Kim (2018) investigated early diagnosis of AD by producing a model that detects the difference between MCI and HA subjects. The model comprises of a deep neural network (DNN) with featurebased inputs relating to the RP of different frequency bands within EEG signals. The model's performance was poor in comparison to other papers analysed, with just 75% accuracy over a small data set of 10 HA and 10 MCI subjects. In comparison to the paper by Zhao and He (2015), the authors also investigate the effect of the number of hidden layers within the network but report the highest accuracy with the largest number of hidden layers tested (L = 4). This difference could be explained by various factors such as the difference in NN type, data set and type of two-class classification.

Fan et al (2018) made use of a complexity measure, multiscale entropy (MSE), to extract features of the EEG signals, in numerical form, for the input to a DL linear regression model. A calculated 46470 features were used which produced a maximum accuracy result of 82%, low compared to the other papers in this section. An interesting data set classification was used which compared HA subjects to three different severities of AD based on increasing clinical Dementia rating (CDR) scores (AD1, AD2 and AD3). The specificity of each AD diagnosis has advantages and disadvantages; it could lead to better diagnosis accuracy results but limits the model to this data set as the CDR scale is not readily available for many other databases. Although the data set was large with 123 subjects, it was also highly unbalanced with only 15 HA subjects. The authors were able to suggest future improvements including altering the MSE method, collecting more data and assessing the interactions between electrodes.

A summary of the key information presented in each discussed paper is detailed in table 1.

Cassani et al (2018) produced a review paper on EEG for AD diagnosis and includes recommendations for the future of this research area. The authors' recommendations focus on the EEG databases, which should be balanced around demographics such as age, gender, number of subjects and education level as well as being as large as possible. The paper also suggests that clearer and more detailed information should be provided about the process of machine or DL. This

Table 1. A summary of the previous directly related papers in the field of DL of EEG signals for the diagnosis of AD, showing the database size and age matching, signal processing methods employed, machine learning architectures used and final classification results.

Reference	Database size and age matching	Signal processing method	Machine learning architecture	Results
Ieracitano et al (2020)	63 HA 63 MCI 63 AD Balanced	Features from Mexican Hat CWT and Bispectrum estimation.	MLP	AD-MCI-HA = 89%
Bi and Wang (2019)	4 HA 4 MCI 4 AD Balanced	2D RGB images by combining spectral topographical maps	DCssCDBM Two hidden layers	AD-MCI-HA = 95%
Ieracitano et al (2019)	63 HA 63 MCI 63 AD Balanced	2D greyscale Periodogram images	CNN One hidden layer	AD-MCI-HA = 80% MCI-HA = 92% AD-HA = 91% MCI-AD = 84%
Kim and Kim (2018)	10 NC 10 MCI Age-matched	Features from RP	DNN Four hidden layers	MCI-HA = 75%
Fan et al (2018)	15 HA 15 AD1 69 AD2 24 AD3 Unknown	Features from MSE Analysis	LASSO Model	HA-AD1 = 42% HA-AD2 = 69% HA-AD3 = 79% AD1-AD3 = 82% AD2-AD3 = 72% AD1-AD2 = 71%
Morabito et al (2016)	23 HA 23 MCI 23 AD Balanced	2D RGB images from Mexican Hat CWT	CNN Two hidden layers	AD-MCI-HA = 82% MCI-HA = 85% AD-HA = 85% MCI-AD = 78%
Zhao and He (2015)	15 HA 15 AD Unknown	Raw Data	RBM Three hidden layers	AD-HA = 92%

AD = Alzheimer's disease, CNN = Convolutional neural network, CWT = Continuous wavelet transform,

DCssCDBM = Discriminative contractive slab and spike convolutional deep Boltzmann machine, DNN = Deep neural network, HA = Healthy aging, LASSO = Least absolute shrinkage and selection operator, MCI = Mild cognitive impairment, MLP = Multi-layer perceptron, MSE = Multiscale entropy, RBM = Restricted Boltzmann machine, RGB = Red, green and blue, RP = Relative power.

is apparent after reviewing the above seven papers, as they all provide different levels of detail surrounding the experimentation and design process and make it difficult to reproduce or build upon their results. The reviewed papers show the possibility of high accuracy classification using a combination of signal processing and DL of EEGs, and there are lots of different methods that are yet to be explored.

3. Materials and methods

3.1. Model design

The overall design of the proposed model within this study is outlined in the experimental flow diagram in figure 1. First, the raw EEG signals were pre-processed and split into epochs of 5 s. Each sample was then subject to signal processing and converted into RGB colour images that were suitable as an input to an optimised DL NN. The resulting image data set was then randomly split into ten folds, from which a discrete portion of the folds were used in the training, validation, and test splits of the DL NN model. The model was then validated using k-fold cross validation and assessed using confusion metrics.

