Variable Interactions in Risk Factors for Dementia

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Abstract—Current estimates predict 1 in 3 people born today will develop dementia, suggesting a major impact on future population health. As such, research needs to connect specialist clinicians, data scientists and the general public. The In-MINDD project seeks to address this through the provision of a Profiler, a socio-technical information system connecting all three groups. The public interact, providing raw data; data scientists develop and refine prediction algorithms; and clinicians use in-built services to inform decisions. Common across these groups are Risk Factors, used for dementia-free survival prediction. Risk interactions could greatly inform prediction but determining these interactions is a problem underpinned by massive numbers of possible combinations. Our research employs a machine learning approach to automatically select best performing hyperparameters for prediction and learns variable interactions in a non-linear survival-analysis paradigm. Demonstrating effectiveness, we evaluate this approach using longitudinal data with a relatively small sample size.

I. INTRODUCTION

In 2014, the UK Alzheimer's Society published a major study on the social and economic impact of dementia in the UK. The report projected that 850,000 people would be living with dementia in the UK by 2015, with associated costs of £26 billion a year [8]. Furthermore, a European study published in 2013 [7] cited the average rate of dementia across 28 countries to be 1.55% of the population. These studies point to dementia as one of the greatest health risks to society today. As a result, systems are required that connect experts from different disciplines to each other, as well as to their target group in society. The In-MINDD project [4], delivers this through the development of the In-MINDD Profiler, a sociotechnical information system which requires multi-disciplinary collaboration between experts in the field of dementia, IT researchers and support staff in order to provision members of society most at risk with state-of-the-art tools for dementia prediction and mitigation.

A. The In-MINDD System

The goal of the In-MINDD project is to develop and validate online tools for middle-aged (40-60) individuals which assess lifestyle factors and how these affect long term brain health, provide personalised strategies for adopting a brain healthy lifestyle and present online supports to help participants implement and adhere to positive change in order to lower the risk of developing dementia. This requires knowledge building

and knowledge sharing, with system-driven decision making, across heterogeneous cultures and health systems.

The development and validation of an online dementia assessment tool requires close collaboration of clinical dementia experts and data scientists. The first element of developing this online system required the definition of a common vocabulary or ontology, in order to share knowledge between relevant stakeholders [18], [19]. Next a multi-factorial model of those elements that pertained to dementia-risk in middle-aged individuals was developed by dementia experts via a systematic literature review [6] and shared with the data scientists via the previously defined system ontology [18], [19]. The system ontology connects the IT and clinical researchers and allows both parties to validate the model - data scientists via the latest machine learning techniques and clinical researchers through more traditional statistical approaches. The result of this work was incorporated into a prototype model for inclusion in the *Profiler* which was developed by the computing researchers. The Profiler presents a user with a series of questions, evaluates the answers to these and hence, provides a brain-health score. Subsequently, the Profiler presents the participant with a series of goals to improve their brain-health and mitigate against those factors from which they are most at risk. This is fundamentally different to systems such as that described in [17] where data is automatically generated in a uniform manner by purpose-built devices. The In-MINDD project is now testing this prototype model to assess if a 6 month intervention with the online profiler and support environment can improve the level of dementia risk in those most susceptible [14].

The previously defined information-system allows iterations and inclusions of improvements in the model, which is the focus of this work. The aim is to improve the predictive power of the model, through machine learning algorithms that offer deeper analyses compared to traditional statistical approaches and can identify latent classes over the course of learning, resulting in candidate risk-factor interactions. These interactions cannot be assessed by data scientists, instead results will be provided to dementia experts for an informed clinical evaluation.

B. Problem Statement and Motivation

As stated in Section I-A, early in the In-MINDD project, dementia researchers identified the 14 most prevalent risk factors from published research which can be used to measure a person's risk of developing dementia [6]. With the exception of age, gender, and years of education, the remaining 11 risk factors are modifiable, meaning if clinical researchers can determine strong risk factors and have the person involved modify their behaviour, they can reduce their likelihood of developing dementia. These modifiable risk factors were identified as cognitive activity, physical activity, mid-life obesity, diabetes, moderate alcohol use (protective), smoking, depression, mid-life hypertension, cholesterol, cardiovascular disease and kidney disease. It is necessary for researchers to determine how these variables interact in order to build an optimum predictive model for identifying and extending life without dementia in middle-aged individuals. For clinical researchers, finding these interactions is a significant task using statistical software packages due to the high number of permutations involved. There is also a need to develop an a-priori hypothesis of interactions before testing these in a relevant dataset.

