



Predicting supply chain risks using machine learning: The trade-off between performance and interpretability

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HIGHLIGHTS

- Machine learning models can be leveraged to accurately predict supply chain risks.
- Choosing more interpretable models may require a compromise in performance.
- Performance is slightly more affected in the case of prediction-related metrics.
- Decision tree models can reveal correlations that influence SCRM decision-making.

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ABSTRACT

Managing supply chain risks has received increased attention in recent years, aiming to shield supply chains from disruptions by predicting their occurrence and mitigating their adverse effects. At the same time, the resurgence of Artificial Intelligence (AI) has led to the investigation of machine learning techniques and their applicability in supply chain risk management. However, most works focus on prediction performance and neglect the importance of interpretability so that results can be understood by supply chain practitioners, helping them make decisions that can mitigate or prevent risks from occurring. In this work, we first propose a supply chain risk prediction framework using data-driven AI techniques and relying on the synergy between AI and supply chain experts. We then explore the trade-off between prediction performance and interpretability by implementing and applying the framework on the case of predicting delivery delays in a real-world multi-tier manufacturing supply chain. Experiment results show that prioritising interpretability over performance may require a level of compromise, especially with regard to average precision scores.

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1. Introduction

Managing risks in supply chains at a local, national, or global scale has increasingly attracted attention from both researchers and practitioners in recent years, owing in part to the worldwide economic uncertainty that began with the 2008 global financial crisis. The field of supply chain risk management (SCRM), which emerged in the early 2000s has now become more than the overlap of directly related areas such as enterprise risk management and supply chain management [1]. As defined in [2], SCRM “encompasses the collaborative and coordinated efforts of all parties involved in a supply chain to identify, assess, mitigate and monitor risks with the aim to reduce vulnerability and increase robustness and resilience of the supply chain, ensuring profitability and continuity”.

The wide range of decisions and actions that are involved in SCRM have led to an equally wide spectrum of solutions proposed by researchers. These can be broadly classified in three categories: (1) multiple-criteria decision analysis techniques; (2) mathematical modelling and optimisation; and (3) Artificial Intelligence (AI) techniques. The first category includes well-established techniques to evaluate different risk-related criteria that affect supply chains, as well as the efficiency of potential solutions, such as analytic hierarchy process [3] or data envelopment analysis [4]. The second category is by far the most common one (as analysed in [2,5]) and includes approached based on stochastic or fuzzy programming and robust optimisation.

AI techniques have received relatively little attention in relation to SCRM or supply chain research, in general. Recently, there has been an AI resurgence due to the availability of increased computing power and large amounts of data, as well as the success of approaches within the broad area of machine learning. This has also led to SCRM researchers considering the potential of

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AI techniques in relation to tasks such as risk identification, prediction, assessment and response [6–9]. However, research is still at early stages, proposing either purely theoretical frameworks that have not been implemented and applied in real-world case studies [6,7], or ad-hoc solutions that are only applicable within the confines of a particular case study [8,9]. Also, these works do not take into account the data imbalance which is inherent in risk-related tasks, since for many risks the ratio of occurrence and non-occurrence is far from balanced.

More importantly, research that uses data-driven AI for SCRM does not take into account the importance of interpretability of results: no conclusions are derived in relation to the reasons behind the results of the machine learning models they create. As stated by Doshi-Velez and Kim [10], most real-world tasks addressed using machine learning techniques cannot be described merely through a result of a single metric, such as classification accuracy. Similarly, Domingos [11] posits that instead of accuracy and computational cost, learners should be evaluated based on how much human effort is saved or how much insight is gained. Some of the reasons cited by Molnar [12] in support of interpretability are directly relevant to SCRM: (1) finding meaning withing and gaining the knowledge captured by machine learning models; (2) detecting bias in models; and (3) increase acceptance of produced solutions. In the context of supply chains, we argue that in order for any results coming out of AI solutions to be useful and able to be integrated in SCRM-related decision-making processes, they have to be interpretable and justifiable.

The research hypothesis investigated in this article is the possibility of utilising machine learning technologies to provide predictive analytics for SCRM that deliver results that are simultaneously interpretable and of a high prediction performance standard. In that sense, we aim to contribute to research in exploring the untapped potential of data-driven AI techniques within SCRM, as identified in [2]. We address the aforementioned limitations by focusing on the particular task of predicting supply chain risks and propose a generic data-driven risk prediction framework that takes into account the special characteristics of SCRM. The proposed framework is then implemented and applied on a real-world case study, investigating a variety of metrics and two well-known machine learning algorithms, one less and one highly interpretable: support vector machines (SVM) and decision tree learning. In the particular case study, risk prediction is achieved through classification. Classifiers are trained to determine whether a delivery is late or not and are then used on unseen data to predict whether future deliveries will be late or not, in other words whether there is a risk of delayed delivery or not. The novelty of the presented research lies not in the employed algorithms which are well-established and whose choice is indicative, but rather in the manner in which such technologies are to be integrated in an SCRM process. Specifically, the contributions of this article are the following:

- A data-driven risk prediction framework that places emphasis on the synergy between AI and supply chain experts, especially with regard to the proper selection of suitable metrics and algorithms according to the goals and priorities of the supply chain. The proposed framework also takes into account imbalanced data, a common occurrence in risk prediction.
- An implementation of the proposed framework that demonstrates the framework's applicability in predicting delays in deliveries within a real-world multi-tier aerospace manufacturing supply chain.
- Quantification of the trade-off between interpretability and prediction performance through experiments for the particular case study.

The remainder of this article is organised as follows. Section 2 offers a concise summary of research efforts that apply data-driven AI techniques for SCRM-related purposes. Section 3 introduces our data-driven risk prediction framework and analyses the individual phases within from both AI and SCRM perspectives. Section 4 illustrates the applicability for SCRM within a real-world supply chain. Then, Section 5 presents results of using the implemented framework to predict delays in deliveries within the studied supply chain. These results are discussed in Section 6, with an emphasis on the trade-off between performance and interpretability, as well as the importance of feature-rich datasets and the effect of imbalanced datasets. Finally, Section 7 concludes and proposes directions for future research at the confluence of AI and SCRM.

2. Related work

The potential of applying big data analytics and machine learning techniques for SCRM has recently been considered in literature. Fan et al. [6] investigate potential big data sources related to supply chains and then propose an SCRM framework that relies on the availability of such data. The framework relies on analysing and monitoring supply chain data to detect emerging risks, maintain relevant risk reports and use these to initiate suitable actions such as replanning the supply chain. He et al. [7] similarly acknowledge the predictive capabilities of big data analytics and incorporate such a component within a generic SCRM process model. However, both of these works are purely theoretical in nature, without implementing or applying the proposed frameworks and models to a real-world case study.

To the best of our knowledge, only three articles in literature have applied big data analytics and machine learning techniques for the purposes of risk identification through detection or prediction within an SCRM process and, hence, are directly relevant to the purposes of this work. These are analysed in Section 2.1. There is also a number of additional articles that are less relevant, in that they investigate similar techniques but for different facets of SCRM, specifically assessing and responding to risks. These are the focus of Section 2.2.

