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ABSTRACT

Near Infrared (NIR) spectroscopy is an analytical technology widely used for the nondestructive characterisation of organic samples, considering both qualitative and quantitative attributes. In the present study, the combination of Multi-target (MT) prediction approaches and Machine Learning algorithms has been evaluated as an effective strategy to improve prediction performances of NIR data from wheat flour samples. Three different Multi-target approaches have been tested: Multi-target Regressor Stacking (MTRS), Ensemble of Regressor Chains (ERC) and Deep Structure for Tracking Asynchronous Regressor Stack (DSTARS). Each one of these techniques has been tested with different regression methods: Support Vector Machine (SVM), Random Forest (RF) and Linear Regression (LR), on a dataset composed of NIR spectra of bread wheat flours for the prediction of qualityrelated parameters. By combining all MT techniques and predictors, we obtained an improvement up to 7% in predictive performance, compared with the corresponding Single-target (ST) approaches. The results support the potential advantage of MT techniques over ST techniques for analysing NIR spectra.

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1. Introduction

Near-infrared (NIR) spectroscopy is a consolidated technology widely applied in several research fields such as food science [3,2,51,45,4,33,43,39,23,39], agriculture [47,30,42,13,17] and

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biomedicine [27,18]. The main advantages of NIR spectroscopy include the possibility of performing rapid, minimally invasive, simple, reagent-free and non-destructive measurements [33,47,42,20]. In this way, NIR analysis is capable of coping with the modernisation of processing industries, improve health diagnoses, reduce costs towards a sustainable alternative in food and raw material characterisation [32,18,41,42].

Essentially, NIR spectroscopy is based on absorption bands derived from overtones and combinations of fundamental vibrations of chemical bonds (mainly C–H, N–H, O–H, S–H,

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C—C and C=O) observed in the 780–2500 nm range of the electromagnetic spectrum [40]. In general, a multivariate statistical approach is mandatory in order to extract the useful information from NIR spectra. In this context, multivariate analysis allows to relate the spectral properties of a representative set of samples with the desired response and to use the resulting model to predict new samples [22,8].

Usually, PLS is the multivariate approach more commonly applied for regression tasks [42]. [33] have highlighted that PLS is the linear-based method more often used for regression models in NIR spectra analysis [33].

In particular, multivariate linear methods (such as PLS) can be considered as a branch of Machine Learning (ML) supervised approaches, which are able to automatically detect patterns in the data and use these patterns to predict future data. However, in some situations changes in the physical and chemical properties of the analysed samples can cause deviations from linearity. Possible non-linear contributions may result from changes in temperature, particle size, viscosity and chemical composition of the samples [47]. In order to solve this issue, non-linear ML methods have been successfully applied to model NIR spectra in several scenarios [5,16,47].

Among non-linear ML algorithms, Support Vector Machines (SVM) has been proven to be an accurate and reliable method, often superior to other regression or classification methods in the field of spectroscopic analysis [45]. [42] described the improvements obtained with SVM compared to PLS. However, the same paper also highlighted that generally, SVM requires a strong optimisation of hyper-parameters and a feature selection step in order to obtain successful results.

Other ML techniques, such as Artificial Neural Networks (ANN), are able to handle non-linear features with a little advantage in prediction accuracy. [30] compared ANN and PLS in the prediction of organic carbon, pH and clay content of soils. The results showed similar performances of both methods, but some laboratory independent validation and on-line evaluation proved the over-performance of ANN models. An important drawback in the use of ANN was highlighted by [51]. Indeed, due to the high number of tunable hyper-parameters and to the limited number of samples usually available in real applications, ANN is hindered by overfitting problem.

In addition, a drawback of non-linear methods is the lack of interpretability. As a matter of fact, the relevance of the different spectral features is a very valuable information for the chemical interpretation of a regression or classification model. Considering model interpretability, [33], reported that linear based machine learning methods are superior to nonlinear ones.

In this context, Random Forest (RF) algorithm can combine both model performances and interpretability, since it generally leads to highly accurate prediction and it can be helpful in spectra interpretation by RF importance. RF was proposed by [10] as a combination of decision tree classifiers in an ensemble. The author describes RF as a trustful method that always converges, avoiding overfitting problem. In the most common RF algorithm, split selection for decision tree building is performed based on the decrease of Gini impurity. This feature value, named as RF importance, provides a relative ranking of the spectral features as described in [38]. Therefore, RF is a useful tool for regression studies and it has a potential for modelling linear and non-linear spectral responses [24]. [6] explored eight different ML algorithms for the prediction of pork meat storage time, with RF achieving the best results.

Although ML non-linear algorithms have been proved to be profitable for spectroscopy, PLS is still a common strategy for modelling spectral data thanks to the ability of dealing with high dimensional and multicollinear data, and the possibility of identifying the relevant predictors with a gain in model interpretability [47,26]. Furthermore, spectroscopic data usually consist of numerous features (wavelengths) and relatively few samples. In this situation, the optimisation of many meta-parameters can easily lead to overfitting [31]. In order to combine the advantages of PLS and ML non-linear algorithms, PLS can be used as a sort of data pre-processing and data compression method to extract the relevant features from spectroscopic data, which can later be used as predictors for non-linear regression methods.

