An extensive experimental setup of ML models for multivariable regression

Veronica Pistolesi, Francesca Poli Computer Science Master Degree, AI curriculum. v.pistolesi6@studenti.unipi.it, f.poli12@studenti.unipi.it

ML course 654AA, Academic Year: 2022/2023 Date: 01/06/2023

Type of project: **B**

Abstract

This project tests the effectiveness of multiple Machine Learning models and libraries on a regression task, namely: Decision Tree, Extra Tree, K-Nearest Neighbors, Support Vector Machine, Random Forest, Bagging and MLP. At the end, we implemented an ensemble model with the top five performing models.

1 Introduction

Our project's aim is to compare the results of different Machine Learning models and libraries applied to a regression task on the ML-CUP22 dataset. Our experiments cover the following models: Decision Tree, Extra Tree, K-Nearest Neighbors, Support Vector Machine, Random Forest, Bagging and MLP. The goal of this comparison is to identify the top models for this task and to try to confirm the expected results given the qualities of each model. Furthermore, an ensemble model consisting of the top five models was selected to be presented to the CUP competition. Some of the previous models were also implemented to carry out a classification task with MONK dataset [3].

2 Method

All the experiments were executed on a laptop with an Intel Core i7-1255U processor, belonging to the x64 architecture, by using Visual Studio Code. The chosen models for the regression task are: DecisionTreeRegressor and ExtraTreeRegressor for the class TREE; SupportVectorRegressor for the class SVM; KNeighborsRegressor for the class NEIGHBORS; RandomForestRegressor, and BaggingRegressor for the class ENSEMBLE; three MLP models for the class NN. The classes were implemented by using scikit-learn [5] library but for the neural networks SciKeras [2] and Skorch [6] were also used. At the beginning, we extracted a development set (70% of the data) and an

internal hold-out test set (30% of the data) from the ML-CUP22 dataset and the input features were normalized with the standard scaler transformation. Then, each model's hyperparameters were subjected to a grid search, with 5-fold cross-validation being used for each configuration of hyperparameters with scikit-learn's GridSearchCV. Models' results were ranked and analyzed according to the Mean Euclidean Error (MEE) score on the validation set (20% of the data), before proceeding with a final model assessment phase. The parameter configurations tested for each model and more details on the implementation choices are presented in Section 3.2.

3 Experiments

3.1 Monk Results

The following steps summarize the experimental setting we adopted for this task:

- 1. One hot encoding to all the input features
- 2. Choice of four model classes for comparison (TREE, SVM, NEIGHBORS, and NN)
- 3. Grid searches on each model's hyperparameters, with stratified 5-fold cross-validation applied to each configuration.

3.1.1 Tree: DecisionTreeClassifier (scikit-learn)

\mathbf{M}	сср	class	criterion	max	max	min	min	min	ACC
	alpha	weight		depth	leaf	samples	samples	weight	(TR/TS)
					nodes	leaf	\mathbf{split}	fraction	
								leaf	
1	0	None	Gini	10	30	1	2	0.01	96,77% /
									89,81%
2	0.01	None	Entropy	100	100	1	2	0	99,41% /
									86,34%
3	0	Balanced	Entropy	10	10	1	2	0.001	95,90% /
									95,37%

Table 1: Best parameters and prediction results with Decision Tree algorithm

3.1.2 SVM: SupportVectorClassifier (scikit-learn)

\mathbf{M}	\mathbf{C}	coef0	gamma	kernel	shrinking	ACC (TR/TS)
1	1	0.5	Scale	Poly	True	100% / 100%
2	1.8	7	Scale	Poly	True	100% / 99,54%
3	0.02	1.5	Scale	Poly	True	93,44% / 97,22%