2.1. Risk identification

Zage et al. [8] address supply chain security risks by proposing methods for identifying deceptive practices within the supply chain, specifically for the e-commerce domain. The approach relies on spectral analysis to analyse large amounts of online transactions to determine traces of the so-called fraudster-accomplice strategy: fraudulent vendors interacting exclusively with real or fake users with good reputation to indirectly and artificially build their a good reputation for themselves. This process leads to developing graph-based metrics which are used by semi-supervised clustering algorithms to accurately determine whether a vendor behaves similarly to a fraudster, based on minimal labelled data.

Ye et al. [9] investigate the applicability of machine learning techniques to identify supply chain disruptions that are rooted in the economic performance of firms within the chain. Publically available financial data, such as asset–liability ratio, are obtained for Chinese firms and for periods before, during and after some form of supply chain disruption took place. Part of these data are used to determine classes of similar firms in terms of financial performance. The rest are used as features in a multi-class SVM classifier, to determine links between financial performance and disruption. The resulting system is capable of determining whether a particular firm exhibits a similar financial profile to ones that have contributed to disruptions in the past.

Both of the aforementioned works [8,9] present ad-hoc solutions specifically aimed at the particular case studies they focus on. Also, their choice of algorithms and metrics does not take into account interpretability or imbalanced datasets. In contrast, our work proposes a generic framework for predicting risks using data-driven AI, which can be applicable to any SCRM-related effort and which places great emphasis on making the right choice of which algorithm and metric to use, based on the supply chain's goals and priorities and the characteristics of available data.

The case of increasing supply chain sustainability through risk mitigation is explored in Mani et al. [13]. In particular, they use data collected through an Internet-of-Things software platform for fleet management and vehicle tracking. Data involves distances travelled, fleet utilisation, vehicle speeds and stoppages and geo-fencing reports. The authors propose several risk-related usages of such data, such as: (1) identifying underutilised vehicles and optimise fleet to reduce carbon footprint; (2) identifying cases of vehicle seizure by law enforcement through geo-fencing data; and (3) using vehicle stoppage data to identify theft and unethical behaviour. However, the focus is on the use of statistical analysis methods, rather than providing some form of predictive capability through learning. Also, while the authors go into great detail about the relevant data, there is no discussion with regard to the implementation and evaluation of the analytics algorithms they employ.

2.2. Risk assessment and response

Machine learning and big data analytics have also been used in relation to risk assessment. The earliest attempt to incorporate machine learning techniques within SCRM is the use of Artificial Neural Networks (ANNs) by Bruzzone and Orsoni [14] to assess production losses. The ANNs are trained in a supervised mode, supplying them with specific scenarios correlating production times, quantities and capacities with corresponding cost estimates. Based on these training data, the ANNs learn how to correlate production characteristics with actual gains or losses, in order to then be able to assess and calculate cost estimates for scenarios with an unknown outcome.

Bayesian networks have shown great potential in modelling for risk assessment, especially in the area of safety risk analysis [15,16]. Two groups of researchers have identified this potential in relation to a specific aspect of the supply chain risk assessment process, namely risk propagation. Garvey et al. [17] use Bayesian networks to model risk dependency graphs which have the ability to adapt when new knowledge is acquired, thus making sure that risk propagation is modelled accurately. Ojha et al. [18] perform a similar analysis of risk propagation using Bayesian networks, automatically learning the interconnections between several risk factors for different supply chain stakeholders and using this knowledge to determine probability of occurrence and cost for risks. Instead of a (parametric) Bayesian network, Shang et al. [19] introduce a nonparametric Bayes model to assess transport time risks in air cargo supply chains. The authors show how the model can be used as the foundation for several data-driven strategic decisions, such as ranking suppliers at the route level, or higher.

A number of studies employ big data analytics in relation to risk response. Papadopoulos et al. [20] employ a data-driven approach to determine factors enabling supply chain resilience, highlighting the importance of swift trust and quality information sharing. Li and Wang [21], on the other hand, use sensor-based data within a food supply chain to dynamically predict the product time-temperature profile; this enables firms to adjust their pricing schemes accordingly, in order to mitigate risk of spoilage before foods are sold. Chen et al. [22] recognise the effect of

incomplete or imperfect knowledge in managing disruption risks and propose a Bayesian learning model to increase robustness in the evaluation of inventory and sourcing strategies of a manufacturer in response to risks. Finally, Zhao and Yu [23] explore the potential of machine learning in the context of supplier selection. Their work relies on data from several past cases of selecting suppliers which are clustered based on similarity and are subsequently fed into a back-propagation ANN to extract rules for optimal supplier selection.

While not directly comparable to our work, the studies summarised in this subsection show the applicability and potential benefits of machine learning and data-driven approaches on a wide array of issues throughout the SCRM process. In general, even though such techniques represent only a small minority of SCRM research (e.g. compared to mathematical programming techniques, as discussed in [2]), their integration in standard SCRM processes can prove advantageous, as evidenced by the results within the studies presented in this section.

3. Data-driven risk prediction framework

To enhance supply chain risk prediction, we propose a framework that relies on the integration of artificial intelligence within the SCRM process. The framework aims for a two-way interactivity and synergy between supply chain and AI experts: any decision by the AI experts depends on specific input by the supply chain, while any models and results produced have to be interpretable so that they can influence SCRM decision-making. Fig. 1 illustrates the framework's process flow. On the right-hand side of the figure, the focus is on traditional tasks included in a standard SCRM process. The left-hand side includes the major tasks that are involved in a data-driven AI methodology. As should be obvious, the framework relies on effective synergies between a team involved in managing risks within a supply chain and a team specialising in data-driven AI. The remainder of this section is devoted to an in-depth analysis of the proposed framework.

3.1. Planning and data collection

The effectiveness of the proposed risk prediction framework relies on two fundamental prerequisite characteristics. First, a risk management plan needs to be created for the supply chain at hand. This plan should clarify the objectives and define the scope of risk management. Indicative questions that would need addressing in terms of objectives include why SCRM is necessary, what outcome is needed and when and which decisions are required and when. In terms of scope, relevant parties should decide whether the focus is the focal firm, risks internal or external to the SC and the specific types of these risks (e.g. environmental, supply, demand, process or control risks according to the classification in [24]). Other important planning decisions indicatively include whether certain types of risk are beyond boundaries and the amount of effort to be allocated in the SCRM process.

The outcome of SCRM planning leads to the second prerequisite of the proposed framework which involves the collection of relevant data. As analysed in [6], data sources in relation to SC are numerous and can be distinguished into internal and external ones. Internal data sources include purchasing, production, delivery and sales records, GPS and container sensor information, firm finances and human resources data. External sources are not directly related to the supply chain and can include newsitems, weather reports, social media activity, national and international policies and so on. Given the multitude of sources, making informed choices targeted at the risks identified in SCRM planning is paramount.