Another issue of great relevance is that, in practical applications, NIR spectroscopy can be used to predict multiple quality parameters of a given sample. In this situation, Multi-target (MT) approaches of Machine Learning (ML) allow to obtain better performances compared to traditional singletarget modelling (ST) [43]. Indeed, MT methods gain advantages from the exploration of inter-target influences and allow to reduce overfitting [9,36,35]. Furthermore, a MT model provides a global comprehension of a given problem by also considering the relationships between the different targets, in addition to the relationships between features and expected predictions (targets) [9]. [37] highlighted that MT prediction has the ability to generate models representing a broad variety of real-world applications, from natural language processing to bioinformatics.

In this context, we propose the usage of MT techniques coupled with ML non-linear algorithms induced on Partial Least Squares regression (PLS) scores to predict statistically correlated targets from NIR spectral data. It is possible to transform NIR data into PLS scores in order to compress hundreds of variables into few relevant features. On the other hand, Multi-targets techniques coupled with base-learners from Machine Learning are capable of dealing with nonlinear behaviour and noisy data with a little prediction error.

In the present work, different MT approaches have been tested: Multi-target Regressor Stacking (MTRS) [46], Ensemble of Regressor Chains (ERC) [46] and Deep Structure for Tracking Asynchronous Regressor Stack techniques from Multi-target (DSTARS) [36]. Each MT technique has been coupled with Support Vector Machine (SVM), Random Forest (RF) and Linear Regression (LR) as base-learners to perform non-linear and linear modelling, respectively. As a case study, the proposed MT approaches have been tested in the prediction of quality-related parameters of bread wheat [21].

Results exposed the advantage of Multi-target strategy reaching an improvement of 7% with ERC and RF. Independent of ML algorithm, in all cases, the usage of MT coupled with SVM or RF increased the predictive performance.

Predictions obtained from RF overcome the SVM performance. Specifically, the MTRS method achieved inferior results when coupled with an LR.

2. Materials and methods

2.1. NIR datasets

The bread wheat samples considered in this study were collected from experimental fields located in different Italian regions and derived from two subsequent harvesting years. On the whole, 391 bread wheat white flour samples were analysed with a Bruker MPA Multi Purpose FT-NIR Analyzer, equipped with an integrating sphere (reflectance mode) and a RT-PbS detector in the 12,500–3600 cm⁻¹ range.

The spectra were acquired following the acquisition parameters defined in [19] (8 cm⁻¹ resolution, 155 sample scans). For each white flour sample, four replicate measurements were performed in different days, each time considering a different 50.0 g aliquot. The sample aliquots were placed into a glass Petri dish and gently shaken to render the packing density as much uniform as possible. The FT-NIR instrument was equipped with a rotating device, in order to acquire an average signal over a relatively wide sample surface. Between the different measurement sessions, samples were stored in sealed plastic bags and put into a plastic box in a dark place at 4 °C. Finally, the spectrum of each sample was obtained as the average of the four replicate measurements.

For each sample, the following quality-related parameters (targets) have been determined using the corresponding reference methods: Hectolitre weight (HlW, kg hl⁻¹), Falling Number (FN, s), Protein content (Prot, % dry matter), Alveographic indexes (W and P/L), and Farinograph stability (Stab, min). Further details about the quality parameters can be found in [11].

The linear statistical correlation between the considered quality parameters (HlW, FN, Prot, W, P/L, and Stab) can be evaluated in Fig. 1, which reports the Pearson correlation coefficient calculated for the different parameters.





Kennard-Stone algorithm applied to the principal components space was used in order to split the whole dataset of NIR spectra into two datasets: the training cross-validation set ($Dataset_{CV}$) composed of 200 samples and the prediction set ($Dataset_P$) of 91 samples. The cross-validation set was used to calculate and optimise the regression models, while the prediction test set was employed to finally evaluate the predictive ability of the traditional ST approach compared to the multi-target ones. This subdivision of the dataset was performed in order to minimise the risk of overfitting.

2.2. Learn-based regressors

The training set of NIR spectra (obtained as reported in Section 2.1) has been firstly analysed by means of PLS. PLS is a multivariate analysis technique in which the independent variables are projected onto a small number of latent variables (LVs) to simplify the relationship between them and a predictive target [33]. The target variable is actively used in assessing the LVs to ensure that the first one is most relevant for predicting the targets. Usually, the relation between the scores extracted from LVs and the targets are built by a linear modelling toward prediction of novel samples. The PLS technique was implemented by SIMPLS algorithm [14]. In particular, the PLS scores have been extracted from the optimal PLS model (i.e., for the model with a number of LVs corresponding to the minimum value of the root mean square error in cross-validation), calculated on the mean-centred training set spectra for each one of the auto-scaled target variables. Subsequently, the test set spectra have been projected on the e PLS latent variable space and the corresponding scores have been calculated. These steps are highlighted in the overview of procedures as 1 and 2, reported in Fig. 2. In this manner, it was possible to extract the useful information contained in hundreds of spectral variables into few latent variables relevant for the prediction of the targets. The ST models were then calculated considering the PLS scores selected for each target, while the ensemble of all the PLS scores was used as input for the subsequent development of multi-target regression models. In particular, the three different MT approaches described in Section 2.3 have been tested (ERC, MTRS and DSTARS).