Table 2: Best parameters and prediction results with SVM algorithm

3.1.3 Neighbors: KNeighborsClassifier (scikit-learn)

\mathbf{M}	algorithm	leaf size	${f metric}$	#neighbors	p	weights	ACC (TR/TS)
1	BallTree	25	Minkowski	10	5	distance	100% / 84,49%
2	auto	10	Minkowski	21	1	uniform	66,86% / 66,20%
3	BallTree	2	Minkowski	18	1	distance	100% / 92,82%

Table 3: Best parameters and prediction results with K-Neighbors algorithm

3.1.4 Neural Networks: MLP (scikit-learn, SciKeras, PyTorch Lightning)

\mathbf{M}	topology	act	η	α	momentum	MSE (TR/TS)	ACC (TR/TS)
1	(4, 4, 4,)	TanH	0.5	0	0.7	0,0 / 0.009	100% / 99,07%
2	(4, 4, 4,)	TanH	0.9	0	0.7	0,006 / 0,037	99,41% / 96,30%
3	(2,)	ReLU	0.7	0.1	0.9	0,057 / 0,06	94, 26% / 93, 98%

Table 4: Best parameters and prediction results with scikit-learn's MLP

\mathbf{M}	topology	act	η	λ	momentum	MSE (TR/TS)	ACC (TR/TS)
1	(8,)	ReLU	0.9	0	0.7	0,003 / 0,015	100% / 97,92%
2	(8,)	ReLU	0.2	0	0.9	0,0002 / 0,0003	100% / 97,92%
3	(12,)	ReLU	0.01	0.001	0.7	0,056 / 0,039	100% / 97,92%

Table 5: Best parameters and prediction results with MLP in Keras

\mathbf{M}	topology	act	η	λ	momentum	MSE (TR/TS)	ACC (TR/TS)
1	(8,)	ReLU	0.1	0	0.1	0,0005 / 0,0027	100% / 100%
2	(8,)	ReLU	0.1	0	0.1	0,0002 / 0,0003	100% / 100%
3	(12,)	ReLU	0.1	0.001	0.3	0,019 / 0,028	98.97% / 98,15%

Table 6: Best parameters and prediction results with MLP in Pytorch

3.2 Cup Results

For this part of the experiments we proceeded as previously stated in Section 2. Details about each model can be found in the related Subsection.

3.2.1 TREE

For both the models *DecisionTreeRegressor* (DTR) and *ExtraTreeRegressor* (ETR) we performed a grid search as previously stated in Section 2. For these models We chose to test the same set of hyperparameter values (see Table 7) while keeping in mind their main difference, namely the splitter criterion which is *best* for DTR and *random* for ETR. The

best two models selected by the search are a DTR with $ccp_alpha = 0.0$, $max_depth = 7$, $min_samples_leaf = 6$, $min_samples_split = 2$ and $min_weight_fraction_leaf = 0.0$ and an ETR with $ccp_alpha = 0.0$, $max_depth = 15$, $min_samples_leaf = 1$, $min_samples_split = 2$ and $min_weight_fraction_leaf = 0.005$. The grid searches lasted for about 3 and 1 minutes, respectively.

Hyperparameters	Values
max_depth	5, 7, 10, 15, 20, 25, 30, 50
min_samples_split	2, 3, 4, 5, 6, 8, 10
min_samples_leaf	1, 2, 3, 4, 5, 6, 8, 10
min_weight_fraction_leaf	0.0, 0.1, 0.5, 0.01, 0.05, 0.001, 0.005, 0.0001, 0.0005
ccp_alpha	0.0, 0.1, 0.5, 0.01, 0.05, 0.001, 0.005, 0.0001, 0.0005

Table 7: Grid search parameters for both DecisionTreeRegressor and ExtraTreeRegressor.

3.2.2 SVM

SupportVectorRegressor (SVR) is not specifically designed for multi-target regression but we were able to use it for this task thanks to MultiOutputRegressor wrapper class available in scikit-learn which extended it to handle multiple target variables. The range of tested hyperparameter values by grid search is shown in Table 8. The best model has C=3, $coef\theta=0.0$, gamma=auto, kernel=rbf and shrinking=False. The training and validation MEE were respectively 1.2855 and 1.4690. The time required for the grid search was around 30 seconds.

