Exploiting Data winning for Authenticity Assessment and Protection of High-Quality Italian Wines from Piedmont*

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ABSTRACT

This paper discusses the data mining approach followed in a project called TRAQUASWINE, aimed at the definition of methods for data analytical assessment of the authenticity and protection, against fake versions, of some of the highest value Nebbiolo-based wines from Piedmont region in Italy. This is a big issue in the wine market, where commercial frauds related to such a kind of products are estimated to be worth millions of Euros. The objective is twofold: to show that the problem can be addressed without expensive and hyper-specialized wine analyses, and to demonstrate the actual usefulness of classification algorithms for data mining on the resulting chemical profiles. Following Wagstaff's proposal for practical exploitation of machine learning (and data mining) approaches, we describe how data have been collected and prepared for the production of different datasets, how suitable classification models have been identified and how the interpretation of the results suggests the emergence of an active role of classification techniques, based on standard chemical profiling, for the assesment of the authenticity of the wines target of the study.

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1. INTRODUCTION

The quality and safety profiles of quality wines represent a peculiar case of the notion of *food integrity*, because of the very high value of a single bottle, and because of the complex chemical profile, requiring therefore specific and robust methods for their univocal profiling/authentication. Vitis vinifera is the unique grape allowed for the winemaking, but many different genetic varieties (e.g. Pinot, Nebbiolo, Merlot, Sangiovese, Sirah and many others) lead to wines with different character and chemical profiles. The industrial processing largely build the wine specificity.

Moreover, the "terroir" (the set of special characteristics that the geography, the geology and the microclimate of a certain region or peculiar location, interacting with grape genetics, express in wine), while bringing to the diversification of the product, complicates significantly the wine chemical profiling under the metabolomics profile and, thus, the process of traceability and identification.

Although specific regulations exist in this matter, and some analytical approaches and protocols are well established for wine tracking and authentication, quality wines are highly subjected to adulteration. Problems with consumption fraud go back a very long time; more or less recently, numerous cases of wine adulteration were worldwide registered, including Austria, France and Italy. Wine fraud is then a big issue worldwide, inducing significant problems for consumers; it also triggers destabilization of the wine market, particularly regarding the quality aspect, with an estimated impact of about 7% of the whole market value.

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Although wine can suffer for different type of frauds and different type of counterfeiting, the predominant problem in wine sector is mislabeling, regarding both the used cultivar of grape and the geographical area of origin; it causes an economical impact estimated to be several million of Euros.

The detection of adulterations or declarations which do not correspond to the labeling are actually official tasks of wine quality control and consumer protection. During the last years, analytical methods have been improved in this field. Some of them (stable isotope ratio analysis by nuclear magnetic resonance, and isotope ratio mass spectrometry) have been adopted as official methods by the European Community (EC). In addition, the use of the corresponding analytical devices is expensive and requires a lot of specialization. Moreover, these methods, while playing a key role in detecting adulterations like addition of water and inadmissible sweetening, represent a challenge also regarding the identification of origin, requiring in this case a very large data base.

Non analytical approaches like olograms, trasponder systems or QR codes only partially address the problem of wine authenticity. Finally, regulations, good manufacturing practices and traceability procedures are not always sufficient to guarantee the wine authenticity, considering the falsification of both varietal and geographic origin. For these reasons, a challenge for wine authenticity is to obtain standard analytical procedures to describe a wine proof-of-identity, which could defend consumers from illegal adulteration practices, as well as from unintentional mislabeling due to mistakes during wine production.

There are then two analytical levels: the *chemical* one, concerning the identification of the most relevant standard chemical analyses to perform, and the *data* one, concerning the determination of the relevant data mining and analysis techniques exploiting the features obtained from the chemical level. From the data mining perspective, the main goal of the work is to show that well-established methodologies of classification, from machine learning, can be suitably adopted to fullfill the task of controlling specific wine adulterations, and consequently be capable of generating added value to the quality control process of such high-quality wines.

In the paper we will discuss the experience gained in the TRAQUASWINE project, having as a major goal the assessment of the authenticity and the protection against fake versions of some of the highest quality (and often top priced) *Nebbiolo-based* wines like *Barolo, Barbaresco* and *Gattinara*. Following Wagstaff's scheme [16], we discuss how data have been collected, how they have been prepared, how suitable classification models have been identified and how the interpretation of the results suggests the emergence of an active role of classification techniques, based on standard chemical profiling, for the assessment of the authenticity of the high-quality wines which were target of the study.

The remainder of the paper is organized as follows: section 2 presents the main objectives and the data collection and preparation steps of the study; section 3 discusses the characterization of the data and of the classification algorithms adopted, by presenting results on different datasets with different sets of features; section 4 compares the approach to related works finally, section 5 presents the conclusions, by pointing out to some future evolutions.