MT algorithms require a learn-based regressor in the stack or chain structure to induce each model. In order to evaluate the benefits induced by linear and non-linear learn-based regressors, MT approaches have been coupled with Linear Regression (LR), Support Vector Machine (SVM) and Random Forest (RF), as shown in Step *a* of Fig. 2.

SVM is a regression algorithm from kernel-based methods. It can be used for solving many types of problems, presenting high accuracy and ability to treat high-dimensional data. Through kernel space transformation, this technique has the flexibility to model diverse data sources [7], increasing the input dimensional space and data separability. In this work, the SVM implementation adopted was the elo71 R package with default parameters.

RF was initially created focusing on classification tasks. This algorithm consists of a collection of structured tree predictors, in which all random vectors are independent, identically distributed and each tree attributes a vote for the most





popular class as the prediction. On the other hand, for regression purposes, the RF is composed of trees depending on a random vector, considering the average result over all trees in the Forest [10]. Each tree is grown using a bagging approach, where different training datasets are formed using bootstrap sampling. The randomForest R package was used with default parameters in experiments.

Therefore, each regression model has been adapted according to the requirements of the three MT techniques (Step b inside Fig. 2).

2.3. Multi-target prediction

The concept of Multi-target (MT) regression is related to the family of predictive problems with multiple continuous response variables, called as targets or outputs, following the assumption of being statistically correlated, i.e. the variation of a target has interference on the behaviour of the other responses [1,48,9,46]. Traditionally, MT tasks have been solved through two different approaches: Algorithm Adaptation and Problem Transformation [9].

Algorithm Adaptation, also named as multi-output adapted, is based on the adaptation of single target regression methods to handle multiple outputs exploring the dependencies among them with an enhancement in the predictive capability. This strategy is performed by modifying the original modelling method, for example changing the optimisation functions (e.g. SVMs) [34,52,50] or the node splitting criteria (e.g. Regression Trees) [29]. This type of approach has been successfully proposed in the last years focusing on different tasks [28,49,9,25,53].

In general, multi-output adapted models generate effective predictors by taking advantage of statistical target dependencies to create a single model. However, this approach could be more challenging since it aims not only to predict multiple targets but also to model and interpret their relationships at once [36]. [52] proposed the modification of task's input space through a virtualisation technique so that a MT task could be modelled as a wider single target problem. The authors used a Support Vector Regression (SVR) and achieved results comparable to ST strategy.

Another approach to address MT scenarios is Problem Transformation. This approach is based on calculating independent ST regression models for each target and finally concatenating the predictions. The calculation of several independent ST models to represent a single MT problem involves a strong increase of computational efforts and in several cases a loss of model interpretation by ignoring the dependencies between targets. On the other hand, this kind of approach offers considerable advantages. The first one is the possibility of applying any base-learner, or even more than one, toward better predictive performance and appropriate problem addressing. Also, adaptation methods improve the modularity and conceptual simplicity, with significantly better predictive performance than state-of-the-art methods [46].

Tsoumakas et al., in [48], proposed the use of random linear targets combinations to explore the relations between output values. This approach increases the original feature space dimension and solves multiple ST problems in the transformed space. The predicted values are used to solve a linear system and obtain the original targets predictions. In the last years, some MT methods have been modified and adapted from the area of multi-label classification [9,46]. Spyromitros-Xioufis et al. [46] proposed two techniques based on Problem Transformation: MTRS (Multi-target Regressor Stacking) and ERC (Ensemble of Regressor Chains). These two techniques are widely known and resulted to be adequate for a diversity of MT scenarios.

Santana et al. [44], proposed the Deep Regressor Stack, an idea similar to MTRS that consists in using targets approximations as additional predicting features in naive deep learning method. However, the drawback of huge memory requirements, as well as inappropriate dimensionality growth, preclude its usage associated with NIR spectra analysis. More recently, Santana et al., in a different work [43], proposed a new method, Multi-target Augmented Stacking (MTAS), addressing multi-target regression to predict twelve poultry meat characteristics. The authors explored the usage of Principal Components Analysis (PCA). Different from [43], in this work, we take advantage of PLS toward exploring DSTARS [36] MT approach. PLS compression leads to the extraction of features that are more relevant to target prediction.

Basically, DSTARS consists in a modification of MTRS built under the assumption that a model induction by deeper layers could offer better predictive performances than just one layer (ST) or two layers (MTRS). Therefore, DSTARS can take

advantage of different linear and non-linear relations among the targets, leading to possible improvements in the prediction performances of complex data, such as NIR spectra.