3.2. Subjects

This study consisted of 141 subjects originating from the Behavioural and Cognitive Neurology Unit of the Department of Neurology and the Reference Centre for Cognitive Disorders at the Hospital das Clinicas in São Paulo, Brazil (henceforth referred to as the 'Brazil' study) (Cassani et al 2017). The Brazil study diagnosed AD and HA subjects according to the National Institute of Neurological Disorders and Stroke and Alzheimer's disease and related disorders (NINCDS) criteria. Typically, within this study AD subjects presented a Mini Mental State Examination (MMSE) score of ≤ 24 and a CDR of 0.5–2, and HA subjects presented an MMSE score of ≥25 and a CDR of 0. The AD subjects were required to have shown functional and cognitive decline over the previous 12 months. The MCI subjects are presented with an MMSE score of ≥24 and a CDR of 0-0.5, with objective evidence of impairment in one or more cognitive domains but retaining independence in functional abilities (Albert et al 2011) (Petersen and Knopman 2006) (Petersen and Negash 2008). In addition, all subjects were required to display absence of conditions that cause cognitive decline specified

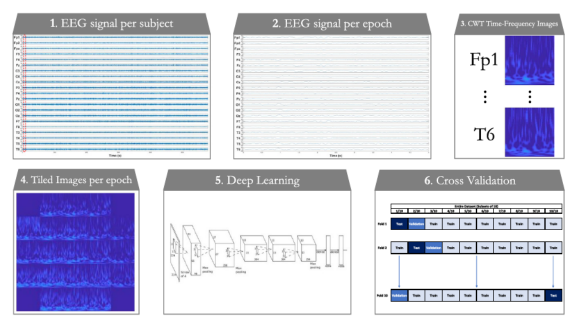


Figure 1. Experimental flow diagram showing a simplified version of the proposed model. The training folds/test folds describe a proportion of the data assigned to train or test the model respectively. Reproduced with permission from Krizhevsky et al (2012).

as diabetes mellitus, kidney disease, thyroid disease, alcoholism, liver disease, lung disease or vitamin B12 deficiency (Cassani *et al* 2017).

The EEG recordings from the Brazil study were collected using a resting-awake eyes-closed method using the Braintech 3.0 EEG device (EMSA Equipamentos Médicos Inc., Brazil) with a sampling frequency of 200 Hz. The electrode positioning consisted of 21 electrode, positioned on the scalp according to the 10–20 layout system (Fp1, Fp2, Fpz, F3, F4, Fz, C3, C4, Cz, P3, P4, Pz, O1, O2, Oz, F7, F8, T3, T4, T5, T6), in which each point corresponds to a brain region. A time series of 587 s (~10 min) was captured for each subject (excluding 2 AD, 1 MCI and 3 HA subjects which had shorter recordings).

3.3. EEG pre-processing

The data were pre-processed by utilising a sequence of different methods. They were first filtered using a 1-60 Hz band-pass, FIR filter with an order of 330 and de-noised using independent component analysis (ICA) and notch filters at 21 and 42 Hz to remove significant oscillatory noise artefacts and its harmonic. They were processed further using the multiple artefact rejection algorithm (EEGLAB plugin for MATLAB®) to automatically classify and remove other noise related ICA components. The de-noised data set was then split into age-matched groups of 52 AD subjects (82.3 \pm 4.7 years of age, MMSE score of 21.0 \pm 4.8), 37 MCI subjects (78.4 \pm 5.1 years of age, MMSE score of 25.4 \pm 2.7) and 52 HA subjects $(79.6 \pm 6.0 \text{ years of age, MMSE score of } 27.5 \pm 1.6).$ It is noted that 22 AD subjects, 19 MCI subjects and 24 HA subjects did not have recorded MMSE scores but were still used for this analysis. Misclassification

of older HA subjects and younger AD subjects can occur when comparing subjects of a large age-range due to the natural progressive neurological degradation during human ageing. Therefore, it is important to use data sets with age-matched subject groups to ensure reliable results (Dukart *et al* 2011).

The time series was then split into 5 s epochs, removing 3.5 s from the start and end of the signal to account for any discrepancies or noise. Using all the raw data available from the 141 subjects over the 21 electrode positions, a total of 340137 samples of 1000 data points were made available for analysis. The number of artefact-free epochs for each subject differed, however all available data were included to provide as many data as possible for analysis.

3.4. Signal processing

For each sample, the data were processed using the CWT, shown by equation (1):

$$W(a,b) = \int_{-\infty}^{\infty} x(t) \frac{1}{\sqrt{|a|}} \psi^* \left(\frac{t-b}{a}\right) dt \qquad (1)$$

where: t = time; $\psi = \text{mother wavelet}$; x(t) = time domain signal; a = range of scales; b = translations; W(a,b) = wavelet coefficients (amplitudes of a series of wavelets), * = complex conjugate.

The CWT function is an analysis method that transforms the raw, one-dimensional input signal into the time-frequency (often called the time-scale) domain. The transform is controlled by a mother wavelet with a zero average which is subject to adaptations over a range of scales (a) and time translations (b), resulting in the output wavelet coefficients (W).