2. THE TRAQUASWINE STUDY

2.1 Main Objectives

The TRAQUASWINE project involved both industrial and academic partners in the creation of a network operating in the wine sector, with the aim to trace and authenticate the origin, quality and safety of Piedmont (Italy) wines, with particular interest to high-value and high-quality *Nebbiolobased* wines. In this context, the principal goals of the project were:

- to define the major characteristics (*chemotype*, with particular interest on bioactive compounds) of Nebbiolo grape and derived wines, identifying molecular markers useful for quality control and wine traceability;
- to establish a comprehensive analytical approach, based on the exploitation of data mining techniques for multivariate analysis to chemotype datasets, useful to the traceability and authentication of high-quality *Nebbiolobased* wines, produced in different area of Piedmont, by diverse wineries.

The selection of chemical parameters and methods was performed by taking into account two main criteria:

- economical cost of the analyses (particularly regarding the required instrumentation);
- the capacity to work also with common chemical parameters, avoiding the need of pluriannual consolidated databases, as often required by classical isotopic methods.

Important feature has been the use of data of different origin, by considering different producers, different areas of production, different typology of wine (i.e., commercial wines on the market, as well as "model" wines explicitly produced for the study) and different aging of the product. A peculiar approach (use of some *Nebbiolo* wines spiked with different percentages of different wines) was finally exploited in order to check the possibility to highlight the illegal adding of a foreign grape cultivar, so detecting potential frauds.

2.2 Data Collection

Commercial samples of Piedmont Nebbiolo-based wines were provided by nine local wineries; producers themselves certified wine's origin and identity. Eight different types of wines at different aging degree (selected on the basis of the mean aging of each type of wine and depending on their commercial availability) were considered. Among them, Barbaresco (BRB), Barolo (BRL), Langhe (LAN), Nebbiolo d'Alba (NEB) and *Roero* (ROE) are typical wine productions of the Alba district in Southern Piedmont, while Gattinara (GAT), Ghemme (GHE) and Sizzano (SIZ) are produced in Northern Piedmont (Novara province). The distribution of samples (spanning a period of 10 wine years) resulted in 18 samples of BRB, 18 samples of BRL, 12 samples of LAN, 18 samples of NEB, 6 samples of ROE, 6 samples of GAT, 18 samples of GHE and 6 sample of SIZ. BRL, BRB, GAT, NEB and ROE were certified as produced from 100%Nebbiolo grape, while the other wines contained percentages of different grape varieties (having however Nebbiolo as the primary grape variety).

Beside such commercial wines, a set of experimental wines (model wines) has been specifically prepared, resulting in 56 additional samples. Among them, 12 samples correspond to wines produced without Nebbiolo grape, and in particular with 100% of some grape varieties partially present in some of LAN commercial wines; we labeled them as NON (*NO Nebbiolo*); 26 samples were prepared as more LAN and NEB wines, for a final count (commercial and experimental) of 22 samples of LAN and 34 samples of NEB; 8 samples were prepared as a correct mix for GHE, resulting in a total of 26 samples of GHE. Finally, the remaining experimental wines were produced to model incorrect (i.e., not allowed by the disciplinary of production) blends for GAT wine, resulting in 10 more samples that we labeled as BLE (*Blend*).

As mentioned earlier, the main goal of the study was to develop and assess analytical techniques (both at the chemical and data levels) to effectively support the authenticity of high-value Nebbiolo-based wines. To this end, we organized the study as follows: we selected some of the most valued 100%-nebbiolo wines as the high quality class to be protected from fakes, and in particular BRL, BRB and GAT; we also selected LAN and BLE as control wines, the former to simulate not allowed blend for BRL and BRB, and the latter for GAT. We obtained a total of 146 samples with 9 types of wines (i.e., the classes) that we have used to train and test our models. We reserved the 12 samples of NON wines as additional control wines, to test the response of the learned models with respect to simulated fake wines with the absence of Nebbiolo grape, and with the additional complication that no such a kind of wines have been used in the training of the models.

2.3 Data Preparation

Wine samples were mainly characterized for their phenolic composition. The phenolic component of wine is very complex and heterogeneous, and is strongly influenced by several factors related to grape polyphenol composition winemaking procedures, wine aging, and storage conditions. In the present study, we decided to employ common and relatively economic analytical techniques, as well as methods easily reproducible. The adopted analytical (chemical) procedures can be categorized as follows:

- *spectrophotometric methods*, which include not specific and low-cost assays, useful to quantify general class of compounds (total polyphenols, total anthocyans and total tannins) and antioxidant activity;
- chromatographic methods, which are more advanced and specific, but are time-consuming and require more expensive instrumentations, which are however generally available in the standardly equipped laboratories for quality control. The chromatographic techniques were employed for the identification and quantification of individual polyphenol compounds in wines.