In the following sections a detailed description of MTRS, ERC and DSTARS algorithms is given.

2.3.1. Multi-target Regressor Stack

MTRS approach is based on additional input features from the output of ST models induction. In this way, considering a dataset composed by $X = \{x_1, x_2, ..., x_n\}$ input features and $Y = \{y_1, y_2, ..., y_d\}$ target variables, MTRS adds the $Y' = \{y'_1, y'_2, ..., y'_d\}$ from ST predictions as inputs, creating a new training dataset $X' = \{x_1, x_2, ..., x_n, y'_1, y'_2, ..., y'_d\}$. The new training dataset is used by each y to train another ST predictors layer, whose outputs are the final predictions.

New income input features are first merged into the first predictors' layer to obtain the output approximations. The approximations compose an augmented dataset toward forming the second level of predictors for the purpose of performing the final induction. Therefore, MTRS introduces inter-target relationships by the use of ST outputs into the modelling, increasing the prediction performance. The training procedure of MTRS is presented in Algorithm 1.

Algorithm 1 (MTRS training algorithm).

1: function MTRS(X, Y, d) $Y' \gets \{\}$ 2: 3: Level₀ \leftarrow {} // ST model induction 4: 5٠ for t = 1 to d do 6: h: $X \rightarrow Y^t$ 7: $Y'^{,t} \leftarrow predict(h, X)$ 8: $Level_0 \leftarrow \{Level_0, h\}$ 9: // Augmented training set definition 10: $X' \leftarrow X \| Y'$ 11: $Level_1 \leftarrow \{\}$ for t = 1 to d do 12: h: X' \rightarrow Y ^t 13: $Level_1 \leftarrow \{Level_1, h\}$ 14: 15: mtrs \leftarrow {Level₀, Level₁} 16: return mtrs

2.3.2. Ensemble of Regressor Chains

The ERC method consists in building a set of randomly ordered chains for each target to produce ST models through a generated sequence [46]. For each chain, initially, a ST model is induced using the first output prediction of the sequence. New models are then induced following the chain order, where each new regressor is trained over the augmented input dataset formed by the original input features and the previous models' predictions. This process is repeated until the end of all chain sequences. After training all models inside the chain, the predicted value of a novel sample is the average value obtained from chain's regressors.

The method prediction for a target y^t , t = [1, d], is the average of the y^t predicted values over all chains. Since the output

predictions are composed of values from different sorted chains, multiple levels of combinations and interdependence between targets are explored. ERC creates all possible target permutations if their number is less than 10 ($d \leq 3$), otherwise, the author suggests to choose ten random combinations.

The ERC's training step is presented in Algorithm 2. The *permute* procedure refers to a function which receives a set of elements and returns all their possible permutations. It is possible to perform all possible permutations without setting this parameter or to specify the desired number of combinations as an argument (in the original ERC's formulation, the mentioned parameter is equal to 10).

Algorithm 2 (ERC training algorithm).

1:	function ERC(X, Y, d)
2:	$targets \leftarrow names(Y)$
3:	if $d \leq 3$ then
4:	$Chains \gets permute(targets)$
5:	else
6:	Chains \leftarrow permute(targets, 10)
7:	for chain in Chains do
8:	$models_{chain} \leftarrow \{\}$
9:	// To build augmented training sets
10:	$X_{auq} \leftarrow X$
11:	for t in chain do
12:	// ST model induction
13:	$h: X_{aug} \rightarrow Y^{t}$
14:	$y_{pred} \leftarrow predict(h, X_{aug})$
15:	// Extend training set with ST predictions
16:	$X_{aug} \leftarrow X_{aug} \parallel y_{pred}$
17:	$models_{chain} \leftarrow \{models_{chain}, h\}$
18:	$erc \leftarrow \{erc, models_{chain}\}$
19:	return erc

2.3.3. Deep structure for Tracking Asynchronous Regressor Stack

DSTARS [36] is based on the hypothesis that deeper layers could emphasise the relationships among targets which are statistically correlated. Instead of using additional features from only one single layer as ST estimators, or strictly two layers as MTRS, DSTARS sequentially builds new output predictions from as many layers as those necessary to minimise the error of a validation set. Iteratively, each best target prediction (evaluated using the validation set) is used as an additional predictive feature, augmenting the dataset. The procedure of the DSTARS algorithm can be split into two main steps: Tracking and Modelling. The former determines the best layer depth of targets variable, while the latter builds the final DSTARS model considering the whole modelling dataset.

The Tracking step, presented in Algorithm 3, starts with the subdivision of the dataset into training and validation sets. The authors recommend the use of a sampling approach, as the k-fold cross-validation, to increase the robustness. During the Tracking step, for each target new

models are calculated using the actual best approximations of all the targets as extra input variables. DSTARS keeps the tracking of the models that generated the best performance on layer *l* for each target y_t . Not all layers need to be used during final modelling for a particular target: in general, targets with low inter-correlations usually require a low layer depth, while some targets may require a greater layer depth. The final configuration is determined through a voting scheme, considering the reduction of the Root Mean Squared Error (RMSE) of the validation set brought by each layer and target combination. The tracking process ends when the decrease of the RMSE value obtained by the addition of a new predictors' layer l + 1 is smaller than an ε parametrised value, leading to convergence.