Hyperparameters	Values
kernel	linear, poly, rbf, sigmoid
С	0.1, 0.5, 1, 2, 3
coef0	0.0, 0.01, 0.1, 0.5
gamma	auto, scale
shrinking	True, False

Table 8: Grid search parameters for SupportVectorRegressor.

3.2.3 NEIGHBORS

For the KNeighborsRegressor (KNR), a preliminary search for the ideal number of neighbors was performed by keeping the other parameters with their default values. Figure 1 illustrates how such parameter below 30 results in a lower MEE score. The range of tested hyperparameter values by grid search is shown in Table 9. The best model has algorithm = auto and $n_neighbors = 16$ and the training and validation MEE were 1.3488 and 1.4555, respectively. The grid search required around 2 minutes.

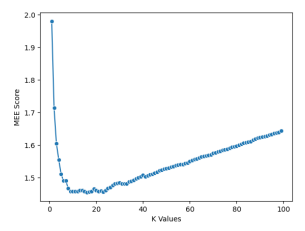


Figure 1: Preliminary search of the optimal number of neighbors for KNeighborsRegressor.

Hyperparameters	Values
n_neighbors	np.arange $(2,30)$
algorithm	auto,ball_tree,
	kd_tree, brute

Table 9: Grid search parameters for KNeighborsRegressor.

3.2.4 ENSEMBLE

The most significant parameter of this class of models is the number of estimators. For this reason, an initial investigation of that number was made for both of the models, leaving the other parameters unchanged, before proceeding with the actual grid search. The upper bound for these investigations was set to 100 due to the limited number of training samples available for this task. All the hyperparameters tested by grid search are shown in Table 10 and Table 11, for RandomForestRegressor (RFR) and BaggingRegressor (BR) resepctively. The best RFR has $ccp_alpha = 0.0$, $max_depth = 10$, $min_samples_leaf = 1$, $min_sample_split = 5$, $min_weight_fraction_leaf = 0.0$ and $n_estimators = 70$ with a training MEE equals to 0.7417 and a validation one equals to 1.4792. The time required for its grid search was around 5 minutes. The best BR has bootstrap = False, $bootstrap_features = False$, $max_samples = 0.3$ and $n_estimators = 70$ with 1.0305 and 1.4753 as training and a validation MEE, respectively. The time required for this grid search was less than 1 minute.

Hyperparameters	Values
$n_{\text{-}}$ estimators	50, 60, 70
max_depth	None, 5, 10, 15
min_samples_split	2, 3, 5
min_samples_leaf	1, 5, 10
$min_weight_fraction_leaf$	0.0, 0.01, 0.05, 0.1
ccp_alpha	0.0, 0.01, 0.05, 0.1

Table 10: Grid search parameters for RandomForestRegressor.

Hyperparameters	Values
$n_{\text{-}}$ estimators	30, 40, 50, 60, 70
max_samples	1.0, 0.1, 0.3, 0.5,
	0.7, 0.8, 0.9
bootstrap	False, True
bootstrap_features	False, True

Table 11: Grid search parameters for BaggingRegressor.