This motivates the need to obtain a deeper understanding of the interactions between risk factors. Neural networks provide possibilities to measure these interactions through examining the *latent* features found in their hidden layers and the weights that combine the visible variables to make up these hidden features. By *latent* features, we mean those *abstract* variables that combine more fine-grained variables into higher-level attributes we are trying to determine. The weights which are the coefficients for the input features are the outputs of our experiments, described in Section IV. The main problem dealt with here is detecting the appropriate latent variables from the very high number of potential multi-attribute combinations that exist and adopting a non-linear dementia survival analysis in the process.

The ideal dataset for this type of research is a longitudinal study which tracks behaviour over a number of years, preferably for those in middle-age. The dataset used was the Maastricht ageing study (MAAS dataset) [3] which was generated from a longitudinal, prospective cohort study which recorded lifestyle and biometric data on middle-aged individuals through 86 types of tests at 3 year intervals over more than 12 years.

C. Contribution

Consider that we have a system in place that connects clinical researchers with participants in a study trial. For this system the data scientist must also access further study trial data to provide a deeper analysis and determine how risk factors interact in the *highest performing* predictive model, subsequently presenting these risk interactions to the clinical researchers. Our primary contribution is in the construction of neural network models for a set of seven tests to determine the best performing of 14 different predictive models for survival analysis. Each model involved a separate experimental run of 512 test configurations, in each case risk interactions were

learned and the weights for each combination of interactions were recorded, to enable the optimal configuration to be found.

D. Paper Structure

The remainder of this paper is structured as follows: in Section II, we describe our approach as a 2-step process of parameter set-up and optimisation; in Section III, we discuss how our system is configured and the different dataset types that served as input; our evaluation, results and analysis are described in Section IV; in Section V, we contrast our work with similar efforts in this area while in Section VI, we provide some conclusions.

II. APPROACH

Two aims are to be solved with our approach: to determine multi-variate feature interactions in dementia survival analysis; and to adopt a non-linear approach as many participants in the Maas dataset have similar attributes but different classifications. Non-linear analyses better model these participant differences and thus, provide superior accuracy in outcome predictions. We use a Multi-Layer Perceptron (MLP) to perform supervised classification and our approach tackles one of the 10 challenges [21] in data mining research: data mining for biological problems.

Supervised classification is a machine learning paradigm which trains an algorithm on data in order to predict a particular outcome; in our case, we perform multi-class classification to distinguish between survival cases and those who encountered *dementia*, *death* or *censorship* (study drop-out) in the dataset. An MLP is an artificial neural network (ANN) that learns abstract features from the data. We hypothesise that these abstract features model the complex dementia risk factor interactions to better predict the outcomes in question. Our approach builds on the outline platform developed in [16] [15], with a new 2-step process to deliver high performing prediction models.

A. Step 1: Determine Hyper-Parameter Settings

The broad purpose of Step 1 is to determine the hyper-parameters for all experiments. Before this step, we have a set of bounds for each hyper-parameter and after this step, all hyper-parameter configurations will be determined (Section III-B elaborates on this). Hyper-parameters are algorithm tuning-parameters that influence how model weights are learned in training according to the optimisation procedure outlined in III-B. The number of hidden layer nodes and learning rate (parameter for magnitude of weight updates) are examples. This must be completed before any form of model optimisation can begin.

B. Step 2: Hyper-Parameter Optimisation

There are five separate procedures in Step 2 that are repeated for *each* of the 256 trials, for each experiment combination described in Section III-A.

- 1) Initialise Architecture: This process uses a hyperparameter configuration from Step 1 to generate the appropriate algorithm configuration and architecture, setting the number of nodes and layers, inputs and outputs, associating all relevant hyper-parameters to the configuration and initialising relevant weights and biases.
- 2) Construct Hypothesis: The hypothesis_function combines the model weights with each sample of dataset variables and incorporates the appropriate activation and classification functions that allow for making predictions. We used an MLP with one hidden layer.

$$z_j^{(0)} = b_j^{(0)} + \sum_{i=1}^n x_i w_{ij}^{(0)}$$
 (1)

$$P(n_j = 1|x, \theta^{(0)}) = a_j = \frac{1}{1 + e^{-z_j^{(0)}}}$$
 (2)

$$z_k^{(1)} = \sum_{j=1}^m a_j w_{jk}^{(1)} \tag{3}$$

$$P(y_k = 1|x, \theta) = \frac{e^{z_k^{(1)}}}{\sum\limits_{k=1}^{K} e^{z_k^{(1)}}}$$
(4)