Algorithm 3 (DSTARS's tracking algorithm).

```
1: function Tracking(X, Y, \varphi, \varepsilon, N<sub>folds</sub>)
 2:
        // Dynamic regressor layer usage count
 3:
         Track \leftarrow [?, d]
         for k = 1 to N_{folds} do
 4:
            // Cross-validation data split
 5:
             \{ (X_{tr}, Y_{tr}), (X_{val}, Y_{val}) \} \leftarrow CV(X, Y, k) \\ errors \stackrel{1..d}{\leftarrow} \infty 
 6:
 7.
                           \leftarrow \infty
            converged 1..d \leftarrow False
 8:
            layer \leftarrow 0
 9:
            T_{tr} \leftarrow T_{val} \leftarrow \{\}
10:
11:
            while !all(converged) do
               T_{tr} \mathrel{\scriptstyle{\textbf{'}}} \leftarrow T_{tr}
12:
                T_{val} \cdot \leftarrow T_{val}
13:
14:
                for t = 1 to d do
                  // ST model induction
15:
                   mod : X_{tr} \parallel T_{tr} \rightarrow Y_{tr}^{t}
16:
                   prd \leftarrow predict(mod, X_{val} \parallel T_{val})
17:
                   // Convergence stopping criteria
18:
                   if RMSE(Y _{val}^{t}, prd) +\varepsilon < errors ^{t} then
19:
20:
                      \textbf{converged} \ ^t \gets \textbf{False}
                      errors t \leftarrow RMSE(Y _{val}^{t}, prd)
21:
22:
                      // Error improvement counting
23:
                      Track[layer, t] \leftarrow Track[layer t] + 1
24:
                      T_{tr}^{t} \leftarrow predict(mod, X_{tr} \parallel T_{tr})
25:
                     T \stackrel{t}{_{val}} , \leftarrow prd
26:
                   else
                      converged t \leftarrow True
27:
28:
               T_{tr} \gets T_{tr} \; \text{'}
29:
               T_{val} \gets T_{val} \; \text{,}
30:
                layer \leftarrow layer + 1
31:
         32:
         // Threshold application: True or False values
         Track \leftarrow Track > \varphi
33:
34: return Track
```

In Modelling step, a model h_{t}^{l} that outputs the t-th target estimation on the l-th layer is included in the final model only if it was used more than the φ percent of times during Tracking. New income samples are sequentially subjected to each layer of regressors and the last layer gives the predicted value for a specific target.

The usual DSTARS's structure starts by Tracking step and goes through the Modelling step presented in Algorithm 4.

Algorithm 4 (DSTARS training algorithm).

1:	function DSTARS(X, Y, φ , ε , N _{folds})
2:	$T \leftarrow Tracking(X, Y, \varphi, \varepsilon, N_{folds})$
3:	dstars \leftarrow {}
4:	$Y_{approx} \leftarrow \{\}$
5:	for $l = 1$ to number of T rows do
6:	$Y_{aux} \gets Y_{approx}$
7:	for $t = 1$ to d do
8:	if T [l, t] = True then
9:	// ST model induction
10:	$mod : X \parallel Y_{approx} \rightarrow Y^{t}$
11:	$Y_{aux}^{t} \leftarrow predict(mod X \parallel Y_{approx})$
12:	dstars \leftarrow {dstars, mod}
13:	$Y_{approx} \leftarrow Y_{aux}$
14: :	return dstars

2.4. Performance metrics

The performances of ST and MT methods have been evaluated on the $Dataset_{CV}$, sampled through a 10-fold crossvalidation approach (CV), and the Prediction set ($Dataset_P$). In the final models, the whole Training set was used to build a single prediction model using the best MT hyperparameters determined in the CV approach, when considering the MTRS, ERC and DSTARS techniques.

Six different performance metrics were used to evaluate the calculated models: Mean Square Error (MSE), Average Relative Error (ARE), Average Root Mean Squared Error (aRMSE), Coefficient of Determination (R^2), average Relative Root Mean Square Error (aRRMSE), and Relative Performance (RP).

MSE corresponds to the mean squared difference between the predicted and the measured values. On the other hand, ARE was used to exposes the magnitude of the difference between the actual quality parameter and the prediction.