3.2.5 NN

For this class of models we decided to implement the same model (MLP) by using different libraries for its implementation. In particular, we used MLPRegressor from scikit-learn, KerasRegressor class from SciKeras, which integrates Keras functionality within scikit-learn, and NeuralNetRegressor class from Skorch, which instead integrates PyTorch functionality within scikit-learn. This allowed us to use in all three cases the GridSearchCV model selection method of scikit-learn, which is common to all CUP exepriments. The three models share the following implementation choices: the mini-batch stochastic method, also known as SGD; Mean Squared Error (MSE) as the loss function; batch size set at 16; maximum number of epochs set at 100; early stopping as the stop condition based on the validation loss with a tolerance of 20 epochs, possible through a 20% validation fraction; and the use of L2 regularization. The default minimum change in the monitored quantity to qualify as an improvement in the early stopping condition is zero but it has been changed into 0.01 in the non scikit-learn MLPs. Even with this kind of stopping condition, a previous testing activity demonstrated how more than 4 layers and 50 units can easily result in overfitting. Because of this, our final grid search does not test a larger number of them. For what concern the activation functions, ReLu and TanH proved to be the most effective while the Linear one was used for the 2-units output layer. For the weights and biases initialization we decided to take advantage of their default values: scikit-learn and SciKeras MLPs uses the Glorot uniform initializer [4], also known as the Xavier uniform initializer, which selects samples from a uniform distribution within the range [-limit, limit]where $limit = \sqrt{\frac{6}{fan_in+fan_out}}$ as the kernel initializer and zero for the bias initialization, while Skorch MLP initializes both weights and biases from a uniform distribution within the range [-limit, limit] where $limit = \sqrt{\frac{1}{fan_in}}$ [1]. In this context, fan_out is the number of output units, whereas $fan_{-}in$ is the number of input units in the weight tensor. The hyperparameter values tested by grid search for the scikit-learn library are shown in Table 12 while for SciKeras and Skorch in Table 13. The best MLPRegressor has $hidden_layer_sizes = (50, 50, 50)$, activation = tanh, $learning_rate_init = 0.01$, momentum = 0.0 and alpha = 0 and the training and validation MEE are 1.4187 and 1.4628, respectively. The grid search required around 12 minutes. The best KerasReqressor has $num_hidden_layers = 1$ (4 layers in total), $h_units = 40$, activation = tanh, $learning_rate = 0.01$, momentum = 0.5 and $weight_decay = 0.0001$ and the training and validation MEE are 1.3976 and 1.4151, respectively. The grid search required around 3 hours. The best NeuralNetRegressor has $num_hidden_layers = 1$ (4 layers in total), $h_units = 40$, activation = tanh, $learning_rate = 0.01$, momentum = 0.0 and $weight_decay = 0.001$ and the training and validation MEE are 1.3627 and 1.4564, respectively. The grid search required around 45 minutes. The MEEs of the last two models correspond to those of the models with the median validation MSE score among five retrainings with relative best parameters and different initializations.

Hyperparameters	Values
hidden_layer_sizes	(50,), (50,50,), (50,50,50,),
	(40,), (40,40,), (40,40,40,),
	(30,), (30,30,), (30,30,30,)
activation	tanh, relu
learning_rate_init	0.001, 0.01, 0.1
momentum	0.0, 0.01, 0.02, 0.03
alpha	0, 0.0001, 0.001, 0.01, 0.1

Table 12: Grid search parameters for scikit-learn's MLP.

Hyperparameters	Values
num_hidden_layers	0, 1
h_units	50, 40, 30
lr	0.008, 0.01, 0.02
momentum	0.0, 0.01, 0.03, 0.1, 0.5
weight_decay	0.0001, 0.001, 0.01, 0.1
activation	ReLU, Tanh

Table 13: Grid search parameters for Skorch and SciKeras's MLP.

3.2.6 Model Assessment

Table 14 summarizes the scores for each of the models. For this part of the experiments we chose to take into account a group of the top models ranked by considering their validation MEE, in particular the first five. Therefore, the resulting ensemble consists of the three *MLP*s implementations, *KNeighborsRegressor* and *SupportVectorRegressor*, the ones highlighted in gray in Table 14. Due to the other models' validation MEE score and the perceived quite high disparity between training and validation MEE scores, which may indicate overfitting of the model, they have not been used for the final ensemble. In order to assess the MEE on the development set and the internal test set, extracted at the beginning of the task (see Section 2), we used the computed average of the predictions of the five selected models. The final results are 1.3113 and 1.4349, respectively.