Equations 1 and 2 are the functions between the input to hidden layers and Equations 3 and 4 are the hidden to classification layer functions. The parameters $\theta^{(0)}$ for the input to hidden layers are $\{b^{(0)}, W^{(0)}\}\$ the bias and weights respectively. Equivalently, $\{b^{(1)}, W^{(1)}\}\$ are the parameters for the hidden to output layers $\theta^{(1)}$. The complete set of these parameters $\{\theta^{(0)}, \theta^{(1)}\}\$ are denoted θ . x is the set of input feature values from $1 \dots m$ where m is the total number of features. n is the set of hidden nodes and a are the activations for these nodes from $1 \dots o$, the total number of nodes in the hidden layer. k refers to a particular classification in the range of output classes 1...K. For Equation 1, weights are multiplied by the input features to give a linear combination before entering the non-linear logistic sigmoid activation function -Equation 2 - to compute the probabilities of a hidden node n being 'on' or 'off' - also known as a node's activation energy. This is the point where we hypothesise interactions are found for the input variables, as during training, each hidden node learns different combinations of features with different sets of weights. The same process is carried out for the hidden to classification layer. The hidden activations a are combined with another set of weights $W^{(1)}$ - Equation 3 - before being input into the softmax activation function (Equation 4), which computes a probability for each of the Kclasses, and the sample is assigned the class for which it has the highest probability. Probabilities of a softmax classifier are not independent as they aggregate to 1, this function is used as dementia, survival, death and censorship are nonindependent outcomes.

3) Build Cost: The cost_function evaluates how well the hypothesis function performs in comparison with the ground truth. We construct a symbolic function to compute the negative log of the probability calculated by the model for the actual class, given the input and model parameters. Known as the negative log likelihood (NLL) and shown in Equation 5; y_k is the ground truth of either: survival, dementia, death or censorship, x is the set of input variables values and θ is the set of model parameters. The output of this function serves as input to the train function where the cost is lowered, giving better predictions for survival analysis. We also add regularisation to this function (term after the addition sign) which penalises large weight values, smoothing the error function and thus, allowing us to find better local minima or ideally, global minima. This will further increase the accuracy for survival analysis classifications.

$$-logP(y_k|x,\theta) + \lambda\theta^2 \tag{5}$$

- 4) Construct Model: At this point, we construct the symbolic functions: update_parameters, train, validate, test and predict. The update function calculates the derivative of the model parameters with respect to the cost to be subtracted from the current weights (weighted by the learning rate α , a hyper-parameter for deciding the magnitude of parameter updates), lowering the cost and enabling better predictions.
- 5) Train Model: This is the optimisation method for the cost function. We use mini-batch stochastic gradient descent (MSGD) with early stopping for this optimisation process. MSGD takes small chunks of the dataset and updates the parameters after computing the cost for the current weights. It is computationally more efficient than updating based on a calculation using the entire dataset (batch gradient descent). In non-convex, non-smooth error functions it also enables the optimisation function to escape poor local minima (which lead to poor predictions of survival) and find better minima and potentially the global optimum for improved predictions. Early stopping helps to avoid over-fitting by adding regular evaluation of the model's performance on held-out validation data and terminating the procedure once the validation performance stops improving. An iteration of MSGD takes this small chunk of the dataset, calculates classification probabilities, then the cost of these predictions before lowering the cost through the update function described in the previous process.

After both steps have been completed, the system then queries a persistent NoSQL store, ordering the results by the best performance on the validation set, and selecting the top performing model for predictions.

III. EXPERIMENTAL SET-UP

The risk factors listed in our introduction are shown in more detail in Table I. After records with missing values were removed, the dataset comprised 840 samples by 25 features, where each sample contained data on an individual aged 50 or over, at baseline. In this section, we outline the mix of 7

TABLE I FEATURES USED IN EXPERIMENTS

No.	Feature	Data Type	Description	Weight
1	p_id	numerical	Anonymised participant ID	n/a
2	age	continuous	Age of participant in yrs.	n/a
3	gender	binary	0 = man / 1 = woman	n/a
4	age_risk	continuous	Dementia risk from age and gender	n/a
5 a	low_ed	binary	Less than 8 yrs. formal education	n/a
5 b	yrs_ed	discrete	Number of years in formal education	n/a
5 c	ed_risk	continuous	Dementia risk from years ed.	n/a
6 a	cog_active	binary	Participant cognitively active @ 50+	-3.2
6 b	hrs_cog_active	discrete	Hrs. cognitively active/day	n/a
7 a	phys_inact	binary	Participant physically inactive @ 50+	1.1
7 b	hrs_phys_inact	discrete	Hrs. physically active/day	n/a
8 a	obese	binary	Participant obese at baseline	1.6
8 b	bmi	continuous	Body mass index of participant	n/a
9 a	mod_alcohol	binary	Low/moderate alcohol consumption	-1
9 b	u_alcohol	discrete	Units alcohol consumed/week	n/a
10	smokes	binary	Participant smokes	1.5
11 a	depressed	binary	Participant is depressed	2.1
11 b	depression_score	discrete	Score on SCL depression scale	n/a
12	hypertension	binary	Has high blood pressure	1.6
13	cholesterol	binary	Has high cholesterol	1.4
14	cvd	binary	Has cardiovascular disorder	1.0
15	kidney	binary	Has kidney disorder	1.1
16	dementia	binary	Developed dementia	n/a
17	died	binary	Participant died during study	n/a
18	in_study_12	binary	Participant was in the 12 yr. follow-up	n/a
19	classes	(derived) categorical	0: survived - no dementia/death; 1: developed dementia; 2: died, no dementia 3: censored - didn't complete study	n/a

different input combinations provided to our modelling process chosen to test performance of the modifiable risk factors.