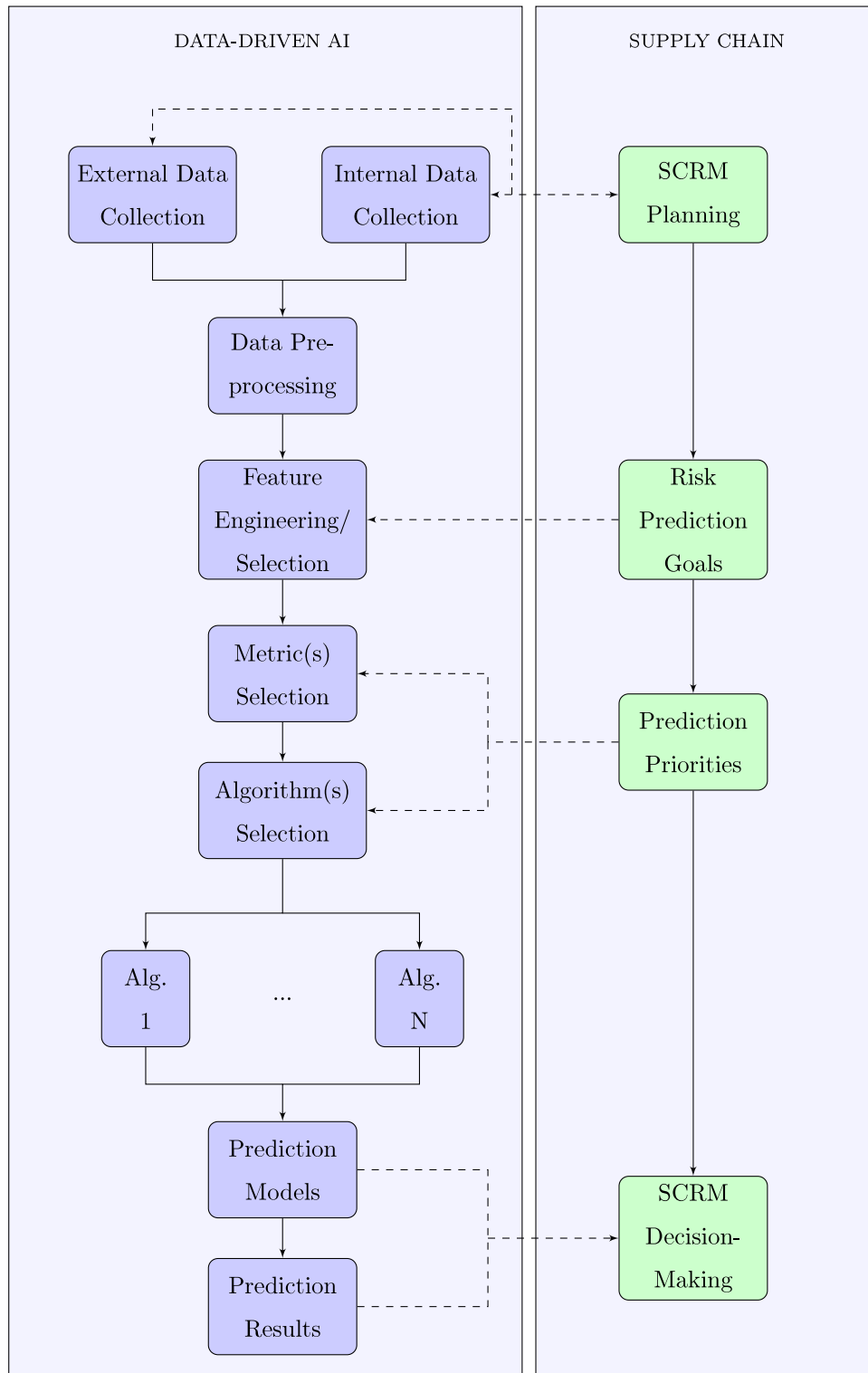


Fig. 1. Data-driven risk prediction framework.

After deciding on particular sources, historical data need to be collected, going as far back as necessary, taking into account the potential targets of the SCRM process and any financial or privacy limitations that may be involved. It may also be necessary to monitor both external and internal data sources, in order to continuously update available information and ensuring that any predictions coming from the framework correspond to the most recent data. This is especially important if the algorithms chosen at a later stage fall into the category of online machine learning,

as discussed later in Section 3.3. For example, if the SCRM process focuses on predicting supplier score trends based on previous performance, it is important to keep predictions up to date to ensure that no supplier is scored based only on their performance in the distant past.

Note that there may be cases where there is also a need for reverse process with regard to data collection and SCRM planning, due to difficulties in obtaining relevant data. Instead of deciding on a particular SCRM plan which then exclusively dictates which

data needs to be collected, data that is already available or easily obtainable is used to determine the scope of the SCRM plan. This is indicated in Fig. 1 by the use of arrows on both sides of the edges that connects planning with data collection.

3.2. Deciding on prediction goals and features

The outcome of the SCRM planning task naturally leads into the next step for the SC side of the framework, which is deciding on the specific goals of the risk prediction process. These may relate to the overall goals and priorities of the supply chain and may be influenced by recent events that may have instigated the need for SCRM in the first place. As with the overall plan, data availability may also affect the choice of risk prediction goals: a data-driven risk prediction framework is only able to predict risks for which relevant data is available. Hence, while SCRM planning may lead to identifying several prediction goals, only those for which relevant data collection (internal or external) has been (or can be) successful may actually be included as targets. The success of this phase (and the risk prediction process as a whole) largely depends on meaningful communication between SCRM and AI experts.

The choice of risk prediction goals directly affects the particular features that will be the focus of the data-driven AI algorithms. Before this step, however, there may be a need for some initial processing of the data collected earlier. Data preprocessing encompasses a wide range of procedures, such as ensuring that data is in a machine-readable format, cleaning data to remove inconsistent or incomplete entries, handling gaps and unknown values in data and scaling, transforming, normalising or discretising data. Note that some of these processes may be dependent on the particular algorithms to be applied, hence they may be delayed until algorithm selection has been performed, as described in Section 3.3.

The preprocessed data is then subjected to the feature extraction, engineering and selection processes [25, Chapter 6]. These are significantly important and may actually make or break a machine learning project [11]. Hence, they directly affect the success or failure of data-driven risk prediction. Feature extraction involves examining the raw attributes in the available datasets and extracting those features that are representative of the dataset and based on which prediction models can be built. Feature engineering involves adding to the initially extracted feature set by combining variables and/or features to create new features that may increase predictive power. Finally, feature selection refers to choosing a subset of the extracted and engineered features to reduce complexity of the learning process and to tackle notorious issues such as the curse of dimensionality [26] and overfitting [27]. The ultimate goal of these processes is to turn a simple set of values in a dataset into a meaningful set of features that has the potential to assist in the risk prediction goals defined previously.

3.3. Linking prediction priorities to metrics and algorithms

If the previous steps focused on making the best out of the available data, the next do the same in terms of available algorithms. Nowadays, there is a vast array of data analytics techniques and machine learning algorithms at our disposal and the question has undoubtedly shifted from using any of these techniques and algorithms to choosing the most suitable ones for the problem at hand. In the proposed SCRM framework, the choice of both algorithms and metrics is inextricably linked to the priorities of the risk prediction task. Below, we analyse some factors that affect this choice.