Considering spectrophotometric methods, the total phenol content is an unspecific parameter; however, due to the complexity of wine's phenolic component, it can be considered an interesting general information. Total anthocyans and total tannins are other peculiar characteristics of wines, related to their organoleptic properties. Antioxidant activity, for which do not exist official methods recommended by international organization for wine analysis , was measured using a rapid chemical model system for the determination of antiradical properties (DPPH assay).

The chromatographic techniques are useful tools to fingerprint wine polyphenols, identifying individual molecules. In this study, different class of compounds were considered: anthocyanins, phenolic acids, hydroxycinnamates and flavonoids. Some of them have been calculated not only as absolute concentrations, but also as relative percentages.

Finally, wine samples were analyzed for the content of resveratrol (both *trans* and *cis* form), important molecule for its beneficial effects on human health, and considered as one of the major compounds responsible for the well-known French Paradox (the fact that French people, having red wines in their dietary habits, have a relatively low incidence of coronary heart disease, while having a diet relatively rich in saturated fats).

3. THE DATA MINING FRAMEWORK

3.1 Data Characterization

From the chemical analyses described in section 2.3, a total of 40 continuous features have been extracted. Besides those described in section 2.3, also wine acidity (the pH) and information about the percentage of some grape varieties have been added. Missing values were present in about 1/3 of the dataset, essentially in association with the latter set of attributes (percentage of grape varieties). As reported in section 2.2, we have then obtained a dataset of 146 instances, each one corresponding to a particular class (i.e. the wine type) out of 9 possible classes, and having 40 numerical (continuous) attributes.

We have organized the evaluation of the data mining experimental study as follows. First of all, we have searched for regularities in the profiles of the available wine chemical analyses by means of some *clustering* algorithms. In particular, we tested EM clustering (both with a free number of clusters and with a number of required clusters equal to the number of classes) and a *K*-mean algorithm (K = 9). As expected, no definite interpretation of the results have emerged from unsupervised data analysis, suggesting that regularities associated to wines should be better captured through supervised methods.

We then considered classification approaches, by having in mind that the goal of the data mining process was mainly to characterize the misclassification of the target wines, that is high-quality wines (BRL, BRB and GAT) and control wines (LAN and BLE). Thus general classification accuracy has been considered of secondary importance. To test classification approaches, we considered three different data sets: dataset D1 containing all the 146 instances with the whole set of 40 features; dataset D2 containing the 146 instances with a reduced set of 15 features, obtained by automatic feature selection; dataset D3 containing the 146 instances with the set of features of D2 manually reduced to 13 attributes, and in particular by removing all the attributes concerning the percentage of grape varieties which survived automatic feature selection.

We have experimented the learning of the following type of classifiers:

1. A Bayesian Network classifier (BN), with learning performed with standard Cooper/Herskovits algorithm [3], allowing a maximum of 3 parents per node, identifying

	Accuracy	Kappa
BN	87%	0.85
MLP	94%	0.93
SMO	92%	0.91

 Table 1: Accuracy and KAPPA statistic for dataset

 D1.

dependencies among the attributes and thus avoiding the Naive Bayes assumption.

- 2. A Support Vector Machine (SVM) based on Sequential Minimal Optimization algorithm (SMO) [10] with a Pearson Universal Kernel (PUK) having Lorentzian peak shape [14], and with Platt scaling in such a way to get a probability distribution over the classes [11]¹.
- 3. A Multi-Layer Perceptron (MLP) with one hidden layer of $n = \frac{f+c}{2}$ hidden units (being f the number of features and c the number of classes)

We also experimented with a lazy learner based on 1-NN classification, and with a Decision Tree learner based on the C4.5 algorithm; since this two types of classifier did not perform well for our task, we do not report the corresponding results in the paper.

The BN, SMO and MLP classifier have been learned using the 146 samples and evaluated by means of a 10-fold cross validation. As mentioned before, the main focus was not on general accuracy (even if definitely an important measure), but rather to check the performance of the classifiers with respect to the classes of high-quality wines (BRL, BRB and GAT) and control wines (LAN and BL), by looking at their possible misclassifications.

We have exploited WEKA's algorithms to perform our experimentation[7], and multi-class classification has been performed as a *one-agaist-all* approach. In the following subsections we will discuss the results obtained on each of the above datasets.