A. Input Combinations

In Table I, we present our extended list of risk factors, either in binary, discrete or continuous form to enable the construction of different models. Relative risk weights are either neutral (set to 1) or assigned a value taken from an meta-analysis of the literature performed by our colleagues in Maastricht [6]. To test the validity of risk factors and learn the relevant interactions, modifiable and non-modifiable combinations were used as input to experiments.

Seven experimental set-ups were constructed, where experiments 1 - 3 were baseline tests without relative risk weights and experiments 4 - 7 incorporated those weights from Table I. For each experiment, 512 trials were run: 256 on standardised data and the same number with unstandardised data. For reporting purposes, it was necessary to provide 2 labels for each experiment. For example, B1 represents Binary Baseline Test 1 and B1-sd represents same baseline test (and inputs) but with a standardised dataset. The dataset was split using a 70:20:10 ratio for training, validation and testing respectively. Data subsets were selected randomly for every test run within each experiment. All experiments but the first (where there was one output node), had 4 output nodes, 1 for each possible class.

1) **B1** and **B1-sd**: First *binary-baseline* experiment on non (B1) and standardised (B1-sd) data.

- **Inputs**: 15 features; unweighted; outcome = dementia (16); 11 binary modifiable factors; adjusted for age (2), gender (3) and education (5a).
- B2 and B2-sd: Second binary-baseline experiment on non (B2) and standardised (B2-sd) datasets.
 Inputs: 15 features; unweighted; outcome = classes (19); 11 binary modifiable factors; adjusted for age (2),
- CB and CB-sd: Continuous-baseline experiments on non (CB) and standardised (CB-sd) datasets.
 Inputs: 15 features; unweighted; outcome = classes (19); 6 binary and 5 continuous modifiable factors; adjusted for age (2) and gender (3) and education (5a).

gender (3) and education (5a).

- 4) BW1 and BW1-sd. first binary-weighted experiment on non (BW1) and standardised (BW1-sd) datasets.
 Inputs: 15 features; weighted; outcome = classes (19); 11 binary modifiable factors; adjusted for age (2), gender (3) and education (5a).
- 5) **BW2** and **BW2-sd**: second *binary-weighted* experiment on non (BW2) and standardised (BW2-sd) datasets. **Inputs**: 15 features; weighted; outcome = classes (19); 11 binary modifiable factors + education risk (5c); adjusted for age (2) and gender (3).
- 6) BW3 and BW3-sd: third binary-weighted experiment on non (BW3) and standardised (BW3-sd) data.
 Inputs: 14 features; weighted; outcome = classes (19); 11 binary modifiable factors + weighted binary modifiable features + age risk (4) and education risk (5c).
- 7) CW and CW-sd: continuous-weighted experiments on

TABLE II HYPER PARAMETERS AND BOUNDS

Hyper-parameter	Bounds (low, high)	Description
batch_size	(1, 50)	number of samples in each mini-batch update
n_hidden_nodes	(2, 20)	number of nodes in the hidden layer
learning_rate α	(0.0001, 0.3)	magnitude of weight updates per iteration
regularisation λ	(0.0001, 0.1)	degree of penalisation for large weights
max_epochs	(100, 2000)	max iterations through the dataset
max_iterations	(5000, 100000)	max number of weight updates performed

non (CW) and standardised (CW-sd) data.

Inputs: 14 features; weighted; outcome = classes; 6 binary and 5 continuous modifiable factors + age risk (4) and education risk (5c).

Each model was optimised on training set; those models with the best performance on the held-out validation data were selected as the best performing and as such had the optimum hyper-parameter configuration; finally the performance of those selected to have the optimal hyper-parameter configurations were evaluated on the further held-out test set, independent to the training and validation data. This ensures bias is not present in the model as it is not evaluated on the data it was trained on, mitigating against over-fitting and leading to better generalisation.

B. Model and Hyper-Parameter Optimisation Procedures

As previously mentioned in Section II MSGD in conjunction with early stopping was the model optimisation procedure for all models. Higher level termination parameters: max_epochs and max_iterations where the bounds can be seen in Table II were also in place if early stopping did not terminate the training. Hyper-parameters were optimised through random search consisting of 256 trials for each experiment. Random search has been shown to outperform grid-search [2] and the number of trials were chosen as per the guidelines also described in [2]. For each trial a configuration was drawn uniformly and randomly according to the bounds shown in Table II.