The signal is fit to match the wavelet, so it is important that an appropriate shaped mother wavelet is chosen for each individual application. There is a plethora of families of mother wavelet choices such as Haar, Daubechies, Coiflets, and Morlet, with the Mexican Hat wavelet proving most popular within this specific field as shown in table 1 (Mallat 2009). Despite this, a family of wavelets called the Morse wavelets, introduced by Olhede and Walden (2002), was chosen for this study (Greco et al 2003). This mother wavelet is a form of analytic wavelet, which are complexvalued wavelets designed for the analysis of modulated oscillations that have useful information in both magnitude and phase (Lilly and Olhede 2010). Due to this unique feature, the Morse wavelet is directly applicable to the complex and non-stationary nature of EEG signals.

3.5. Image data set generation

The output of the CWT function can be displayed visually as a time-frequency map. In this situation, a plot called a scalogram is created which plots frequency against time, with the energy of the CWT coefficients indicated by the colour of the plot.

When computing the CWT over many EEG samples, it is important to determine a standard for the colour bar scale. The colours presented on the plot are governed by the scale choice, including the maximum and minimum values and the scale base (i.e. logarithmic or linear). The scalogram plot, by design, plots absolute values of the wavelet coefficients, so a minimum value of 0 was chosen. To obtain the maximum value on the scale and the type of scale, the maximum value of the coefficients for each sample were plotted on a linear scale and a logarithmic scale. These plots indicated that a logarithmic scale would be more appropriate for this task as the data contained a high quantity of low maximum coefficient values in contrast to a low quantity of high maximum coefficient values, as shown in figure 2. By rounding the highest maximum coefficient to the nearest whole number, a maximum scale value of 6×10^2 was chosen that could encompass all the signals.

The colour map used for the time-scale representation of the coefficients was 'parula', with the default number of unique colours set at 256. This value was kept relatively low to reduce image complexity and therefore computational expense. A dark blue colour represents an intensity value of 0 (absolute coefficient value = 0), and a light-yellow colour represents an intensity value of 1 (absolute coefficient value = 6×10^2), changing colour logarithmically between these values (Mathworks 2020).

The generated scalogram plots were then saved and combined into tiled images based on the 10–20 system. The images relating to each electrode in a 5 s epoch were combined using the orientation in figure 3 to produce topographical images based on the 10–20 system.

The resulting image data set consisted of 16197 images, of which there were 6020 AD, 4289 MCI and 5888 HA images.

3.6. DL model

The DL model used was a modified AlexNet architecture that was optimised for three-class classification, shown in figure 4. AlexNet, is a deep CNN that consists of eight main layers, five of which are convolutional layers and the remaining three as fully connected layers (Krizhevsky *et al* 2012).

The architecture had to be altered by reducing the weights on the final fully connected layer from the standard 1000 classes to three. A variety of parameters and hyperparameters were optimised throughout the training of this model and were altered one-by-one over a range of values to provide a model which produced the best output accuracy performance.

3.6.1. DL parameters

This section will detail the DL parameter and hyperparameter choices that were optimised to tune the model. The tuned model provided the best results in terms of accuracy, loss and generalisation performance. These were manually adjusted and tested sequentially to obtain the final model. The order of optimisation was chosen by starting with the most impactful parameters first and ending with fine tuning parameters, relating to the performance determined by the final classification accuracy.

The optimiser choice was tuned first, comparing three common optimisers: stochastic gradient descent (SGD), moving average of squared gradients (RMSProps) and adaptive moment estimation (ADAM). Optimiser choice is important as it is used to update the weights of the model during training. SDG is a gradient descent method whereas RMSProp and ADAM are both adaptive techniques. In short, RMSProp is an extension of SGD and ADAM is a further extension of RMSProp (Ruder 2017). As expected, the ADAM optimiser produced the best accuracy performance (Kingma and Ba 2015).

The learning rate is one of the most impactful parameters as it describes how often the weights of the model are updated and is therefore important to be paired with the optimiser. Four learning rates were tested, 1×10^{-3} , 1×10^{-4} , 5×10^{-5} and 1×10^{-5} , with a learning rate of 1×10^{-4} producing the best accuracy performance.

The learning rate can either be kept constant or reduced over time with a policy. This policy can allow greater control over the weights at the later stages of training for fine tuning and avoidance of overfitting. The reduction is split into the period and the factor. The period describes the rate of change and the factor describes the magnitude of the reduction. During this step, factors of 1/2, 1/3 and 1/5 were combined with periods of 10 and 5 epochs and were

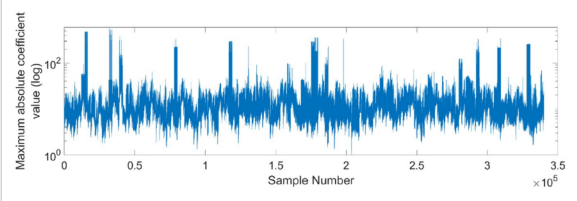


Figure 2. A line graph of the maximum absolute coefficient value on a logarithmic scale against sample number (n = 340137).