3.2 Complete Feature Set

In this part we discuss the results obtained by using the whole set of features obtained after the chemical analyses performed on the available wine samples (dataset D1). Table 1 reports the general accuracy and KAPPA statistic of the three classifiers. It can be noticed that the general accuracy (and its significance as measured by KAPPA statistics) is rather high, with the MLP classifier being the most accurate and slightly better than SMO. Figure 1, figure 2 and Figure 3 report the prediction statistics with respect to the classes of interest for BN, MLP and SMO classifier respectively. We also report Matthews Correlation Coefficient (MCC) and F1-measure, with MCC being more significant in our case, since the number of "positive" instances (those of the target class) are much less that the number of "negative" instances (those of all the other classes). Concerning control wines (LAN and BLE), we can notice that classification accuracy is very good, but more importantly, no control wine is misclassified as a high-quality one; on the other hand,

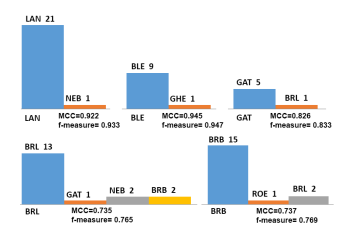


Figure 1: Predictions BN classifier, dataset D1

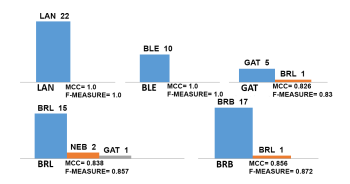


Figure 2: Predictions MLP classifier, dataset D1

high-quality wines (BRL, BRB and GAT) are also recognized very well, with no misclassification predicting control wines. From the diagrams, we can also notice that each misclassification is also not really surprising: for instance in every classifier, BRB is misclassified either as BRL (a very similar wine from the point of view of origin, production and grape composition) or as ROE, which is another 100% Nebbiolo wine. MLP has a perfect performance on control wines (no misclassification). SMO has a misclassification on BLE (but with the other control class LAN) and a perfect classification on BRL (definitely the most valued wine of the set).

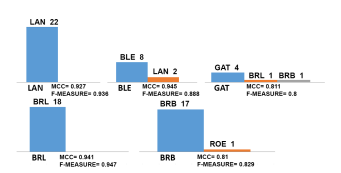


Figure 3: Predictions SMO classifier, dataset D1

¹The regularization parameter (a.k.a complexity) has been set to 10, since we are dealing with a multi-class problem with 9 different classes, and it is a good practice to set the parameter close to the number of classes [9].

	I	Probabi	lity di	stributi	on of m	isclassif	ied prec	lictions	
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE
LAN	0	0	0.02	0.914	0	0.056	0.024	0.002	0.002
BLE	0.893	0	0	0	0	0	0	0	0.107
GAT	0	0.055	0	0.001	0.944	0	0	0	0
	0	0.827	0	0.015	0.158	0	0	0	0
	0	0.019	0	0.918	0.015	0.048	0	0	0
BRL	0	0.001	0	0.976	0.006	0.017	0	0	0
	0	0	0	0.002	0.351	0	0.647	0	0
	0	0	0	0	0.299	0	0.701	0	0
	0	0	0	0.001	0	0.898	0.101	0	0
BRB	0	0	0	0	0.971	0	0.029	0	0
	0	0	0	0.192	0.736	0	0.072	0	0

Table 2: Probability distribution of misclassifications, dataset D1: BN classifier.

	P	Probability distribution of misclassified predictions									
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE		
GAT	0.003	0.277	0	0.266	0.449	0.001	0.003	0	0		
	0	0	0	0.585	0.408	0.007	0	0	0		
BRL	0	0	0	0.559	0.437	0.003	0	0	0		
	0.089	0.600	0	0.008	0.302	0.001	0	0	0		
BRB	0	0.003	0	0.001	0.933	0.003	0.059	0	0		

Table 3: Probability distribution of misclassifications, dataset D1: MLP classifier.

Considering the classes of interest, we also computed, for each classifier, the probability distribution on the predictions that are misclassified. Table 2, table 3 and table 4 show the results for classifiers BN, MLP and SMO respectively. The prediction performed by the classifier corresponds to the entry shown in **boldface** (the largest probability of prediction). By inspecting such distributions, we confirm the good performance of the classifiers, with MLP and SMO being almost equivalent and sligthly better than BN. All classifiers agree with a very high confidence that a high-quality wine cannot be confused with a control wine (see last two columns of the tables in the rows of high-quality wines). We can also notice that distributions obtained from MLP and SMO are in general less skewed than those of BN, showing more "caution" in the actual prediction, but also very good performance on predicting control wines. For instance SMO, having two misclassification on BLE, predicts BLE as the second most probable class and estimates a probability close to 0 to highquality wines (exactly 0 for GAT).