IV. EVALUATION AND ANALYSIS

All experiments were run on a Dell Optiplex 790 running 64-bit Windows 10 Home with an Intel Core i7-2600 quadcore 3.40 GHz CPU and 16.0GB of RAM. The code for the experiments was developed in Python using the Enthought Canopy (1.5.4.3105) distribution of 64-bit Python 2.7.9 and developed in PyCharm 4.5 IDE. The code makes use of NumPy 1.9.2, Theano 0.7.0 and their dependencies. The AUC and χ^2 test of independence were calculated in JMP Version 11. The time taken to train a total of 3,584 models for unstandardised and standardised baseline, modifiable, modifiable with education and modifiable with age and education risk and continuous tests was 14.79 hours.

A. Evaluation Metrics

The evaluation measures used, where tp (true positives) are the sum of values predicted positive that are positive, tn (true negatives) are the sum of values predicted negative that are negative are as follows:

- **Precision**. The positive predictive value which is a proportion of values predicted positive that are actually positive: $\frac{tp}{n_predicted_positive}$.
- **Recall**. The *sensitivityltrue positive rate* (TPR) which is the proportion of true positives to the actual number of positives in the dataset: $\frac{tp}{n_actual_positive}$.
- True Negative Rate (TNR): Also called *specificity*, the rate at which the classifier identifies actual negative cases: $\frac{tn}{n_actual_negative}$.
- F1 Score. This is a harmonic *mean* of precision and recall: $2 * \frac{precision*recall}{precision+recall}$.
- Accuracy. The proportion of correctly classified instances: $\frac{tp+tn}{n_predictions}$.
- AUROC. Using varying thresholds, this function evaluates True Positive vs False Positive trade-off.
- $\mathbf{p} > \chi^2$. This chi-squared test is used to show that model predictions are significant where the rejection threshold is p > 0.05.

B. Results

In this section, we provide the results of our main research goal: the determination of those risk factors and their combinations which best predict *survival* in the dementia study. Our evaluation will focus on our use of a series of statistical tests to determine the highest performing predictive models. We can only present the results of interactions found as in reality, these interactions can only be evaluated by our research collaborators, dementia specialists at the University of Maastricht and this work is currently in progress.

1) Survival Analysis Results: Our analysis focuses on the survived classification versus all other hazards, grouping predictions of death, dementia and censorship. Models performed well when classifying survival and censorship, gave moderate results for death but poor results when predicting dementia. This negative performance is due to class imbalance, while having sufficient numbers of survival, death and censorship cases (roughly 400 censored samples) numbers for dementia (60) were too low.

Of the baseline experiments CB-sd performed best in the measures of F1, Accuracy, AUROC, and Significance with values of 0.6888, 0.6667, 0.7326 and 0.0002 respectively. CB performed best in the Precision and TNR measures with scores of 0.6296 and 0.7872 while B2 performed best for recall at

TABLE III
ANALYSIS SUMMARY STATISTICS

Summary Statistics for Survival Analysis							
	Precision	Recall(TPR)	TNR	F1	Accuracy	^a AUROC	$^{ m bc}{f p}>\chi^{f 2}$
B2-sd	0.4727	0.8125	0.4423	0.5977	0.5833	0.6922	0.0187
СВ	0.6296	0.4595	0.7872	0.53125	0.6429	0.7021	0.0024
CB-sd	0.6078	0.7949	0.5556	0.6888	0.6667	0.7326*	0.0002
BW1	0.6364	0.4118	0.84	0.5	0.6667	0.6921*	0.0221
BW1-sd	0.48	0.75	0.5	0.5854	0.5952	0.6797	0.0237
BW2-sd	0.6383	0.75	0.6136	0.6897	0.6786	0.7016*	0.003
BW3	0.5849	0.9118	0.56	0.7126	0.7024	0.7635	< 0.0001
BW3-sd	0.4642	0.7429	0.3878	0.5714	0.5357	n/a	0.33
CW	0.4262	0.9286	0.375	0.5843	0.5595	0.7299	0.0043
CW-sd	0.5263	0.9091	0.4706	0.6667	0.6429	0.7693	0.0002

- ^a Calculated by fitting logistic model (LM) between actual vs. predicted classifications.
- *LM parameter estimates unstable, due to low instance of a class in test set.
- b Degrees of freedom:3 in all but BW2-sd and BW3-sd, DOF: 6.
- ^c Findings corroborated with LM as some instances of χ^2 had cell counts < 5.