The aRMSE (Average Root Mean Square Error) is the mean of the Root Mean Square Error (RMSE) values obtained from each target. This last acts as a baseline and allows the measurement of the improvement over a shallow predictor. This metric has been used in various MT works [9] to compare non-homogeneous targets distributions. The aRRMSE, presented in Eq. (1), is computed averaging the *d* targets Relative Root Mean Square Error (RRMSE), and it was applied in this research to evaluate the improvement over ST models. The RRMSE for a given target *t* is the RMSE normalised by the average of the corresponding t.

$$aRRMSE = \frac{1}{d} \sum_{t=1}^{d} \sqrt{\frac{\sum_{k=1}^{k} (y_t^k - \hat{y}_t^k)^2}{\sum_{k=1}^{N_{test}} (y_t^k - \overline{y}_t)^2}}$$
(1)

The Coefficient of Determination (R^2) explains the proportion of the total variation associated with the dependent variable that is predictable from the model. The closer the R^2 values are to 1, the greater is the amount of variation of the dependent variable which is predictable by the regression model [12].

The Relative Performance (RP) compares for each target the aRRMSE of each MT method with the aRRMSE of the corresponding ST model. In this sense, it measures the increase (if RP > 1) or decrease (if RP < 1) in model performances [46]. The RP formulation is presented in Eq. (2).

$$RP_d(M) = \frac{aRRMSE(ST)}{aRRMSE(M)}$$
(2)

The aRRMSE supports the comparison of possible method superiority through the application of the Friedman's statistical test with significance level at α = 0.05. The null hypothesis states that the performances of the MT methods are equivalent regarding the averaged aRRMSE per dataset. Any time the null hypothesis is rejected, the Nemenyi post hoc test can be applied, stating that the performance of two models is significantly different if the corresponding average ranks differ by at least a Critical Difference (CD) value. When multiple models are compared in this way, a graphic representation can be used to represent the results with the Critical Difference (CD) diagram, as previously proposed in [15].

Finally, the results obtained from ST and MT approaches and expressed with the error metrics cited above were analysed by means of Principal Component Analysis (PCA) in order to obtain a graphical overview of the whole model performances and evaluate the effectiveness of the different MT methods.

3. **Results and discussion**

3.1. Multi-target and Single-target comparison

In order to gain a first overview of the performance of the ST approach compared with MT, Table 1 reports the results obtained with linear regression coupled with Single-target and Multi-target methods. In this context, it must be highlighted that the linear regression of the PLS score values in the ST approach corresponds to the usual PLS regression. Table 1 shows that DSTARS and ERC resulted to be the best performing methods, with the lower aRRMSE value and the higher R² value obtained on the test set.

In general, the potential improvement induced by the different MT approaches is highlighted in Fig. 3, which reports the RP improvement for each algorithm (RF, SVM and LR). ERC led to improvements in all model performances obtaining the best average RP (1.015) from both data sets. The second best average RP was obtained by DSTARS (1.010); it did not improve the predictive performance only for RF in Dataset_{CV}. However, DSTARS overcame ERC results for LR algorithm in Dataset_B obtaining 1.021 of RP. The worst results were achieved by MTRS with an average RP equals to 0.985. This approach was only able to contribute using SVM in Dataset_{CV}.

Concerning the different learn-based predictors, SVM took advantage from MT approaches with an increase of model performances, except for the case of the prediction of the test set (Dataset_P) coupled with MTRS. RF presented improvements with ERC, slightly boost coupled with DSTARS in $Dataset_{P}$ and no contributions over Dataset_{CV}. LR obtained the best RP with DSTARS, minor gains when combined with ERC and no improvements with MTRS.

In order to obtain a general overview of the performances of all the tested models, a PCA model was calculated on the results considering the metrics related to error performance from all the considered combinations of techniques and algorithms, and for both the CV and P datasets.

The optimal number of principal components (PC) has been found to be equal to 3, retaining the 97.72% of the total variance. Considering the behaviour of the loadings, essentially PC1 accounts for the errors obtained in prediction (Dataset_P), PC2 is related to the errors in cross-validation (Dataset_{CV}) and PC3 accounts for the differences between the error metrics. Fig. 4 reports the biplot of PC1 and PC2 feature space, explaining 81.90% of the total variance. Concerning the predictive performance of the learn-based regressors, LR generally led to a higher error in prediction for both ST and MT approaches, while RF is the algorithm giving the best prediction results. Considering the comparison between singletarget and multi-target approaches, generally ERC led to the higher enhancement of model performances, except for LR algorithm where the results obtained with DSTARS, ERC and ST are almost the same or inferior (similar to what observed in Fig. 3). In addition, it is possible to highlight that SVM coupled with ERC or DSTARS is the most stable algorithm since it led to lower error values in both cross-validation and prediction.

In order to emphasise a possible superiority of the different combinations of algorithms and MT strategies, Friedman's statistical test and the Nemenyi post hoc test have been applied to the averaged (P dataset) aRRMSE values. Fig. 5 shows the Critical Difference (CD) diagram obtained from the statistical test results. The different models are connected when statistically significant differences are not observed between them ($\alpha = 0.05$ and CD = 7.07). Lower rank values indicate the most accurate (lower aRRMSE) methods, while higher values indicate the less accurate ones.