3.3.1. Interpretability

The first and foremost issue is whether the risk prediction process is aimed primarily at maximising prediction performance (e.g. in terms of accuracy) or understanding which features contribute to a risk becoming a reality. In the former case, machine learning algorithms that have a proven track record of performance in a wide variety of settings and data input should be chosen, regardless of whether they adopt a black box or a white box approach in terms of their models. However, black box approaches have to be excluded if the priority is to understand and explain why the model leads to one prediction instead of another. The case study presented in Section 4 explores this trade-off in more detail.

3.3.2. Data velocity and volume

Another issue affecting the choice of machine learning algorithms is related to the input data sources, which may either be historical, updated at a slow velocity, or real-time, changing as soon as new information is available. Online machine learning algorithms fit more with the latter case, while batch learning may be more suitable for historical data. The decision is again tied to prediction priorities: if there is a need for an active approach that monitors live data related to risks and issues prediction alerts, then online learning has to be used; this would not be the case, if, for instance, the goal is to assess past iterations of supply chain interactions to plan future steps.

Similarly, the size of available data also contributes to the decision of which machine learning algorithm to employ. Algorithms that can be parallelised can work more efficiently with large-scale datasets. Note that, at this point, large data size refers to the number of samples and not the number of dimensions. Reducing the number of dimensions is handled in previous steps, e.g. through feature selection, as discussed in Section 3.2.

3.3.3. Metrics

Equally important to the choice of algorithms is the choice of metrics. Especially in a risk prediction setting, the chosen metric must be directly linked to the desired outcome of the prediction process. For instance, if we are using a classification methodology to predict whether a risk will manifest, then the prediction model can lead to one of four outcomes, which may have different levels of importance, depending on prediction priorities. Specifically:

- True positive (TP): the model correctly predicts the risk manifesting. This allows the supply chain to prepare for such an outcome early enough to mitigate consequences and any resources spent for this are justified. Hence, true positives have to be maximised and the metrics chosen must be able to capture this.
- True negative (TN): the model correctly predicts that the risk will not manifest. These should also be maximised, but their importance may be lower than TPs.
- False Positive (FP): the model incorrectly predict that the risk will manifest, which means mitigation resources have been spent unnecessarily. If saving resources is prioritised higher than capturing all likely cases of the risk manifesting, then a metric that lends more importance to FPs than TPs.
- False Negative (FN): the model incorrectly predict that the risk will not manifest. This may be the most undesirable outcome, since, in essence, the risk prediction goal is not achieved. In some cases, the priority may be to minimise FNs, even at the expense of an increase in FPs.

Given these, the choice of a standard metric like accuracy ($\frac{TP+TN}{PP+PN}$, with $PP = TP+FP$ and $PN = TN+FN$) may not be suitable, since high values of accuracy may be achieved when TNs are high

but FNs are low, leading to a misleading view of the actual merit of the predictor. Also, since the algorithms' learning capability is iteratively evaluated using metrics, an unsuitable metric may also push learning towards an undesirable direction.

3.3.4. Imbalanced data

The choice of algorithms and metrics is all the more important when data is imbalanced, which is a common occurrence in risk prediction settings: in historical data, cases where risks occur are less than cases where risks did not manifest; in a binary classification setting, the former represent the so-called minor class, with the latter occupying the major class. He and Garcia [28] summarise methodologies for handling imbalanced data in three main categories: sampling methods to rebalance the dataset, cost-sensitive learning algorithms that penalise misclassification more for minor classes and kernel-based or active learning methods. They also argue against the use of accuracy metrics in an imbalanced setting, due to their potential to be deceiving and proposed various alternatives such as ROC or precision–recall curves.

3.4. Making decisions based on prediction models and results

The algorithms and metrics chosen in the previous step yield one or more trained prediction models which can then be fed with available data to produce results. These are then used to influence SCRM-related decision making. Indicatively, decisions that can be influenced may revolve around the following broad risk categories:

- Supplier risk: supplier selection decisions based on score trends (e.g. how often a supplier delivers on time)
- Demand risk: determining customer volatility in terms of order quantities or dates and assisting in peak-and-trough analysis
- Capacity risk: addressing volatile needs due to seasonal trends or because of conflicting customer order books
- Process/product risk: assessing product complexity (e.g. likelihood of right first time) and creating product profiles (e.g. runners, repeaters or strangers)

4. Case study

To illustrate the applicability of the proposed framework, we explore the case of SCRM within a real-world multi-tier aerospace manufacturing supply chain, with partners in Europe and Asia. In this section, we explain how we employed the proposed framework, presenting a detailed account of each step. For most data-driven AI tasks, we relied on scikit-learn v. 0.20.2 [29]. For resampling, we used imbalanced-learn v 0.4.2 [30]. All experiments were performed on a Windows® 10 system with an Intel® Core™ i7-4770 CPU at 3.40 GHz and 16 GB RAM. The source code of our implementation is available at <https://github.com/gmparg/FGCS-SCRM>.

4.1. Dataset and SCRM planning

Through initial discussions, a general understanding of potential risk prediction use cases was gained. However, due to the limited availability of easily accessible data and the inability to conduct further data collection processes, we settled on a dataset containing information on around 500,000 product deliveries from tier 2 suppliers to the tier 1 supplier in the supply chain for a six-year period (2011–2016). For each delivery, the following data are available:

- Tier 1 supplier: site name and id
- Tier 2 suppliers: supplier name and id
- Products: part number and description; unit price
- Orders: purchase order number, date and line number; quantity ordered; due date; original requested delivery date; currently accepted delivery date
- Deliveries: receipt date; quantity received; quantity rejected; purchase order line delivery status

While the overall SCRM activities of the particular supply chain expand across a wide spectrum of potential risks, for this particular case study the focus was on risks related to suppliers. Particularly, the plan communicated by partners involved managing risks related to suppliers being unable to fulfil future orders or delivering a product late and using such information to determine supplier score trends based on the number of successful (early or on-time) deliveries per month. Based on this plan, we opted to explore the case of predicting whether a supplier-related risk will occur. To achieve this, we employed binary classification algorithms to predict whether future deliveries of a particular supplier will be late or not.

The dataset was provided in CSV format, which can be fed directly as input to scikit-learn implementations. A number of preprocessing tasks were necessary. First, we removed any incomplete entries, e.g. deliveries where some of the relevant dates or statuses were not included or were invalid. Also, we removed redundant variables, specifically the full name of a supplier and the full description of a part, since alphanumeric identifiers for both suppliers and parts are also available. Then, all available data had to be converted to numerical form. In some cases, such as quantities and unit prices, no conversion was necessary. However, dates had to be split into three numbers (year, month and day), while alphanumeric values such as supplier identifiers or part numbers were converted to unique numerical identifiers. Finally, the resulting numerical data were normalised, scaling individual samples (i.e. deliveries) so that they have unit norm. We used the least squares norm (also known as L2-norm).