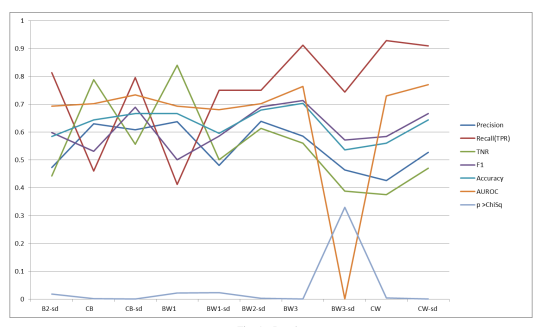


Fig. 1. Results

0.8125. For all experiments, BW2-sd had the highest Precision of 0.6383, CW had the highest Recall of 0.9286, BW1 had the best true negative rate of 0.84, BW3 had the highest F1, Accuracy and Significance at 0.7126, 0.7024 and < 0.0001 respectively. CW-sd had the highest AUROC of 0.7693. B1 and B1-sd, using only *dementia* diagnosis as outcome, predicted every participant as 'not dementia' and were therefore omitted, a result expected due to the previously described class imbalance. BW2 is also omitted from the results table due to poor performance. In summary, the performance of weighted experiments was superior to those of baseline experiments. It is important to note that the first 3 measures (Precision, Recall and TNR) are individual scores while the next 3 (F1,

Accuracy and AUROC) are different aggregations of these first 3 measures and as such, our working assumption is that the latter 3 measures carry more significance. Across the 3 aggregate measures, we feel that AUROC offers most significance as it provides the model with the best trade-off between true and false positives. This is followed by Accuracy as it delivers the model with the highest number of correct classifications.

2) Risk Factor Interactions: It is from these high performing models presented in Table III that we can extract the variable interactions illustrated in Table IV. These interactions are found from the top-3 runs of the BW3 experiment discussed in Section III-A and illustrate combinations of risks weighted

TABLE IV
TOP PERFORMING RISK FACTOR INTERACTIONS

	BW3 Interactions - 3 Top Ranked Models					
BW3- RankA - Trial 160 of 256						
Node	Weight/Rank	Positive Interactions	Negative values			
H2	1st: 0.6411	cog_act	age_risk			
Н3	2nd: 0.0945	educ_risk, phys_inact, depressed	diabetes, di_alcohol, hyperT			
H1	3rd: 0.0837	educ_risk, cog_act, obese, diabetes, hyperT	phys_inact, di_alcohol			
Н6	4th: 0.0667	cog_act	diabetes, di_alcohol, smokes, cholesterol, kidney			
H4	5th: 0.0562	phys_inact, smokes, cholesterol, cvd	educ_risk, diabetes, depressed			
H5	6th: 0.0549	educ_risk, obese, smokes	cvd, di_alcohol			
BW3-F	BW3-RankB - Trial 249 of 256					
H7	1st: 1.3669	educ_risk, phys_inact, hyperT, cholesterol	age_risk, kidney			
H5	2nd: 0.8725	kidney	cog_act, hypertension, di_alcohol			
H8	3rd: 0.5933	diabetes, depressed, cholesterol	age_risk, educ_risk, phys_inact, hyperT			
H4	4th: 0.2600	educ_risk, obese, hyperT	depressed			
H1	5th: 0.0485	age_risk, diabetes, depressed, cholesterol	cog_act, obese			
Н6	6th: -0.0425	cog_act, obese, depressed	di_alcohol, cvd			
Н3	7th: -0.2031	educ_risk, cog_act, phys_inact, diabetes, smokes,	age_risk			
		hyperT				
H2	8th: -0.2654	diabetes, smokes, hyperT	cog_act, depressed, cvd			
BW3-RankC - Trial 44 of 256						
Н3	1st: 0.7534	obese, smokes	age_risk			
Н6	2nd: 0.7006	di_alcohol	age_risk			
H7	3rd: 0.0042	age_risk, educ_risk, smokes, hyperT	obese, di_alcohol			
H5	4th: 0.0037	age_risk, educ_risk, depressed, hyperT	cog_act, di_alcohol			
H10	5th: 0.0030	age_risk, educ_risk, hyperT	cog_act			
H8	6th: 0.0024	age_risk, educ_risk, depressed	di_alochol, cvd			
H4	7th: 0.0020	age_risk, educ_risk, hyperT, cvd	diabetes, smokes			
H2	8th: 0.0019	educ_risk, depressed	di_alcohol, kidney			
H9	9th: 0.0016	age_risk, smokes	diabetes, di_alcohol			
H1	10th: 0.0015	age_risk, educ_risk, hyperT, cholesterol	cog_act			

above average for the negative and positive weights for each node, as shown in Table IV. Put simply, the top interaction was that which occurred most often: cvd (cardiovascular disease) and $di_alcohol$, occurring in all three models. Other interactions like diabetes and smokes occur often together but also in the context of other variables so need further validation. For the remainder of this section, we focus on the discussion and analysis of the approach and results which delivered these interactions, as clusters of Risk Interactions can only be evaluated by clinical researchers.