According to the results reported in Fig. 5, no statistically significant differences were observed when comparing ERC + RF, DSTARS + RF, ERC + SVM, DSTARS + SVM, MTRS + RF, ST + RF. ST + SVM, DSTARS + LR and MTRS + SVM.

Table 1 – Average aRRMSE and R ² perform	nances of ST, ERC, MTRS and DSTA	RS approaches coupled with linear regressio	n
obtained on Dataset _{CV} and Dataset _P .			

	ST		ERC	ERC		MTRS		DSTARS	
	aRRMSE	R ²	aRRMSE	R ²	aRRMSE	R ²	aRRMSE	R ²	
Dataset _{CV} Dataset _p	0.7224 0.7454	0.4378 0.4008	0.7121 0.7366	0.4541 0.4143	0.7238 0.7664	0.4366 0.3680	0.7148 0.7304	0.4506 0.4230	

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Fig. 3 - Relative Performance (RP) comparison of RF, SVM and LR predictor over all datasets (Dataset_{CV} and Dataset_P).



Fig. 4 – PC space obtained from techniques (ST, MTRS, ERC and DSTARS) combined to algorithms (RF, SVM and LR) over all datasets (CV and P) by Mean Squared Error (MSE), Average Root Mean Squared Error (ARMSE), Average Relative Root Mean Squared Error (aRRMSE) and Average Relative Error (ARE).

Nevertheless, comparing the aRRMSE value of each regression algorithm, all the LR solutions obtained inferior results, including the ST + LR (PLS core). MTRS technique did not lead to significant prediction improvements; in some cases, the results obtained with this technique were inferior to those gained using the ST approach, which in turn was a

competitive solution when coupled with RF and SVM non-linear algorithms.

3.2. Algorithms prediction improvements

MT prediction problems are composed of targets that present different prediction complexity and distinct inter-correlation with each other. Section 2.1 reported the results obtained from Pearson linear correlation between the targets, which showed a considerable variation.

In order to evaluate the error reduction achieved by MT approaches separately for each target, the *RP*_t values obtained by comparing the *RMSE* performance between MT techniques and ST have been calculated for each different target. The results are shown in Fig. 6, where the *RP*_t values are reported as a heat map. For each combination of MT approach and lean-based regressor, a reddish colour is related to a decrease of model performance compared the corresponding ST method, while a greenish colour is related to an increase of model performances.

For both datasets, the predictive performances of *Hectolitre Weight* target were enhanced by the use of MT approaches, except for the case of MTRS + RF and MTRS + LR. In these last configurations, the target was predicted with inferior result when compared to ST method with RP_t 0.96 and RP_t 0.99, respectively. Similar behaviour was also observed for *P/L*, which has been improved with respect to ST strategy, and partly for *Stability*, for which improvements were observed only in cross-validation. Conversely, *Protein* and W showed lower error reduction, mainly in the case of MTRS, independently from ML algorithm. However, it has to be considered that the Coefficient of Determination of *Protein* target obtained by PLS (ST + LR) was equal to 0.90 (R^2), while for the same target the R^2 value in prediction obtained with DSTARS + RF was equal to 0.91. Therefore, for some targets

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Fig. 5 – Comparison of the averaged aRRMSE values from all technique and algorithm according to the Nemenyi test. Groups of methods that are not significantly different (α = 0.05 and CD = 7.07) are connected.





standard PLS was sufficient to obtain satisfactory results, and the use of MT approaches can lead only to slight improvements.

On the other hand, DSTARS + LR improved the performance of *Hectolitre Weight* from 0.50 (R^2) to 0.56 (R^2). This result meets the importance of exploring different levels of intra-target correlation to perform the target prediction. An important advantage of DSTARS is the possibility to create a MT strategy through the correlation between the targets and to reveal it by the depth of Tracking procedure. For this reason, DSTARS was capable of facing this issue obtaining results superior to ST by dealing with the different inter-correlation between the targets as shown in Fig. 7.

Fig. 7 shows the RMSE values associated with the model layers of each target. In some cases, for example Protein and P/L, an additional layer is not required to produce good predictors. However, some targets such as Stability and Falling

Number required an additional layer to improve the overall prediction. This figure exposes the RMSE of Modeling and Testing sets split in the kernel of DSTARS strategy as described in Section 2.3.3. The obtained representations meet the heat map representation of Fig. 6, since the targets with smaller improvement from MT approach require a shallow layer structure of DSTARS. This type of investigation was not supported by ERC technique; however, this latter allowed to achieve a better prediction error reduction, as exposed in Section 3.3.

3.3. Target prediction improvements

Different prediction improvements were obtained for each target. In Table 2, the RMSE values obtained for all the tested models are reported where, for each target, the best predictive performance is highlighted in bold. For cross-validation, no



Fig. 7 – DSTARS depth of layers obtained by the use of SVM as base-learner to predict the samples from Dataset_{CV}.