-	Fp1	Fpz	Fp2	
F7	F3	Fz	F4	F8
Т3	C3	Cz	C4	T4
T5	P3	Pz	P4	T6
-	01	Oz	O2	-

Figure 3. A 5×5 tiled image created from individual electrode position images.

compared against a stationary learning rate with no policy. This resulted in a chosen policy of factor = 1/3 and period = 10 epochs.

The batch size describes how many sample images are used in each iteration of the training. Smaller batch sizes result in noisy updates and are computationally expensive but they offer a good regularising effect. In contract, larger batch sizes decrease processing time but may have poorer generalisation to unseen data. Batch sizes of 50, 100 and 150 were chosen as an initial starting point, moving on to batch sizes of 16, 32, 64 and 128 due to MATLAB's computational efficiency when working with powers of 2. Although a smaller batch size produced superior accuracy results, a batch size of 64 was used as it produced results that had a good compromise between the advantages of both small and large batch sizes, running each fold in approximately 30 min using a GTX 1050Ti GPU.

Validation patience is a method of early stopping and is governed by the number epochs used to train the model. Patience is beneficial as it reduces the chance of overfitting and gives the model enough time to generalise. It does this by stopping the model a certain number of epochs after it has reached its lowest validation loss. The validation patience of ten allowed the model to reach its minimum validation loss whilst avoiding overfitting. One epoch is comprised of using each sample in the whole data set once.

Weight learn rate factor (WLRF) and bias learn rate factor (BLRF) were the last parameters that were

optimised and relate directly to the final fully connected layers of the network. They determine the relationship between the weight and bias learning rates and the global learning rate, used to fine-tune the result. Values of 10, 20, 30 and 40 for both WLRF and BLRF were tested which varied the final accuracy by ~0.1%. From this, values of 20 for both WLRF and BLRF was chosen as it produced the best combined validation accuracy result.

A summary of each parameter, its value and the justification for the choice can be seen in table 2.

3.7. Reporting metrics

To assess the performance of the proposed model, Confusion Matrices and receiver operating characteristic (ROC) curves were generated, in addition to the calculation of four different confusion metrics (Hossin and Sulaiman 2015).

Confusion matrices show the relationship between the predicted classes of each image in the test data set (predicted output) in relation to the true class (ground truth) of each image in the test data set (true output), given by the predefined label. In general terms, the results along the diagonal show correct predictions and off-diagonal values show the incorrect predictions. Within this, there are four bins each prediction can belong to:

- True positive (TP)—Predicting the sample as positive when it is actually positive (i.e. AD classified as AD).
- True negative (TN)—Predicting the sample as negative when it is actually (i.e. MCI classified as MCI and HA classified as HA).
- False positive (FP)—Predicting the sample as positive when it is actually negative (i.e. MCI/HA classified as AD). Also known as a Type I error.
- False negative (FN)—Predicting the same as negative when it is actually positive (i.e. AD classified as MCI/HA). Also known as a Type II error.

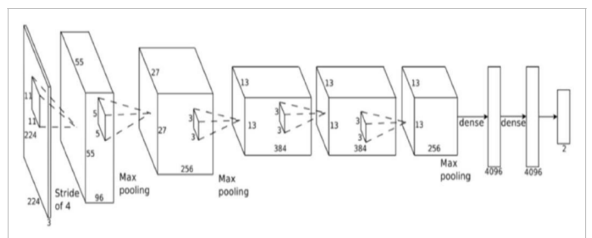


Figure 4. A schematic of the AlexNet DL architecture, showing the tensor size as cuboids, kernel size as dashed pyramids and informative descriptions of areas with max pooling and stride lengths, culminating in an example binary classifier. Reproduced with permission from Krizhevsky et al (2012).

Table 2. A summary of the optimised parameters, detailing the parameter name, chosen value and justification of the choice.

Parameter	Value	Description	
Architecture	AlexNet	Shallowest investigated network but produced the fastest and most accurate results.	
Optimiser	ADAM	Proven in experiment and literature to be a high performing optimiser for image classification learning.	
WLRF	20	Optimised to provide the highest accuracy values.	
BLRF	20	Optimised to provide the highest accuracy values.	
Mini batch size	64	Maintains a good relationship between the usage of GPU memory and model fine detail.	
Validation frequency	177	Provides a validation point at the end of every epoch.	
Validation patience	10	Provides enough time for the validation accuracy results to stabilise.	
Initial learning rate	1e ⁻⁴	Set to allow for a learning rate curve that asymptotes to zero roughly following a smooth curved trajectory.	
Learn rate schedule	Piecewise	Allows the learning rate to be altered as the model progresses.	
Learn rate drop period	10	Reduces learning rate by 1/3 every ten epochs which were optimised for this	
Learn rate drop factor	0.33	application to avoid overfitting.	