C. Discussion and Analysis

BW3-sd aside, all models were found to be significant under a chi-squared test of independence. Significance of BW3 performed poorly at 0.33 meaning predictions achieved could be due to chance. We conclude poor performance of BW3-sd is likely due to the standardisation process. Weighted features had continuous values, but further analysis shows they are technically ordinal - either 0 or the risk weight - hence, do not follow a normal distribution. Feature scaling (subtracting the min value and dividing by the range) instead of standardising (subtracting the mean and dividing by the standard deviation) would provide greater benefit in this context and will be explored in future work. As a result, all BW models apart from BW2 performed better when unstandardised, in contrast to continuous models which performed better when standardised. The opposite effect found for BW2 is likely due to the continuous age which is on a much wider scale than the other

features and thus, we conclude that standardising the dataset improved performance.

Models built from continuous data performed better than those only including binary features, where binary features pertain to the participant either having or not having a risk factor. Weighted models also performed better than their nonweighted counterparts. Binary models only begin to outperform the continuous baseline when education risk is added in BW2. Continuous data provides far more information than binary with the result that CB outperforms BW1 until education and age/sex risks (significant predictors of dementia) are added in BW2 and BW3 respectively. The negative performance B2 shows risk weightings have a significant role to play in survival analysis in the absence of continuous data. BW2 performs better than BW1 and likewise BW3 performs better than BW2, with increasing AUROC for each. Further information is given to each model, first incorporating education risk and subsequently then incorporating age/sex risk. The key conclusion here is that continuous data and relative risks improve the predictive accuracy of the model. This is important, as currently the In-MINDD model developed by the clinicians only contains dichotomous has/does not have variables for each particular risk factor.

When considering the aggregate measures, BW3 performs best with highest F1, Accuracy and Significance, the only model with F1 and Accuracy > 0.7. Therefore, BW3 was the most accurate classifier, correctly classifying the highest number of instances (Accuracy) and performing best at identi-

fying and predicting positive instances (F1). The significance $(p > \chi^2)$ of < 0.0001 shows that BW3 achieved predictions that are the most significant and the least likely to be due to chance. CW-sd achieved the best AUROC score of 0.7693 and a close to best significance level of 0.0002. Thus, we observe that CW-sd had the greatest trade-off between true and false positives, a very important factor in survival analysis. Also CW-sd predictions are *very* unlikely due to chance (0.0002). The difference in AUROC for BW3 and CW-sd is very small at 0.0058 or 0.58%. BW3's greater performance is likely due to relative risk weights being designed for dichotomous risk factor cut-offs rather than continuous scores. On the other hand, continuous data was present for only 5 out of 11 risk factors in the dataset and if continuous measures were used for all features then continuous models would likely surpass binary models on all scores.

CW has the highest TPR of 0.9286, meaning out of the instances it predicted positive, >92% were *actually* positive - an excellent score - but offset by poor Precision of < 0.5, worse than the binary baseline. Using the positive values in the test set, it identified <50% which is reflected in its poor F1 score of 0.5843 and lower level of significance. We conclude this is likely due to lack of standardisation. BW1 has highest true negative rate at 0.84, but has a poor TPR of < 0.5. Although performing better than other models in predicting hazards (negative cases), the low TPR shows us this model could not perform well in modelling *survival* cases. This is undesirable as <50% of those it predicted positive, were actually positive, with many of these as false positives.

In summary, the important conclusion from the results is that even when the relevant risk weights are omitted and only dichotomous values are present, the model is still significantly predicting *survival*, showing that these particular factors are valid in the prediction of *survival* versus other hazards. Furthermore, models which include relative risk weights from the literature experience an improvement in predictive accuracy. Our results show that continuous data can again improve the model. Although the final binary model outperforms the continuous model in some measures, as continuous data is only present for 5 of the 11 inputs we are confident that when continuous measures are added for all inputs that it will outperform all binary models. With respect to input interactions we have successfully presented those automatically learned by the MLP.

V. RELATED RESEARCH

Although there is potential for artificial neural networks (ANN) in survival analysis, it is not in widespread use for clinical applications. Neural networks in the context of health analytics have been shown to be at worst, on par with traditional regression approaches [16], [10] and at best outperform these methods [20]. Health benefits accrued from using ANNs in medical interventions have been shown to have significant impact in their application, where notable examples include cervical cytology and the early detection of acute myocardial infarction (heart attack) [9]. Therefore, the motivation for

the application of these methods to the field of dementia is significant.