As a final investigation on D1, we tested the classifiers using a test set TS composed by the 12 instances of NON wines resulting as discussed in section 2.2. Differently from the wine samples used for training the classifiers, the wine samples in TS do not contain any part of Nebbiolo grape; this implies that acceptable predictions can be those selecting our control wines (LAN and BLE) which are the ones containing parts of the grape varieties contained in the wine samples of TS; in particular, LAN is also a preferred predictions than BLE, because of the way NON wine samples have been prepared. The results are shown in figure 4. SMO shows very reasonable predictions, by identifying very precisely the only class (LAN) containing percentages of the grapes present in the test cases of TS. By considering all the results, we can conclude that, on dataset D1, SMO appears to be the best globally performing classifier.

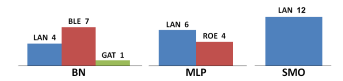


Figure 4: Predictions of NON wines, dataset D1

3.3 Automatic Feature Selection

The second round of experiments we have performed takes into account the relevance of the available features of the wine samples, with respect to our target classification task. To this extent, we constructed and evaluated the same set of classifiers considered in section 3.2, but on a dataset with a reduced set of features. In particular, we have performed a correlation-based feature selection with sequential forwardbackward selection on the original feature space [12, 8], resulting in the selection of 15 analytical attributes out of 40^2 . In particular, all the features representing grape variety percentage, but those of Nebbiolo and Uva Rara were removed by feasture selection, together with a set of redundant information about polyphenols that were initially provided both as absolute concentrations and relative percentages. As already mentioned, the resulting dataset will be refferred as D2. Table 5 reports the general accuracy and KAPPA statistic of the three classifiers for dataset D2. Results show again a good and significant global accuracy, this time with SMO slightly better than MLP. Figure 5, figure 6 and Figure 7 report the prediction statistics with

 $^{^{2}}$ An approach based on a PCA transformation of the feature space has also been investigated in the study, with results and considerations similar to those reported in the present papers for the other alternatives.

	P	Probability distribution of misclassified predictions									
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE		
BLE	0.004	0	0	0.117	0.016	0	0.013	0.599	0.251		
	0.005	0	0	0.154	0.018	0	0.027	0.620	0.176		
GAT	0.036	0.254	0	0.010	0.624	0	0.073	0.002	0.001		
GAI	0.014	0.342	0	0.021	0.187	0	0.433	0.002	0.001		
BRB	0	0	0	0	0.989	0	0.011	0	0		

Table 4: Probability distribution of misclassifications, dataset D1: SMO classifier.

	Accuracy	Kappa
BN	83%	0.80
MLP	89%	0.87
SMO	90%	0.88

Table 5: Accuracy and KAPPA statistic for datasetD2.

respect to the classes of interest for BN, MLP and SMO classifier respectively. Concerning control wines, we can

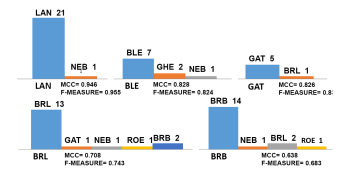


Figure 5: Predictions BN classifier, dataset D2

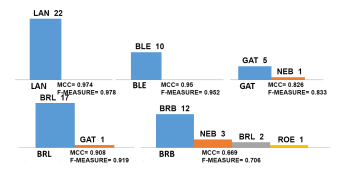


Figure 6: Predictions MLP classifier, dataset D2

notice again a very good classification with high accuracy and no control wine misclassified as a high-quality one; on the other side, high-quality wines are also recognized well (SMO has a perfect prediction on GAT), with no misclassification predicting control wines. As in the previous case, misclassifications can in general be suitably justified: the most interesting result is provided by SMO, where BRL and BRB are each other confused in the corresponding misclassification (which is something definitely acceptable).

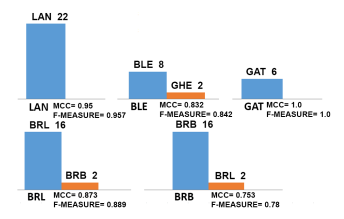


Figure 7: Predictions SMO classifier, dataset D2

As in the case of dataset D1, we also computed, for each classifier, the probability distribution on the predictions that are misclassified, by focusing on the classes of interest. Table 6, table 7 and table 8 show the results for classifiers BN, MLP and SMO respectively. Again, the prediction performed by the classifier corresponds to the entry shown in boldface.