Model	Validation MEE	Training MEE	GAP
MLP (SciKeras)	1.4151	1.3976	0.02
KNR	1.4555	1.3488	0.11
MLP (Skorch)	1.4564	1.3627	0.09
MLP (scikit-learn)	1.4628	1.4187	0.04
SVR	1.4690	1.2855	0.18
BR	1.4753	1.0305	0.44
RFR	1.4792	1.7417	0.74
DTR	1.7264	1.1848	0.54
ETR	1.8040	1.5208	0.28

Table 14: Models' rank based on their validation MEE score.

4 Conclusion

Our research on this multi-target regression task reveals that Neural Networks (NN), K-Nearest Neighbors (KNN) and Support Vector Machines (SVM) are the most effective classes of models, while Ensemble and Trees tend to perform worse according to the MEE score. Even if a model performance could depend on many factors, this kind of behaviour confirms what we expected from a theoretical point of view due to the adaptability, capacity for generalization, and suitability for various data structures of NN, SVM and KNN. Ensemble techniques and Decision Trees, on the other hand, could have a propensity for overfitting and restrictions in modeling complex relationships.

For a complete plot perspective, please refer to Appendix A.

Our implementation code is fully replicable and available on GitHub at https://github.com/VeronicaPistolesi/MachineLearning. Our team's name is Ladydebugs and the final output for the CUP competition is in Ladydebugs_ML-CUP22-TS.csv file.

Acknowledgments

We agree to the disclosure and publication of our names, and of the results with preliminary and final ranking.

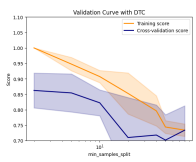
References

- [1] Linear layer initialization in pytorch. https://pytorch.org/docs/stable/generated/torch.nn.Linear.html#torch.nn.Linear. Accessed: April, 2023.
- [2] Adrian Garcia Badaracco. Scikit-learn wrapper for keras. https://github.com/adriangb/scikeras/, 2022. Accessed: April, 2023.
- [3] Dheeru Dua and Casey Graff. UCI machine learning repository, 2017.
- [4] Xavier Glorot and Yoshua Bengio. Understanding the difficulty of training deep feed-forward neural networks. In Yee Whye Teh and Mike Titterington, editors, *Proceedings of the Thirteenth International Conference on Artificial Intelligence and Statistics*, volume 9 of *Proceedings of Machine Learning Research*, pages 249–256, Chia Laguna Resort, Sardinia, Italy, 13–15 May 2010. PMLR.
- [5] F. Pedregosa, G. Varoquaux, A. Gramfort, V. Michel, B. Thirion, O. Grisel, M. Blondel, P. Prettenhofer, R. Weiss, V. Dubourg, J. Vanderplas, A. Passos, D. Cournapeau, M. Brucher, M. Perrot, and E. Duchesnay. Scikit-learn: Machine learning in Python. *Journal of Machine Learning Research*, 12:2825–2830, 2011.
- [6] Marian Tietz, Thomas J. Fan, Daniel Nouri, Benjamin Bossan, and skorch Developers. skorch: A scikit-learn compatible neural network library that wraps PyTorch, July 2017.

Appendix A

Please note that with *learning curves* we refer to plots with MEE scores on the y-axis and training set sizes on the x-axis, whereas with training curves we refer to plots with MEE scores on the y-axis and **epochs** on the x-axis.

Monk plots



Training score

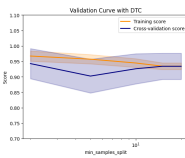
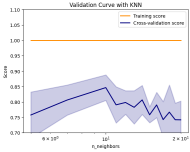


Figure 2: Learning curve of Decision Tree on MONK 1.

Figure 3: Learning curve of Decision Tree on MONK 2.

Figure 4: Learning curve of Decision Tree on MONK 3.