The majority of ANN survival analyses focus on postsurgery patient mortality or relapse, in areas such as pancreatic cancer [1], breast cancer [10] and liver cancer [20]. ANNs have been applied to dementia, [11], [12], but as far as we can identify from the literature, none adopt a survival analysis paradigm with respect to dementia as explored in this work. Furthermore, all discussed research compares ANNs to methods such as Cox's regression or non-neural machine learning algorithms. This research evaluates ANNs built on modifiable risk factors with those including non-modifiable factors. We add to the research through the comparison of non-modifiable and modifiable risk factor models for dementia and by using a survival analysis paradigm. Finally, none of the aforementioned works analyse latent features learned in the hidden layers to extract input features interactions, which are clinically relevant, as these model risk factor interactions and increase a classifier's predictive power. This insight could provide further understanding to clinicians as to why ANNs would better model survival and can perhaps identify novel interactions from the data which might not be attainable from traditional means. Variable interactions learned during an ANN survival analysis are analysed in [5], but again this in the context of surgery survival and not dementia, which is a further focus of this work.

Exploring [11] further, two (MLP and radial basis function) neural nets are found to perform worse than Support Vector Machines (highest), Random Forests and Linear Discriminant Analyses. ANNs are complex to train, so there are several likely reasons for poor performance, as discussed by the authors. Primarily, ANNs are highly sensitive to tuning (hyper) parameters used to initialise and inform the training procedure, vastly effecting model quality. [11] only optimise the number of hidden nodes, where all other settings were those "commonly used in data mining applications" i.e. not fitted to the data. Furthermore, [11] use a grid-search optimisation procedure, in contrast to random search for all hyperparameters - employed by this research - a methodology shown to outperform grid-search [2]. Finally, their cross-validation approach is sub-optimal for hyper-parameters evaluation and choice. Their strategy essentially splits the data in two - a portion for training and evaluating hyper-parameters and a held-out cross-validation set for evaluating the classifier's final performance. When data on which the model is trained is also used to evaluate hyper-parameter performance, this effectively over-fits the hyper-parameters to the training data. Instead, a three-way split is advised, building the model on training data, evaluating hyper-parameters on held-out cross validation data and an independent test set to evaluate classifier performance on completely unseen data. The latter approach adopted in this work, as well as an automated random search method for hyper-parameter optimisation which has been shown to improve a classifier's predictive power [2]. Furthermore, [11] do not apply ANNs in the context of dementia survival, but instead for dementia classification and none of the latent

features learned by the network were analysed.

In [13], the authors model latent classes (behaviours) in dementia analysis, showing the approaches relevance and according to the authors, the sole study to do so for dementia. The analysis attempts to determine sub-behaviours across six domains: church-attendance; smoking; alcohol use; social interaction; and physical exercise. LCA requires a number of further steps after the identification of classes. First, LCA is applied and tested for a variety of possible latent class numbers (similar to our number of nodes), then multi-nominal regression is applied to assign a sample to the relevant class (behavioural sub-category) before finally running regression again (for each identified class) to evaluate survival probabilities. In our work we use an MLP to automate this process of learning latent variables. An MLP essentially aggregates all steps and where latent variables are identified, samples activate relevant latent variables (hidden nodes) and classification probabilities are identified, all during the course of training. In addition we have extracted non-linear continuous interactions between sub-categories with the MLP in contrast to the linear LCA, thus providing more representative abstract classes.

VI. CONCLUSIONS AND FUTURE WORK

Dementia is a major health concern with the projected numbers likely to consume countries' health budgets in years to come. While many specialists in this field together with clinical researchers are involved in longitudinal studies in an effort to determine the prominent interactions between risk factors, this is a difficult task, given the numbers of risk factors, the permutations possible, and the application of weightings to each risk factor. While the In-MINDD project connects clinical researchers to trial participants through a cloud-based profiler, it is only by connecting the third stakeholder, the data scientist, that we can deliver greater impact.

In this research, we describe our neural network service which using a series of validation functions, supports the profiler by determining the set of interacting risk factors in the highest performing prediction models with varying degrees of modifiable and non-modifiable risk-factors. As part of our evaluation, we present the results of experiments for 10 predictive models, each of which had 512 test configurations across 7 parameters to determine the highest performing model. Our goal was two-fold, first build a non-linear predictive model of dementia survival with a Multi-Layer Perceptron and second examine its hidden features to establish clusters of risk factors which we could then provide to clinical research colleagues for evaluation. This builds on existing research as although ANNs have been applied in a survival analysis paradigm before - as far as we are aware - they have not been applied to dementia survival analysis. Furthermore, in the dementia context the hidden layers have not been analysed to extract candidate clusters of risk factors, which has been presented in this work. Finally, this work adds a procedure which automates the optimisation and selection of the MLP's hyper-parameters using random search in contrast to previous works which often did not optimise all hyper-parameters, or did so in a

manual grid-search fashion. The result of this is a predictive model which significantly predicts dementia survival and sets of candidate risk factor interaction which we will provide to dementia experts who can evaluate these from a clinical standpoint.

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