4.2. Feature engineering and selection

The next step in the process was to create a feature set with the maximum prediction potential. To do this we first had to decide on a particular prediction goal, in relation with the SCRM plan of managing supplier-related risks. The goal chosen was to predict, for a particular supplier, whether a delivery will be late or not. Based on this goal, we first manually assessed the variables within the dataset to determine relevant features. This resulted in excluding only tier 1 supplier site and id, since for the particular tier 2 supplier selected, only one tier 1 supplier site is involved. We also removed any variables that could lead to data leakage: receipt date, quantity received, quantity rejected and purchase order line delivery status; this information would not be available at the time of prediction. This process resulted into 15 features: unique part id, quantity, unit price, purchase order year, month and day, due year, month and day, original request year, month and day and currently accepted delivery year, month and day.

As a target variable, we selected the delivery status variable, which takes one of the following three values: early (8 or more business days early), on time (up to 7 business days early or up to 3 business days late) and late (4 or more business dates late). For simplification, we adapted this to two values: 1 to denote deliveries that are 4 or more business days late and 0 for all others. In this way, the goal of predicting whether a delivery is late or not amounts to a binary classification problem. Note that, as expected, our dataset is imbalanced, since 23% of the deliveries contained are late, while the rest are on time or early; in the case

of some suppliers the imbalance is even more pronounced, with only 10% of the deliveries being late.

Initial experiments using the initial feature set led to prediction results that were only marginally better than those achieved with the Zero Rule classifier. This classifier is a more suitable baseline for imbalanced datasets, since it always chooses the majority class, which means that in our case it classifies all deliveries as not late. To improve results, we decided to apply feature engineering on some of these features to create additional ones. Specifically, for each date in the original feature set, we also added features referring to the week day (from 1 to 7, corresponding to Sunday through Saturday), week number (from 1 to 52) and season (from 1 to 4, corresponding to winter, spring, summer and autumn). We also created features by taking the difference between each distinct pair of dates (with the 4 dates – purchase order, due, request, currently accepted delivery – yielding 6 distinct pairs in total). As a result of this process, the feature set size increased to 33 features.

Following these manual processes, we run automated feature selection to rank features based on their importance and exclude the lowest-ranked ones. We used a number of different approaches implemented in scikit-learn, specifically: removing features with low variance, univariate feature selection by ranking features based on the ANOVA F-value, mutual information and χ^2 test, recursive feature elimination and feature selection using the Extra-Trees algorithm. In general, all approaches ranked features similarly. We deferred any decision on cutoff thresholds until after selecting and running learning algorithms, so that we could see how leaving out some of the features affects prediction performance (see Section 5).

4.3. Metrics and algorithms selection

The final important decision to make was to determine how we would evaluate prediction results and which learning algorithms would deliver these results. Since no specific prediction priorities were communicated to us at this point, we could not make any safe assumptions regarding prioritisation of positive and negative predictions. The choice of metrics which prioritise one over the other, such as precision ($\frac{TP}{TP+FP}$) or recall ($\frac{TP}{TP+FN}$) might introduce unjustified bias, so we opted to use metrics that balance between different outcomes and ones that are suitable for imbalanced datasets like the one in our case study. Specifically, we consider the following:

- F_1 score, which is equal to $2 * \frac{\text{precision} * \text{recall}}{\text{precision} + \text{recall}}$, essentially the harmonic average between precision and recall, attributing equal importance to them.
- Average precision $AP = \sum n(R_n - R_{n-1}) * P_n$, with R_n and P_n denoting precision and recall at the n th threshold, respectively.
- Matthews correlation coefficient, which is defined by the following equation: $MCC = \frac{TP * TN - FP * FN}{\sqrt{(TP+FP)(TP+FN)(TN+FP)(TN+FN)}}$. This is another balanced measure that is especially suitable for imbalanced datasets [31].
- Confusion matrix, in order to obtain a direct view of how numbers shift between true/false positives and negatives.

In what concerns the choice of algorithms, as argued in Section 1 we have to consider the trade-off between performance and interpretability. While the requirement that algorithms must make correct predictions in as many cases as possible is undeniable, we also need these results to be explainable so that they can influence SCRM decision-making. To investigate this trade-off, we selected two algorithms. We first chose support vector machines (SVM) with an RBF kernel, as an example of highly performant

learners [32]. Note that any other algorithm that is known to perform well in binary classification problems can be chosen, such as the now ubiquitous neural network-based learning algorithms. Our choice is indicative and is informed by our need for an off-the-shelf solution with as few tuning parameters as possible. On the interpretable side of the trade-off, we chose decision tree learning, since classification trees are generally easily understood by non-experts, either in their graphical form or converted to a set of IF-THEN rules [25].

4.4. Model generation

Before feeding the dataset to the selected algorithms, we split it into training and test sets, holding out 20% of the available data for the test set. Note that this was done separately for each supplier in our dataset. For the training process, we used stratified 5-fold cross validation to ensure that each class is correctly represented in all folds. This is especially important in our case, due to data imbalance. In the case of SVM, we used a grid search to determine optimal parameters C and γ (penalty parameter and kernel coefficient). For decision tree learning, we run two experiments, one with default parameters that allow an arbitrarily large tree and one where we limit the tree depth to 6 and the number of leaf nodes to 13. This is aimed at ensuring interpretability of the resulting models [12]: a depth of 6 translates to up to 6 decision points, while limiting leaf nodes ensures that the tree will not grow too large, since a tree with such depth can contain maximum $2^6 = 64$ nodes.

Since the dataset is imbalanced, we also explored whether techniques specifically designed to address data imbalance could improve the generated models. We tested over-sampling techniques (random, SMOTE [33] and ADASYN [34]), under-sampling techniques (random, cluster centroids and Tomek's links [35]), combination of over- and under-sampling (SMOTE+Tomek [36]) and penalised models, with weights assigned to classes in an inverse proportion to the number of samples they contain (low weight to major class). Most of these techniques had an insignificant effect to the generated models, with the exception of those optimising recall, as analysed in the next section.

5. Results

In this section we first present the results of the feature selection process, followed by the prediction results using the SVM and decision tree models produced for the case study. All results presented refer to a single supplier, the one with the largest number of entries in the dataset (36 677 in total). Out of these, 5058 (13.8%) are late deliveries, while the rest are either early or on time. The validation test comprises 29 341 entries (80%), with the remaining 7336 (20%) constituting the test set.

5.1. Feature selection

In general, applying feature selection techniques to the set of 33 features did not have a very significant effect to prediction results. Out of the explored techniques, recursive feature elimination seemed to lead to slightly higher scores for most metrics. We used the implementation provided in scikit-learn which also supports cross validation. To rank features, we used a standard linear SVM with $C = 1$ and average precision as metric.