Validation Curve with KNN Training score

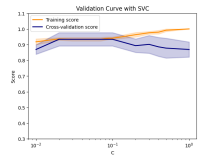
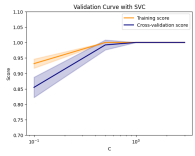
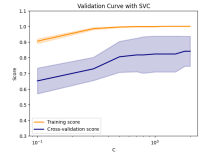


Figure 5: Learning curve of KNN on MONK 1.

KNN on MONK 2.

Figure 6: Learning curve of Figure 7: Learning curve of KNN on MONK 3.





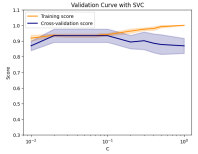


Figure 8: Learning curve of Figure 9: Learning curve of Figure 10: Learning curve of SVM on MONK 1.

SVM on MONK 2.

SVM on MONK 3.

MLP (scikit-learn)

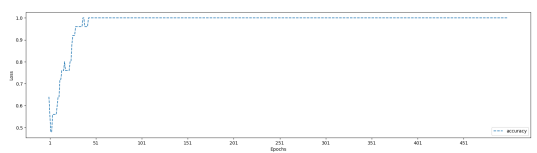


Figure 11: Accuracy in training curve of scikit-learn's MLP on MONK 1.

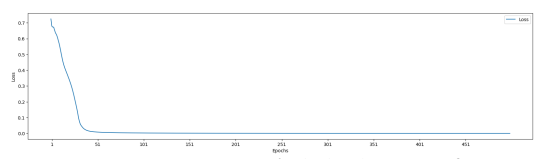


Figure 12: Loss in training curve of scikit-learn's MLP on MONK 1.

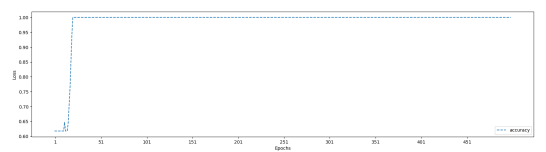


Figure 13: Accuracy in training curve of scikit-learn's MLP on MONK 2.

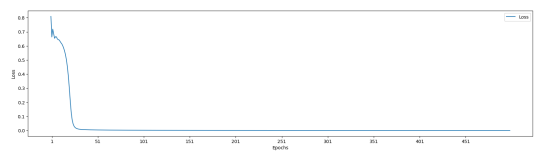


Figure 14: Loss in training curve of scikit-learn's MLP on MONK 2.

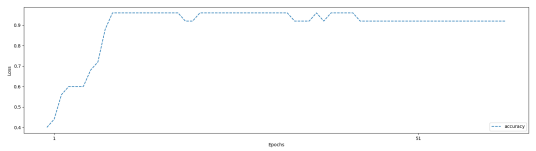


Figure 15: Accuracy in training curve of scikit-learn's MLP on MONK 3.

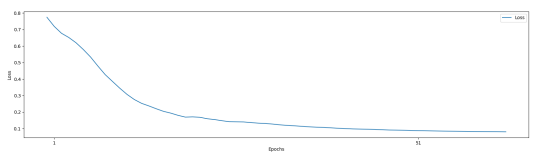


Figure 16: Loss in training curve of scikit-learn's MLP on MONK 3.

MLP (SciKeras)

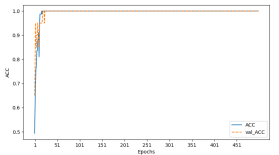


Figure 17: Accuracy in training curve of SciKeras' MLP on MONK 1.

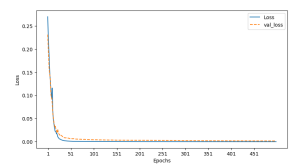


Figure 18: Loss in training curve of SciKeras' MLP on MONK 1.

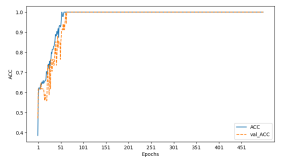


Figure 19: Accuracy in training curve of SciKeras' MLP on MONK 2.