These definitions relate directly to the calculation of the per class accuracies shown in equation (2):

$$Accuracy = \frac{TP + TN}{TP + FP + FN + TN}$$
 (2)

where: TP = True positive; TN = True negative; FP = False positive; FN = False negative

3.8. Model validation

The proposed model was validated using k-fold cross-validation (k=10) using the ten splits of the image data set. This method is a statistically based evaluation method which is applied to a DL model with a limited data set to show its expected performance on new data. The data set is shuffled and split into folds randomly, reducing the chance of the order of the data affecting the model results.

For each fold, 8/10 folds were used for training, 1/10 fold was used for validation and 1/10 fold was used for testing. Using the results from the ten folds, the average (mean) and standard deviation (std) of the reporting metrics were calculated.

4. Results

For each subject in the database, the EEG signals were split into sample epochs of 5 s for each electrode. For each sample, time-frequency maps were created using the CWT for each electrode (number of images produced from data analysis = 340137), then the electrode images were combined using the 10–20 system to create one image per epoch (number of images = 16197). The images, an example of which is shown in figure 5, were randomly split into ten folds, comprising of 8/10 folds training data, 1/10 fold validation data and 1/10 fold test data, ensuring that there was the correct proportion of classes within each split. These were then fed into an optimised AlexNet DL model to predict the classes associated with each image. The model was split into ten folds and assessed using k-fold cross validation for robustness, producing results in approximately 30 min per test.

The results of the ten-fold cross validation tests are shown in table 3 and present an overall test accuracy of 98.9 \pm 0.4%. The confusion matrices of fold 1–10 are shown in figures 6(a)–(j), respectively.

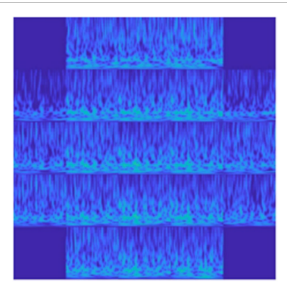


Figure 5. An example output image, showing a 5×5 tiled image of CWT scalograms as detailed in figure 3.

Table 3. Model performance of each fold, detailing each class accuracy and the overall accuracy, including the mean and standard deviation of each category, shown to one decimal place.

Fold	AD class accuracy (%)	MCI class accuracy (%)	HA class accuracy (%)	Overall accuracy (%)
1	99.5	97.7	98.5	98.6
2	99.0	97.9	99.3	98.8
3	99.2	98.1	98.3	98.6
4	99.8	98.1	99.0	99.1
5	99.0	97.2	97.5	98.0
6	99.3	98.6	99.0	99.0
7	99.7	99.1	99.0	99.3
8	99.5	99.1	99.3	99.3
9	98.5	98.1	99.2	98.6
10	99.3	99.1	99.2	99.2
Mean	99.3	98.3	98.8	98.9
Std	00.4	00.6	00.5	00.4

To show additional information about the model, the validation loss and accuracy values were used as a comparison to the test accuracy. The average validation loss value, 0.07 ± 0.02 , is very low, which showed that the model was fit to the data well. Across the folds, the average validation accuracy was $98.8 \pm 0.3\%$ which is similar to the overall average test accuracy, indicating that the model has been trained appropriately to stop over or underfitting.

In relation to class specific classification, there was an accuracy for AD of 99.3 \pm 0.4%, MCI of 98.3 \pm 0.6% and HA of 98.8 \pm 0.5%. This shows a very small bias towards the HA and AD classes, indicative of the bias within the original data set. The overall range of 1.0% shows that this bias is minimal, and the skewed data set has been accounted for in the model.

The low standard deviation values across all the accuracy values also shows robustness and consistency between the ten validation tests.

5. Discussion

This paper has presented a classification method with an overall three-class classification accuracy for AD vs MCI vs HA of 98.9 \pm 0.4%. The resulting classification accuracy showed minimal bias between class predictions over a large database of 52 AD, 37 MCI and 52 HA subjects. The proposed method extracted artefact free, 5 s long EEG signals and analysed them using the CWT signal processing method with a Morse mother wavelet. Image maps created from the output scalogram graphs were then used as the input to an optimised DL model based on the AlexNet architecture for three-class classification. The model was assessed using a ten-fold cross validation method, presenting results that improved upon current reviewed literature within the proposed field.

During the development of this model, alternative signal processing techniques and pre-trained DL models were researched and tested. Networks such as IN Publishing J. Neural Eng. 18 (2021) 046087 C J Huggins et al.