A problem can be noticed in the BN classifier, where the first misclassification of BRB has a a prediction on BLE with a 29% probability, even if the most probable predicted class is NEB. MLP is showing no misclassifications for control wines, even if there are 6 misclassifications of BRB; however, three of them have exactly BRB as the second most probable prediction (with a 48% probability in one case and more than 30% in the others), while the predictions of control wines are either 0 or very close to 0. Concerning SMO, the confusion of BLE with GHE is justified by the fact that samples corresponding to such instances represented mixtures very close to the blend allowed for GHE, while confusion of BRL with BRB and vice versa are justified by the very high similarity of the two wines (as already mentioned).

Finally, we tested the classifiers learned using D2 using the test set TS of NON wines, resulting in the predictions graphically reported in figure 8. BN's prediction appears



Figure 8: Predictions of NON wines, dataset D2

		Probabi	ility dis	tributio	on of mi	sclassifi	ed pred	ictions	
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE
LAN	0	0.002	0.017	0.917	0	0.001	0.058	0	0.004
	0.704	0	0	0	0	0	0	0	0.295
BLE	0.620	0	0	0	0	0	0	0	0.380
	0.061	0	0	0.613	0	0	0	0	0.326
GAT	0	0.017	0	0.101	0.873	0.001	0.008	0	0
	0.006	0.580	0	0.007	0.406	0	0	0	0
	0	0	0	0.727	0.272	0	0	0	0
BRL	0	0.006	0.001	0.066	0.270	0.467	0.188	0	0
	0	0	0	0	0.442	0	0.558	0	0
	0	0	0	0	0.227	0	0.772	0	0
	0	0.007	0	0.515	0.056	0	0.128	0.006	0.287
BRB	0.002	0.001	0	0.080	0.649	0	0.254	0	0.013
	0	0	0	0	0.968	0	0.032	0	0
	0	0	0	0.009	0	0.991	0	0	0

Table 6: Probability distribution of misclassifications, dataset D2: BN classifier.

	F	Probabi	lity di	stributi	on of m	isclassif	ied pre	diction	s
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE
GAT	0.002	0.270	0	0.711	0.002	0	0.014	0.001	0
BRL	0.068	0.475	0	0.008	0.422	0	0.026	0	0
	0	0.012	0	0.637	0.001	0	0.349	0	0
	0	0	0	0.575	0.003	0.41	0	0	0.011
BRB	0	0.003	0	0.516	0	0.001	0.48	0	0
DIUD	0.007	0	0	0	0.691	0	0.302	0	0
	0	0.015	0	0.001	0.952	0.023	0.009	0	0
	0	0	0	0.195	0.007	0.773	0	0	0.025

Table 7: Probability distribution of misclassifications, dataset D2: MLP classifier.

	Accuracy	Kappa
BN	79%	0.75
MLP	82%	0.79
SMO	89%	0.87

 Table 9: Accuracy and KAPPA statistic for dataset

 D3.

more reasonable than in the case of D1 (no high-quality wine prediction, even if NEB predictions are not very well justified); MLP's predictions have a problem in the two cases where the sample is recognized as a BRB (in both cases with a rather high probability); SMO's predictions are quite good, even if NEB predictions shows an identification problem on the corresponding samples.

To summarize, also in this situation (i.e., using D2 for training), SMO resulted to be performing in general better than the other classifiers.

3.4 Automatic and Manual Feature Selection

A third round of experiments concerned the manual elimination, from the dataset D2, of the information concerning the percentage of grape variety. This resulted in another dataset (D3) with a total of 13 attributes. Table 9 reports the general accuracy and KAPPA statistic of the three classifiers for dataset D3. The accuracy of each classifier slightly reduces again with respect to the previous cases, and SMO results better than MLP, with the lowest accuracy obtained by BN (as in the use of D2). Figure 9, figure 10 and Figure 11 report the prediction statistics with respect to the classes of interest for BN, MLP and SMO classifier respectively. As in the previous cases, no control wine is predicted

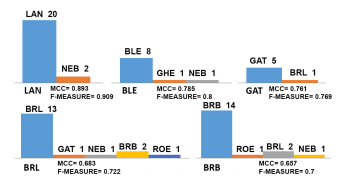


Figure 9: Predictions BN classifier, dataset D3

as a high-quality wine and vice versa. SMO has exactly the same (good) performance as in the case of D2, and also BN produce almost the same results. MLP slightly augment the number of misclassifications, but is still performing better than BN. Thus, the complete elimination of the grape percentages from the dataset does not seem to influence too much the cross validation performance of the classifiers we considered (it is worth noting that in 1/3 of the samples such data were already missing).