Fig. 2 shows the results of the recursive feature elimination process. The plotted scores are the prediction scores (with highest score 1) resulting using cross validation. The highest score is achieved for the subset containing 26 features. The 7 features that are eliminated are the following: year the purchase order was raised (ranked 2nd); year of delivery accepted by tier 1

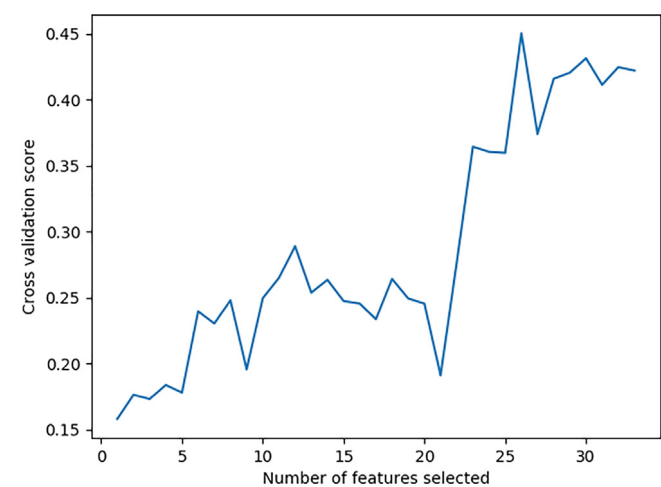


Fig. 2. Scores for each subset of 33 features.

supplier (ranked 3rd); order quantity (ranked 4th); part number (ranked 5th); day the purchase order was raised (ranked 6th); the difference between the delivery dates originally requested and accepted by tier 1 supplier (ranked 7th); and day the delivery was due (ranked 8th). For the remainder of this section we present results using both the original 33 features and the 26 remaining features after selection.

5.2. SVM models

As mentioned earlier, we conducted grid search using scikit-learn in order to find optimal values for the SVM parameters C and γ . C is the penalty of misclassification, with higher values meaning a higher penalty is imposed, hence the model is stricter. Values of γ express the influence that each particular training sample has on the model, with lower values meaning increased influence and higher values meaning less influence. Given the fact that we consider several different metrics, we run grid searches for each metric separately and for both feature sets (with 33 and 26 features).

Fig. 3 shows indicative results of the grid search process, using average precision and F_1 score as metrics. For average precision, very low C values are not optimal and results generally improve as values of γ increase. Highest average precision is achieved for $C = 1$ and $\gamma = 10^4$, which means that, in this particular case, precision is not enforced by imposing high penalties on misclassification, but by limiting the influence that each sample has on the model. For F_1 , results are similar, although differences are more pronounced. Scores are much lower for very low C values and while results improve as γ increases, they actually deteriorate for the highest value ($\gamma = 10^4$). Highest values are for $C = 10^4$ and $\gamma = 10^4$. Since F_1 is the harmonic mean between precision and recall and given the results on precision, we can draw the conclusion that higher misclassification penalty is required to achieve high recall while maintaining high precision. Results are similar between the two feature sets, with the smaller set achieving higher scores for a slightly higher range of parameters, which confirms the results of the recursive feature elimination process.

As should be expected, different optimal parameters were obtained through the grid search process for each metric and for the two feature sets. Tables 1 and 2 offer a comparison of these results. Each line corresponds to the optimal parameters calculated using one of the metrics. Values in bold are the highest achieved for each metric.

Table 1
Prediction scores using SVM and 33 features.

| Params | | Test scores | | | | | | Classification | | | |
|--------|----------|--------------|--------------|--------------|--------------|--------------|--|----------------|------|-----|-----|
| C | γ | AP | F_1 | Recall | MCC | Acc | | TP | TN | FP | FN |
| 1 | 10^4 | 0.835 | 0.764 | 0.712 | 0.646 | 0.939 | | 721 | 6169 | 155 | 291 |
| 10^3 | 10^3 | 0.632 | 0.771 | 0.740 | 0.738 | 0.940 | | 749 | 6144 | 180 | 263 |
| 10^4 | 10^3 | 0.618 | 0.765 | 0.766 | 0.728 | 0.935 | | 775 | 6086 | 238 | 237 |

Table 2
Prediction scores using SVM and 26 features.

| Params | | Test scores | | | | | | Classification | | | |
|--------|----------|--------------|--------------|--------------|--------------|--------------|--|----------------|------|-----|-----|
| C | γ | AP | F_1 | Recall | MCC | Acc | | TP | TN | FP | FN |
| 1 | 10^5 | 0.851 | 0.681 | 0.557 | 0.664 | 0.928 | | 564 | 6244 | 80 | 448 |
| 10^2 | 10^4 | 0.651 | 0.791 | 0.753 | 0.775 | 0.943 | | 774 | 6141 | 183 | 238 |
| 10^3 | 10^4 | 0.643 | 0.782 | 0.771 | 0.747 | 0.941 | | 780 | 6120 | 204 | 232 |

Table 3
Prediction scores using random oversampling and SVM.

| Params | | Test scores | | | | | | Classification | | | |
|-------------|----------|-------------|-------|--------|-------|-------|--|----------------|------|------|----|
| C | γ | AP | F_1 | Recall | MCC | Acc | | TP | TN | FP | FN |
| 33 features | | | | | | | | | | | |
| 0.01 | 10^4 | 0.263 | 0.419 | 0.973 | 0.376 | 0.627 | | 985 | 3617 | 2707 | 27 |
| 26 features | | | | | | | | | | | |
| 10 | 10^2 | 0.465 | 0.642 | 0.931 | 0.608 | 0.857 | | 942 | 5342 | 982 | 70 |

In general, SVM prediction models achieve good results across different metrics, although results are lower for balanced metrics and recall, than average precision. Also, highest values for average precision are achieved for lower penalty values (parameter C), while for all other metrics, higher penalty values are required. The results also show how misleading accuracy can be as a metric in scenarios with imbalanced datasets. If read without analysis, accuracy scores indicate that, regardless of parameter values, models are highly successful, achieving 94% accuracy. However, if we look at the TP and FN values, the reality is that the best-performing model misses $237/1012 = 23.4\%$ of late delivery cases. This is reflected more accurately in the scores using all other metrics.

Results improve slightly using the features that remain after the feature selection process. Highest MCC score is improved the most, by 5%, while highest recall is almost unchanged. The model with parameters $C = 10^3$ and $\gamma = 10^4$ achieves the lowest number of false positives, predicting correctly 77.1% of late deliveries. Parameters $C = 1$ and $\gamma = 10^5$ lead to the highest precision, with 87.6% of predictions of late delivery being correct. Finally, parameters $C = 10^2$ and $\gamma = 10^4$ lead to the best possible compromise between different classifications.

Using resampling techniques to tackle data imbalance does not lead to significant improvements in prediction performance, apart from recall. Table 3 shows the results of using random oversampling before running a grid search optimising recall. While almost perfect recall is achieved for both feature sets (predicting correctly 97.3% and 93.1% of late deliveries using 33 and 26 features, respectively, it is at the expense of the other metrics. Specifically, for the full feature set, 2707 samples are misclassified as late deliveries, leading to a precision of only 18.5%. For the 26-feature set, results are slightly more balanced, with 982 samples misclassified as late deliveries and a precision of 49%. This explains why random oversampling does not improve metrics such as F_1 or MCC, since they take into account both false positives and false negatives.