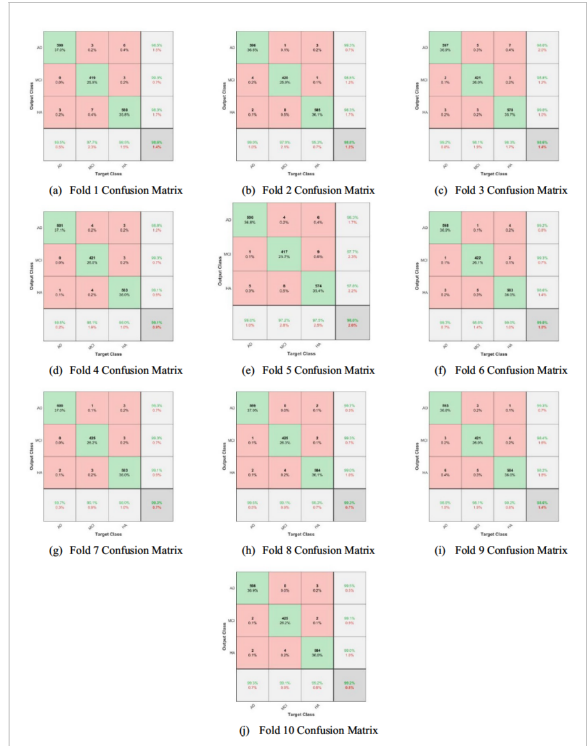


Figure 6. (a)–(j) Confusion matrices showing the model predictions (output class) against the ground truth (target class) from folds 1–10 respectively, relating to table 3.

'ResNet-18' and 'GoogLeNet' were investigated and compared to 'AlexNet' leading to surprising results. ResNet-18 has 18 convolutional layers and makes use of residual learning every two convolutional layers to skip over layers of the network if required. These address the vanishing gradient problems which occurs when adding a large number of layers to a network (He *et al* 2015). GoogLeNet is even larger, with 22 convolutional layers. GoogLeNet introduced

a unique model that uses inception modules to reduce the depth of the model by collating convolutional resulting using parallel layers of 1×1 , 3×3 and 5×5 kernel filters. This resulted in a model that has 12-time fewer parameters than AlexNet, which has 5000000. It also contains two auxiliary classifiers that offer a regularising effect on the network (Szegedy *et al* 2014). Despite the increased complexity of both GoogLeNet and ResNet-18 compared to

AlexNet, they produced models that took longer to train and had worse accuracy results. This is suggested to be due to more overfitting present due to higher divergence in training and validation loss plots.

Alternative signal processing methods could have been used in this method such as short-term Fourier transform (STFT). The output spectrogram from STFT analysis is very similar to the scalogram from CWT analysis, with the main difference being the resulting frequency and time resolutions. STFT uses a fixed window size resulting in time-frequency graphs with uniform time and frequency resolutions. A trade-off between the frequency and time resolutions must be made when choosing the fixed window size for STFT, which is not required for CWT. CWT uses variable, scaled window sizes to create graphs with non-uniform time and frequency resolutions. The non-uniform resolutions allow lower frequency trends in the data to be seen at longer time intervals and higher frequency trends to be seen at shorter time intervals. Due to the non-stationary nature and varied frequency range of EEG signals, CWT was chosen as the most appropriate signal processing method.

In comparison to the literature reviewed in section 2, the results improve on the accuracy of HA vs MCI vs AD presented by the most recent study from Ieracitano et al (2020), who produced a combined Bispectrum and CWT feature model, with a maximum average accuracy of 89%. Their paper uses more subjects (n = 189) in comparison to the subjects in this study's database (n = 141) and is balanced between classes which means the data are better suited for DL applications. The results are also calculated over a range of ten tests which gives reliability to their results. The results also improve on the maximum accuracy found in literature, from Bi and Wang (2019) at 95% accuracy of HA vs MCI vs HA using an alternative method and data set. The input to their bespoke machine learning model uses a unique combination of spectral topography maps to produce this result, however, lacks depth as the subject pool is very small (n = 16) in comparison to our data set. Both papers included a statistical analysis of the results, such as ROC curves and confusion matrices, further allowing direct discussions and comparison between the papers.

The other papers in section 2 by Kim and Kim (2018), Fan et al (2018) and Zhao and He (2015) are all binary classifiers. As this research study produced three-class classification of AD vs MCI vs HA, it has a clear impact advantage. The signal processing methods of RP from Kim and Kim (2018) and MSE from Fan et al (2018) are less computationally complex compared to CWT. Conversely, their respective DNN and LASSO classification models were more complex than the proposed CNN in this study. All three papers also use databases that are much smaller compared to the database used in this study, despite this still produce lower accuracy results.

Despite the promising results presented, our study has limitations. The recording of scalp EEG signals was conducted in a controlled environment, requiring clinical space and trained professionals which can be time consuming and expensive. Bias is an important topic in medical AI and DL and should be reduced as much as possible. By removing human interaction in the signal processing and classification stages, this study has attempted to avoid bias; however, there were still areas for improvement. The database required initial input of labels for this supervised learning method, which means that there was a possibility of errors within the diagnosis via the standard clinical criteria explained in section 3.2 (Batum et al 2015). The educational level of the subjects, which is a known contributor to AD, has not been assessed in the data pre-processing. The similarities and differences between the 5 s epochs within the 10 min samples have also not been assessed, which could affect the results. Furthermore, it is unrealistic to compare these results directly to the in vivo diagnostic accuracy of AD (77%) as the results within this report rely on pre-defined labels using gold standard techniques. To be used in a clinical setting, this method would need to be validated using additional data, ideally progressing onto a model that produces significant results regardless of the patient's demographical information such as age, location, and educational history.