	Pr	Probability distribution of misclassified predictions									
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE		
BLE	0.990	0	0	0	0	0	0	0	0.009		
BLE	0.962	0	0	0	0	0	0	0	0.037		
BRL	0	0	0	0.002	0.199	0	0.799	0	0		
DILL	0	0	0	0.004	0.358	0	0.638	0	0		
BRB	0	0	0	0	0.936	0	0.063	0	0.013		
	0	0	0	0.001	0.936	0	0.063	0	0		

Table 8: Probability distribution of misclassifications, dataset D2: SMO classifier.

		Probab	ility dis	stributio	on of mi	sclassifi	ed pred	ictions	
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE
LAN	0	0	0	0.566	0	0.002	0.002	0.43	0
	0	0	0.005	0.955	0	0.001	0.032	0.001	0.005
BLE	0.611	0	0	0.002	0	0	0	0	0.387
	0.037	0	0	0.632	0.001	0	0	0.015	0.314
GAT	0	0.017	0.001	0.08	0.893	0	0.008	0	0
	0.209	0.419	0.013	0.004	0.354	0	0	0	0
	0	0	0.001	0.718	0.281	0.	0	0	0
BRL	0.011	0	0	0	0.481	0	0.508	0	0
	0	0	0	0	0.237	0	0.763	0	0
	0	0.037	0.051	0.151	0.313	0.324	0.119	0.005	0
	0	0	0	0.009	0	0.991	0	0	0
BRB	0.072	0	0	0.035	0.846	0	0.044	0	0.004
	0	0	0	0	0.979	0	0.021	0	0
	0.01	0.012	0.002	0.611	0.086	0	0.186	0.009	0.084

Table 10: Probability distribution of misclassifications, dataset D3: BN classifier.

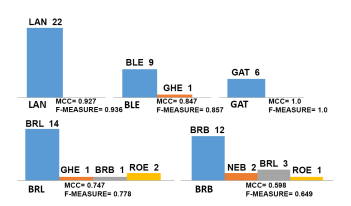


Figure 10: Predictions MLP classifier, dataset D3

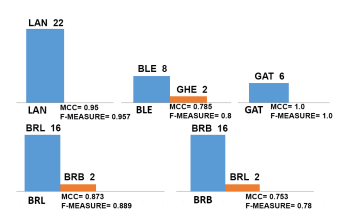


Figure 11: Predictions SMO classifier, dataset D3

Again, as in the case of other datasets, we computed, for each classifier, the probability distribution on the predictions that are misclassified, by focusing on the classes of interest. Table 10, table 11 and table 12 show the results for classifiers BN, MLP and SMO respectively (prediction performed by the classifier corresponding to the boldface entry). It can be noticed the very good performance of SMO, that in case of BRB and BRL (the only high-quality wines having misclassifications) shows a null probability of predicting control wines and in one prediction of BRB (the second misclassification of BRB) has a 31% probability of predicting the correct wine (against a 64% probability of confusing it with the similar BRB). Finally, predictions on BLE (the only misclassified control wine) shows a null or close to null probability of predicting high-quality wines.

Also in this case we finally tested the classifiers learned with D3, using the test set TS of NON wines; results are summurized in figure 12. These results shows that SMO is definitely more robust and stable with respect to the others in presence of control wines on which the classifier was not trained; this proposes, together with the previous results, SMO as the best performing approach on D3 as well.

To conclude, the results of the experiments suggest that standard chemical profiling of Piedmont *Nebbiolo-based* wines, coupled with data mining classification techniques, can be a powerful tool to authenticate high-quality and high-value

		Probab	ility di	stributi	on of m	isclassifi	ied prec	lictions	
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE
BLE	0.471	0	0.076	0	0.005	0	0	0	0.447
	0.862	0.095	0.001	0.001	0.039	0	0	0.002	0
BRL	0.383	0.012	0	0	0.014	0	0.518	0	0.072
	0	0	0.21	0	0.064	0.69	0.036	0.001	0
	0	0	0.118	0	0.278	0.59	0.014	0	0
	0	0	0.025	0.641	0.233	0.095	0	0.001	0.004
	0	0	0.014	0.748	0	0.233	0	0.001	0.005
BRB	0.004	0.109	0	0.008	0.543	0	0.335	0	0
DIID	0	0.041	0.001	0	0.591	0.002	0.365	0	0
	0	0.004	0	0.009	0.878	0.002	0.107	0	0
	0	0	0.035	0.327	0	0.626	0	0.002	0.01

Table 11: Probability	[,] distribution	of misclassifications,	, dataset D3:	MLP classifier.
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	Probability distribution of misclassified predictions								
True Ground	GHE	GAT	SIZ	NEB	BRL	ROE	BRB	LAN	BLE
BLE	0.99	0	0	0	0	0	0	0	0.01
	0.747	0	0.059	0.01	0	0	0.002	0	0.182
BRL	0.001	0	0.043	0	0.16	0	0.795	0	0
	0.002	0	0.052	0.001	0.309	0	0.636	0	0
BRB	0.001	0	0	0	0.936	0	0.063	0	0
	0	0	0	0.001	0.937	0	0.063	0	0

Table 12: Probability distribution of misclassifications, dataset D3: SMO classifier.