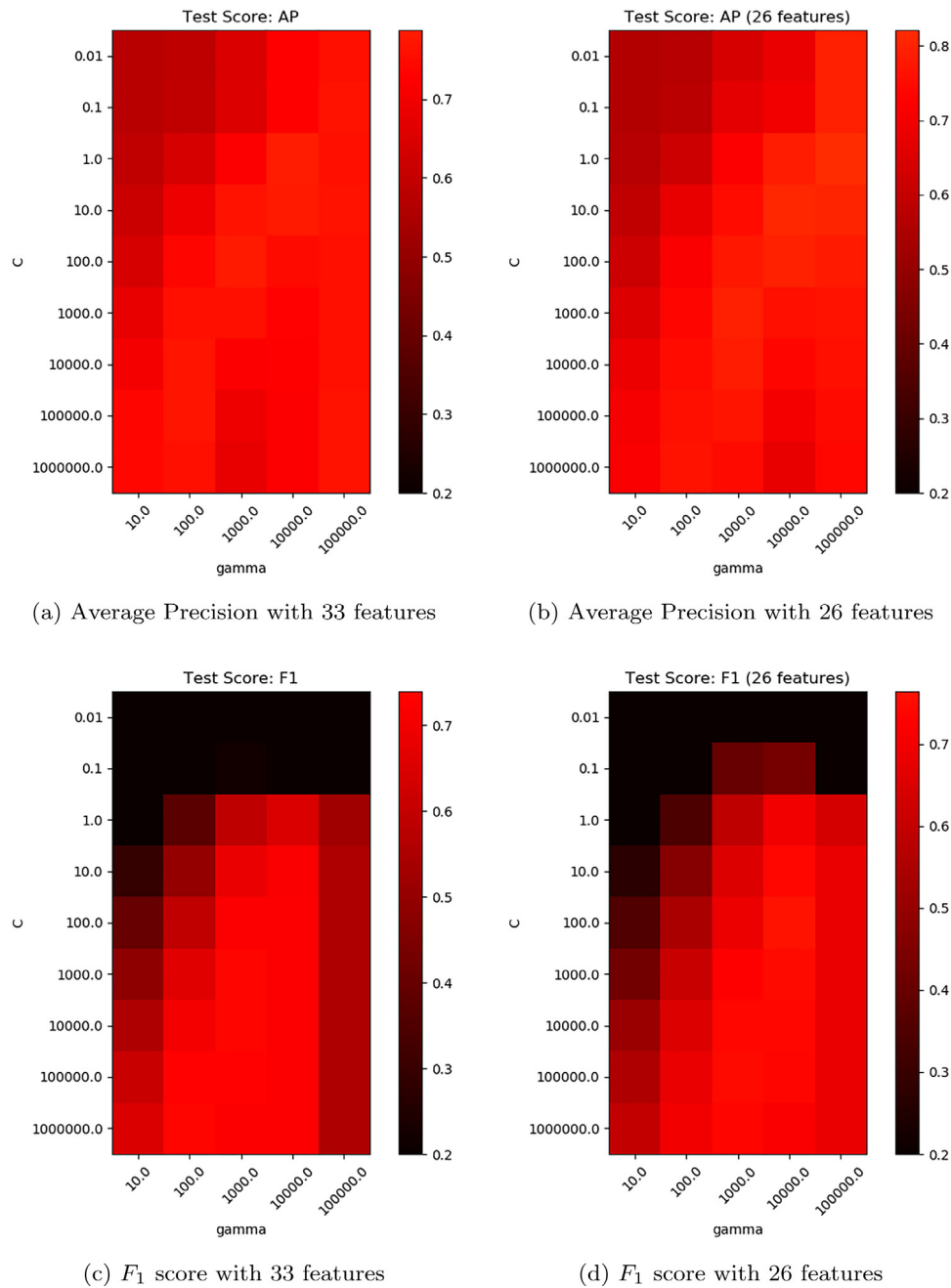


Fig. 3. Indicative grid search results for SVM parameters.

5.3. Decision tree models

For decision trees, we first executed the decision tree classifier implemented in scikit-learn with default parameters. This classifier is an optimised version of the CART algorithm uses the Gini impurity measure to decide on the quality of each split in the tree: each split must minimise the probability that a random element in the sets created by the split is misclassified if classified according to the label of the majority of the elements. Also, no limit is imposed on the size and structure of the resulting tree. Results using both feature sets are shown in Table 4.

Results are roughly comparable to those using SVM, slightly worse in terms of average precision but with slightly better F_1 scores. However, the resulting trees are considerably large: for 33 features the resulting tree has a maximum depth of 29 and 2267 total nodes, while for 26 features depth is 28 and total nodes

Table 4

Prediction scores using decision trees.

| Features | Test scores | | | | | Classification | | | |
|----------|-------------|-------|--------|-------|-------|----------------|------|-----|-----|
| | AP | F_1 | Recall | MCC | Acc | TP | TN | FP | FN |
| 33 | 0.666 | 0.800 | 0.817 | 0.768 | 0.944 | 827 | 6097 | 227 | 185 |
| 26 | 0.693 | 0.816 | 0.806 | 0.787 | 0.949 | 816 | 6153 | 171 | 196 |

are 2401. Since interpretability is hindered when trees grow too large, we limited the classifier to a maximum depth of 6 and maximum 13 leaf nodes. This led to trees with 25 nodes in total, such as the one in Fig. 4. Results using restricted parameters are shown in Table 5.

In the tree shown in Fig. 4 the feature that contributes to the best split ($X[24]$) is the difference between the due date of the delivery and the date that was accepted by the tier 1 supplier.

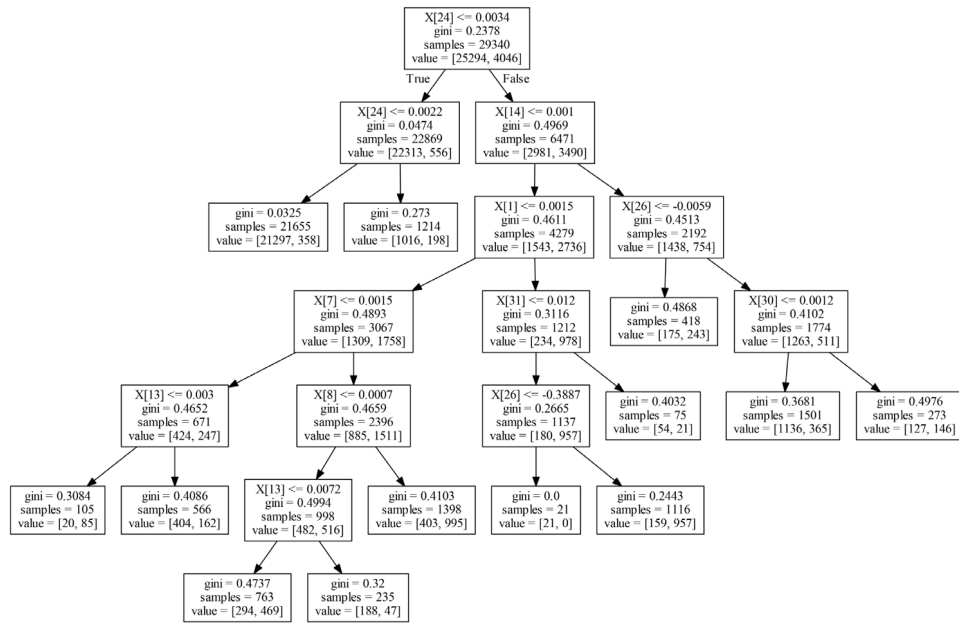


Fig. 4. Decision tree classifier using restricted parameters.