6. Conclusions

In this study, a signal processing method combined with an optimised DL model has been developed that presents accuracy results of 98.9%, currently higher than any results presented in literature within this field.

Other methods were tested and discussed, including spectrogram time-frequency images, alternative DL architectures GoogLeNet and ResNet-18 and a variety of different hyperparameter values. These different strategies did not produce superior accuracy results but provided a useful comparison to the final model.

This study has shown some very promising results but further work is required to progress this research, with the end goal to produce a clinically accurate tool that could be used to aid healthcare professionals in providing better diagnosis for elderly patients with symptoms that are typical of MCI or AD. Recommendations for future work could encompass the following:

Test this model as a two-class problem by comparing the AD vs MCI accuracies against the MCI vs HA accuracies. This leads to a hypothesis that a lower number of classification categories would result in improved accuracies.

- Explore alternative ways to create RGB images from EEG signals, possibly by combining other signal processing methods quantifying changes in these signals related to AD.
- Explore additional pre-trained networks or bespoke DL architectures.
- Use the model on other databases (out of sample data) to further understand its reliability and accuracy.
- Test the use of automatic hyperparameter selection using methods such as Bayesian optimisation.
- Create an 'International Standard' data set for training and testing similar systems for the diagnosis of AD.

The findings presented in this study have significantly added to the continued knowledge surrounding DL of EEG signals for AD diagnosis. The increased accuracy results show promising outcomes for future applications of DL to the classification and diagnosis of AD. This could eventually help combat the resource-intensive and human dependant methods that are currently used, ultimately providing a quantitative probability value for the diagnosis of a patient.

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References

- Albert M S et al 2011 The diagnosis of mild cognitive impairment due to Alzheimer's disease: recommendations from the national institute on aging-Alzheimer's association workgroups on diagnostic guidelines for Alzheimer's disease Alzheimers Dementia 7 270–9
- Batum K, Cinar N, Sahini Ş, Cakmak M A and Karsidag S 2015 The connection between MCI and Alzheimer disease: neurocognitive clues *Turk. J. Med. Sci.* 2105 1137–40
- Bi X and Wang H 2019 Early Alzheimer's disease diagnosis based on EEG spectral images using deep learning Neural Netw. 114 119–35
- Britton J W, Frey L C, Hopp J L, Korb P, Koubeissi M Z, Lievens W E, Pestana-Knight E M and Louis E K S 2016 Electroencephalography (EEG): An Introductory Text and

- Atlas of Normal and Abnormal Findings in Adults, Children, and Infants 1st edn (Chicago, IL: American Epilepsy Society)
- Capecci E, Morabito F C, Campolo M, Mammone N, Labate D and Kasabov N 2014 Feasibility study of using the NeuCube spiking neural network architecture for modelling Alzheimer's disease EEG data *Italian Workshop on Neural* Networks (WIRN) 2014 (Salerno)
- Cassani R, Estarellas M, San-Martin R, Fraga F J and Falk T H
 2018 Systematic review on resting-state EEG for Alzheimer's
 disease diagnosis and progression assessment *Dis. Markers*2018 1–26
- Cassani R, Falk T H, Fraga F J, Cecchi M, Moore D K and Anghinah R 2017 Towards automated electroencephalography-based Alzheimer's disease diagnosis using portable low-density devices *Biomed. Signal Process.* Control 33 261–71
- Craik A, He Y and Contreras-Vidal J L 2019 Deep learning for EEG classification tasks: a review *J. Neural. Eng.* 16 1–28
- Dauwels J, Vialatte F and Cichocki A 2010 Diagnosis of Alzheimer's Disease from EEG signals: where are we standing? *Curr. Alzheimers Res.* 7 487–505
- DeTure M A and Dickson D W 2019 The neuropathological diagnosis of Alzheimer's disease *Mol. Degener.* 14 1–18
- Dodge S and Karam L 2017 A study and comparison of human and deep learning recognition performance under visual distortions (arXiv:1705.02498)
- Dukart J, Schroeter M L and Mueller K 2011 Age correction in dementia—matching to a healthy brain *PLoS One* 6 1–9
- Fan M, Yang A C, Fuh J-L and Chou C-A 2018 Topological pattern recognition of severe Alzheimer's disease via regularised supervised learning of EEG complexity Front. Neurosci. 12 1–10
- GBD 2016 Dementia Collaborators 2019 Global, regional, and national burden of Alzheimer's disease and other dementias, 1990–2016: a systematic analysis for the global burden of disease study 2016 *Lancet Neurol.* 18 88–106
- Greco A, Costantino D, Morabito F C and Versaci M 2003 A Morlet wavelet classification technique for ICA filtered SEMG experimental data *Proc. IJCNN 2003* (Portland: IEEE)
- He K, Zhang X, Ren S and Sun J, 2015 Deep residual learning for image recognition 2016 IEEE Conf. on Computer Vision and Pattern Recognition (Las Vegas)
- Hossin M and Sulaiman M N 2015 A review on evaluation metrics for data classification evaluations *Int. J. Data Min. Knowl. Manage. Process* 5 1–11
- Ieracitano C, Mammone N, Bramanti A, Hussain A and Morabito F C 2019 A convolutional neural network approach for classification of dementia stages based on 2D-spectral representation of EEG recordings Neurocomputing 323 96–107
- Ieracitano C, Mammone N, Hussain A and Morabito F C 2020 A novel multi-modal machine learning based approach for automatic classification of EEG recordings in dementia Neural Netw. 123 176–90
- Jack C R Jr et al 2016 A/T/N: an unbiased descriptive classification scheme for Alzheimer disease biomarkers Neurology 87 539–47
- Jack C R Jr et al 2018 NIA-AA research framework: toward a biological definition of Alzheimer's disease Alzheimers Dementia 14 535–62
- Kim D and Kim K 2018 Detection of Early Stage Alzheimer's Disease Using EEG Relative Power with Deep Neural Network IEEE Engineering in Medicine and Biology Society Annual Int. Conf. (Honolulu, Hawaii)
- Kingma D P and Ba J L, 2015 ADAM: a method for stochastic optimisation Int. Conf. on Learning Representations (San Diego)
- Kramer M A, Chang F-L, Cohen M E, Hudson D and Szeri A J 2007 Synchronization measures of the scalp electroencephalogram can discriminate healthy from Alzheimer's subjects Int. J. Neural Syst. 17 1–9