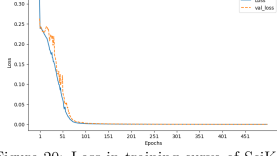


Figure 20: Loss in training curve of SciKeras' MLP on MONK 2.

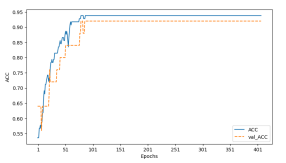


Figure 21: Accuracy in training curve of SciKeras' MLP on MONK 3.

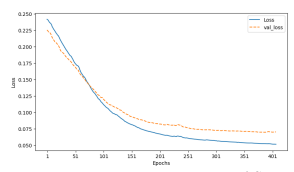


Figure 22: Loss in training curve of SciKeras' MLP on MONK 3.

MLP (Skorch)

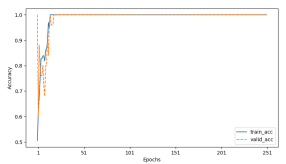


Figure 23: Accuracy in training curve of Skorch's MLP on MONK 1.

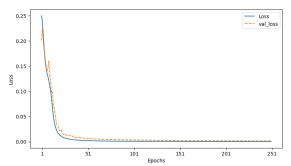


Figure 24: Loss in training curve of Skorch's MLP on MONK 1.

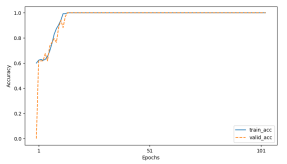


Figure 25: Accuracy in training curve of Skorch's MLP on MONK 2.

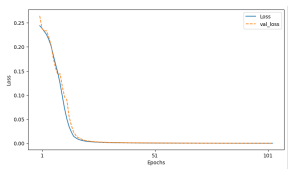


Figure 26: Loss in training curve of Skorch's MLP on MONK 2.

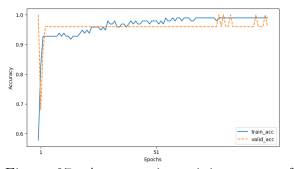


Figure 27: Accuracy in training curve of Skorch's MLP on MONK 3.

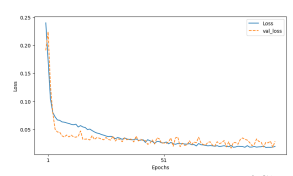


Figure 28: Loss in training curve of Skorch's MLP on MONK 3.

CUP plots

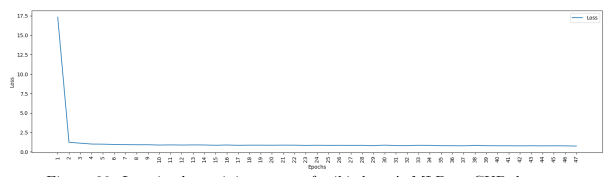


Figure 29: Loss in the training curve of scikit-learn's MLP on CUP dataset.

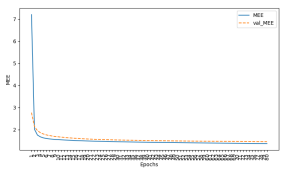


Figure 30: MEE in training curve of Skorch's MLP on CUP dataset.

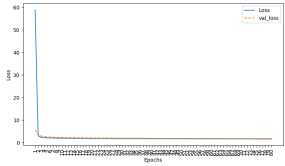


Figure 31: Loss in training curve of Skorch's MLP on CUP dataset.

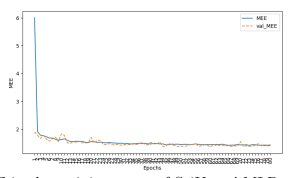


Figure 32: MEE in the training curve of SciKeras' MLP on CUP dataset.

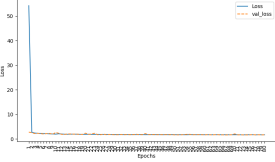


Figure 33: Loss in the training curve of SciKeras' MLP on CUP dataset.