Dataset	Technique	Algorithm	RMSE					
			HlW	FN	Prot	W	P/L	Stab
Dataset _{CV}	ST	RF	2.715	56.816	0.406	66.053	0.517	5.89
Dataset _{CV}	MTRS	RF	2.831	60.835	0.410	73.469	0.515	6.12
Dataset _{CV}	ERC	RF	2.615	58.563	0.397	68.285	0.494	5.78
Dataset _{CV}	DSTARS	RF	2.634	60.285	0.397	66.810	0.521	5.79
Dataset _{CV}	ST	SVM	2.720	57.565	0.420	66.768	0.497	6.07
Dataset _{CV}	MTRS	SVM	2.616	56.449	0.458	71.179	0.487	5.86
Dataset _{CV}	ERC	SVM	2.584	57.401	0.432	66.767	0.479	5.83
Dataset _{CV}	DSTARS	SVM	2.563	56.788	0.417	67.313	0.482	5.93
Dataset _{CV}	ST [*]	LR	2.859	58.744	0.391	68.187	0.525	5.82
Dataset _{CV}	MTRS	LR	2.875	59.840	0.404	72.922	0.510	5.51
Dataset _{CV}	ERC	LR	2.726	59.603	0.403	69.854	0.511	5.50
Dataset _{CV}	DSTARS	LR	2.656	60.154	0.419	70.096	0.514	5.53
Dataset _p	ST	RF	2.643	64.064	0.382	74.097	0.517	5.51
Dataset	MTRS	RF	2.702	64.593	0.386	76.650	0.515	5.59
Dataset	ERC	RF	2.558	62.935	0.383	74.080	0.506	5.42
Dataset	DSTARS	RF	2.638	64.601	0.387	72.587	0.516	5.49
Dataset	ST	SVM	2.726	64.045	0.389	75.541	0.532	5.81
Dataset	MTRS	SVM	2.669	64.630	0.394	74.805	0.522	6.05
Dataset	ERC	SVM	2.633	63.978	0.385	73.915	0.519	5.79
Dataset	DSTARS	SVM	2.628	64.045	0.385	73.634	0.532	5.81
Datasetp	ST [*]	LR	2.979	66.759	0.419	74.251	0.532	5.68
Dataset	MTRS	LR	2.994	66.467	0.434	82.968	0.514	6.06
Dataset	ERC	LR	2.769	65.570	0.414	77.101	0.515	5.83
Dataset	DSTARS	LR	2.696	66.759	0.419	72.665	0.532	5.68

one technique significantly outperforms the others. On the other hand, ERC technique turned out to be superior for the majority of targets in prediction set. For $Dataset_{CV}$, SVM algorithm obtained the lowest prediction error for more targets (HlW, FN and P/L), but in the experiments conducted over $Dataset_P$ the RF algorithms obtained the lowest RMSE for all targets.

In order to compare the tested combination of MT approaches and base-learners with usual PLS, the RP values

have also been calculated considering ST + LR results as reference. The obtained RP values from $Dataset_P$ are reported in Table 3.

All the RP values are superior to usual PLS (RP values greater than 1), except for MTRS built with LR models. This result can be due to a naive MTRS modelling obtained by linear regression, since the linear inter-correlations had already been explored by PLS scores extraction. In other words, all linear dependencies between the targets were explored in the

	LR	SVM	RF	Average
MTRS	0.97	1.02	1.03	1.01
ERC	1.01	1.04	1.07	1.04
DSTARS	1.02	1.03	1.05	1.04
Average	1.00	1.03	1.05	
* Average calculated from	n ML algorithms.			
** Average calculated from	n MT techniques.			

first step of the proposed approach, and the addition of another linear inference layer does not lead to an improvement. On the other hand, by exploring several different sequential chains and different layers depth, ERC and DSTARS could improve the performances. Also, the combination of non-linear ML algorithms and MT techniques allowed to improve the performance up to 7% (ERC + RF).

Taking into account the results reported in the present study, we suggest the use of Multi-target over PLS scores modelled by a ML algorithm. Our solution was able to treat some drawbacks of PLS by non-linear ML modelling. Indeed, RF could be used to report extra information from RF importance and requires fewer hyper-parameters than SVM. In the case of a demand for high predictive power, ERC is the recommended choice.

4. Conclusion

In the present study, we explored the possibility of taking advantage from multi-target approaches for the simultaneous prediction of different quality-related parameters with NIR spectroscopy. In particular, two different aspects related to the analysis of spectral data have been jointly considered: the benefits of non-linear modelling and the possible advantages of multi-target prediction. Moreover, by means of PLS, the data dimensionality was reduced and a machine learning algorithm was applied to deal with non-linearities in NIR spectra. In addition, the results were further confirmed by a robust validation procedure, allowing to avoid overfitting. Finally, by the use of a Multi-target strategy, it was possible to overcome the actual predictive performance by accounting for the relationships between the targets. Considering all the evaluated quality parameters, this procedure allowed to obtain an increase in the predictive performance up to 7%.

Declaration of Competing Interest

The authors declare they have no significant competing financial, professional, or personal interests that might have influenced the performance or presentation of the work described in this manuscript.

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