- Krizhevsky A, Sutskever I and Hinton G E 2012 ImageNet classification with deep convolutional Adv. Neural Inf. Process Syst. 25 1–9
- Lilly J M and Olhede S C 2010 On the analytic wavelet transform.

 IEEE Trans. Inf. Theory 56 4135–56
- Mallat S 2009 A Wavelet Tour of Signal Processing: The Sparse Way 3rd edn (Amsterdam: Elsevier)
- Mathworks 2020 Parula (available at: www.mathworks.com/ help/matlab/ref/parula.html) (Accessed 08 November 2020)
- Morabito F C, Campolo M, Ieracitano C, Ebadi J M, Bonanno L, Bramanti A, Salvo S D, Mammone N and Bramanti P, 2016 Deep convolutional neural networks for classification of mild cognitive impaired and Alzheimer's disease patients from scalp EEG recording IEEE 2nd Int. Forum on Research and Technologies for Society and Industry Leveraging a Better Tomorrow (RTSI) (Bologna)
- Olhede S C and Walden A T 2002 Generalized Morse wavelets IEEE Trans. Signal Process. 50 2661–70
- Parra M A, Butler S, McGeown W J, Nicholls L A B and Robertson D J 2019 Globalising strategies to meet global challenges: the case of ageing and dementia J. Glob. Health
- Petersen R C and Knopman D S 2006 MCI is not a clinically useful concept *Int. Psychogeriatrics* 18 393–414
- Petersen R C and Negash S 2008 Mild cognitive impairment: an overview CNS Spectr. 13 45–53

- Prince M, Wimo A, Guerchet M, Ali G-C, Wu Y-T and Prina M 2015 The Global Impact of Dementia: Summary Sheet. (London: Alzheimer's Disease International)
- Rossini P M et al 2020 Early diagnosis of Alzheimer's disease: the role of biomarkers including advanced EEG signal analysis. Report from the IFCN-sponsored panel of experts Clin. Neurophysiol. 131 1287–310
- Ruder S 2017 An overview of gradient descent optimization (arXiv) 1 1–14
- Sabbagh M N, Lue L-F, Fayard D and Shi J 2017 Increasing precision of clinical diagnosis of Alzheimer's disease using a combined algorithm incorporating clinical and novel biomarker data *Neurol. Therapy* 6.83–95
- Stanford Vision Lab 2010 (available at: http://image-net.org/ challenges/LSVRC/2010/) (Accessed 24 December 2019)
- Szegedy C, Liu W, Jia Y, Sermanet P, Reed S, Anguelov D, Erhan D, Vanhoucke V and Rabinovich A 2014 Going deeper with convolutions 2015 IEEE Conf. on Computer Vision and Pattern Recognition (CVPR) Boston
- Weller J and Budson A 2018 Current understanding of Alzheimer's disease diagnosis and treatment F100Research, 7(F1000 Faculty Rev: 1161) pp 1–9
- Zhao Y and He L 2015 Deep learning in the EEG diagnosis of Alzheimer's disease *Computer Vision—ACCV 2014 Workshops* 1st edn, ed C V Jawahar and S Shan (Berlin: Springer) pp 340–53