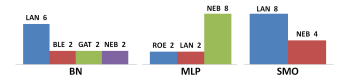


Figure 12: Predictions of NON wines, dataset D3

wines. All the tested classifiers performed rather well with respect to the objectives of the work, with BN being the more problematic in some situations, and MLP comparable in performance with SMO, the latter showing a better robustness with respect to possible fake wines.

4. RELATED WORKS

Wine classification and identification through data mining and pattern recognition using chemical descriptors have been approached several times in different settings. Several paper are reported in literature, focused on different analytical targets and techniques, where however the complexity of the data mining task is often proportional to the complexity of the chemical or sensory analyses. Moreover, employed data mining techniques cover essentially all the possible classification or discriminant analysis methods, showing that no "silver bullet" is available in general [2, 15].

Authors in [5] investigate aroma compounds of Galician white wines using a dataset of 42 samples; they show that PCA was not able to accurately separate all the wine varieties, thus different machine learning techniques (SVM, Random Forest, MLP, k-NN and Naive Bayes) were evaluated, obtaining a perfect classification accuracy using the Random Forest algorithm. Differently from our study, the target was maximizing general classification accuracy, and no commercial wine was analyzed.

PCA and Extreme Learning Machine (ELM) were also applied to detect grape varieties: the PCA algorithm was adopted to process chemical components of publicly available wine datasets and classification was performed using the ELM. The experimental results show that the proposed model was useful for general wine classification [17].

Sophisticated analytical methods (even coupled with specific sensory devices such as electronic nose or tongue) have been often used to obtain wine characterization; however, as in our case, [1] shows that simple and nonselective techniques (as UV-visible spectrophotometric methods), in combination with classification approaches, can be demonstrated successful. Here the emphasis is again in improving general classificational accuracy (in this case of several Spanish wines).

Finally, another recent work proposing support vector machine as very promising in this context is presented in [4]; the task here is however completely different, since the goal is to predict (with a suitable score) human wine tasting preferences (that are however closely related to the wine type characteristics).

5. DISCUSSION AND FUTURE WORKS

In this paper we have reported of the results of a study exploiting classification for the assessment of the authenticity of some high-value Italian wines. The problem has been addressed without expensive and hyper-specialized wine chemical analyses, and by learning and suitably evaluating different standard classifiers on the resulting chemical profiles. The proposed approach can be regarded as an instance of the 3-phases Wagstaff's scheme [16]: *Necessary Preparation* has involved the definition of the analyses for data collection and preparation, Machine Learning Contribution has regarded the selection of a supervised strategy and of a suitable set of classifiers, while Impact has concerned the evaluation of the classification results with respect to the study's objectives. The conclusion is that an SVM classifier like SMO emerges as very promising in this context, both for detecting wrong or illecit blends (not necessarily of low quality, but not compliant with regulations as in the case of LAN and BLE), as well as for avoiding incorrect introductions of unrelated cultivars (as in the case of NON). Moreover, feature selection has pointed out that the number of chemical parameters, needed to get good performances with respect to the authentication objective, can be significantly less than the whole set of parameters initially tought as necessary: performances on datasets D2 and D3 of SMO are in fact comparable, and in some sense even better than those on dataset D1. For instance, in D1, GAT has been misclassified by SMO as BRL and BRB, which are wines similar to GAT, but from different location (Northern vs Southern Piedmont area). Therefore, we can conclude that even a reduced number of chemical analyses (and so a less expensive strategy), together with a suitable classification methodology can be really effective in the considered task.

A weak point of the study could be the not very large number of available samples; however, compared with similar studies, the sample size used in the work is definitely adequate and larger than the usual ones. The practical problem is that obtaining such samples is rather demanding, even if stardard analytical techniques are used. We plan to exploit approaches for generating syntethic data (from the real ones available), in such a way to extend the evaluation of the classifiers to a larger set of data. We have already obtained some interesting preliminary results by learning, using PC algorithm [13], class-specific Linear Gaussian Bayesian Networks [6] as generative models for such synthetic data. The experimental evidence suggests that a data mining classification tool, on wine chemical profiles, can be suitably proposed to address the wine quality control issue.

6. ACKNOWLEDGMENTS

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