Table 5

Prediction scores using decision trees with max_depth=6 and max_leaf_nodes = 13.

| Features | Test scores | | | | | Classification | | | |
|----------|-------------|----------------|--------|-------|-------|----------------|------|-----|-----|
| | AP | F ₁ | Recall | MCC | Acc | TP | TN | FP | FN |
| 33 | 0.533 | 0.704 | 0.728 | 0.655 | 0.915 | 737 | 5980 | 344 | 275 |
| 26 | 0.516 | 0.683 | 0.651 | 0.636 | 0.916 | 659 | 6066 | 258 | 353 |

Table 6

Prediction scores using random oversampling and decision trees.

| Features | Test scores | | | | | Classification | | | |
|--|-------------|----------------|--------|-------|-------|----------------|------|------|-----|
| | AP | F ₁ | Recall | MCC | Acc | TP | TN | FP | FN |
| Unrestricted trees | | | | | | | | | |
| 33 | 0.698 | 0.821 | 0.823 | 0.792 | 0.950 | 833 | 6139 | 185 | 179 |
| 26 | 0.674 | 0.805 | 0.810 | 0.773 | 0.946 | 820 | 6118 | 206 | 192 |
| Trees with max_depth=6 and max_leaf_nodes=13 | | | | | | | | | |
| 33 | 0.493 | 0.670 | 0.883 | 0.630 | 0.880 | 894 | 5562 | 762 | 118 |
| 26 | 0.453 | 0.630 | 0.928 | 0.595 | 0.850 | 939 | 5295 | 1029 | 73 |

This leads to roughly 78% of the training samples classified as not late if the difference is less than or equal to 21 days (the value corresponding to 0.0022 before normalisation), which is incorrect for only 556 samples. For the remaining 23% the decision depends on combinations of additional features. For instance, when the season of the due date is not winter (corresponding to taking the left branch at node where $X[14] \leq 0.001$), the purchase order month is January to June (left branch of $X[1] \leq 0.0015$), the date that was accepted by the tier 1 supplier is not in the last 9 weeks of the year (left branch of $X[31] \leq 0.012$) and the delivery date originally requested by the tier 1 supplier is within a year of the currently accepted one (right branch of $X[26] \leq -0.3887$), then the delivery is predicted to be late. This is correct for 957 of the 1116 samples.

Using resampling techniques in combination with decision trees yields a mixture of results. For unrestricted decision trees, prediction performance is slightly improved overall for 33 features but is slightly lower for 26 features. In both cases, trees are significantly larger than without random oversampling: depth of 36 and 2711 nodes for 33 features and depth of 37 and 2681

Table 7

Summary of best prediction scores for different classifiers and metrics.

| Classifier | AP | F ₁ | Recall | MCC | Acc |
|------------|-------|----------------|--------|-------|-------|
| SVM | 0.851 | 0.791 | 0.973 | 0.775 | 0.943 |
| DT | 0.698 | 0.821 | 0.823 | 0.792 | 0.950 |
| RDT | 0.533 | 0.704 | 0.928 | 0.655 | 0.916 |

nodes for 26 features. For decision trees restricted to a maximum depth of 6 and maximum 13 leaf nodes, results are similar to SVM: recall is significantly improved for both 33 and 26 features, at the expense of the other metrics. Results are summarised in Table 6.

6. Discussion

In this section we discuss the presented results in relation to interpretability and dataset characteristics. Section 6.1 discusses the tradeoff between prediction performance and interpretability, as well as the potential knowledge that can be gained from interpretable models to influence SCRM processes. Then, Section 6.2 summarises the effects of feature engineering and selection, as well as data imbalance, as perceived in the presented case study. To facilitate discussion, Table 7 contains only the best scores, from the ones reported in the tables of Section 5, that were achieved for each metric using SVM, unrestricted decision trees (DT) and restricted decision trees (RDT).

6.1. Interpretability

Results in Table 7 show that decision trees without restrictions are capable of achieving comparable results with SVM, performing slightly worse with regard to average precision and recall but doing slightly better in the case of balanced metrics, F₁ score and MCC. Imposing restrictions on decision trees helps quantify the trade-off between performance and interpretability. For the particular dataset, a classifier whose results can be easily interpreted achieves 37% lower average precision, 15% lower MCC score, 11% lower F₁ score and 5% lower recall.

The question that the SCRM decision-makers have to consider is whether the potentially lower performance of interpretable

models is acceptable, given the fact that predictions can be interpreted. The decision is case-specific and is entirely dependent to the particular set of data, SCRM plan and prediction priorities. If, for instance, the importance placed on precision is high because there are significant costs in treating a delivery as late when this will not be the case eventually, then the added value of interpretability may be less appealing. If, on the other hand, the primary goal of the SCRM plan is to understand what factors may contribute more to a delivery being late, then some reduction in prediction performance may be deemed acceptable.

With regard to knowledge that can be gained from interpretable machine learning models, the presented case study shows that decision trees can be informative, revealing correlations of feature values that lead to one or the other outcome. Decision trees by design can deliver such interpretation, whereas this is not possible with SVM models. The particular dataset and features could only yield mainly date-related information, but richer datasets could potentially be even more helpful. This knowledge about correlation patterns in the dataset can then be taken into account in SCRM decision-making efforts, keeping in mind that correlation does not necessarily imply causality. The importance of tree depth should also be obvious by this example: explaining decisions starts becoming too complex after 4 or 5 splits.

6.2. Datasets and imbalance

The importance of a feature-rich dataset is also illustrated in the presented case study. The feature engineering process yielded additional features based on date-related information; however, further feature combinations were not possible, since the initial feature set was quite limited. Also, feature selection in such a dataset does not lead to significant improvement in performance; however, it can definitely prove useful in identifying the most important features and reducing the size of the feature set to increase execution time for complex tasks such as grid search for SVM.

Finally, the significant effect of imbalanced datasets is evident throughout the presented results, especially with regard to performance metrics. Accuracy is proven to be not only uninformative but potentially misleading. Balanced metrics such as F_1 and MCC scores present a more conservative but fair evaluation of prediction performance. However, no one metric is the best choice for any SCRM prediction task: it is the task and particularly the goals set within the SCRM planning process that should lead to the correct choice of metrics. As in the case study, recall is more useful when the goal is to minimise missed cases of late delivery, while precision is preferable when incorrectly predicting a delivery to be late is especially undesirable. These considerations should be taken into account in any SCRM-related prediction effort that involves events and risks that are less likely to occur.

7. Conclusions and future work

In this work, we proposed a risk prediction framework for SCRM that utilises data-driven AI techniques and relies on the collaboration and interactivity between AI and supply chain experts. The framework emphasises the need for linking choices of metrics and algorithms to SCRM goals which may prioritise interpretability over prediction performance or vice-versa. It also illustrates the difficulties of working with imbalanced datasets, which may feature in SCRM-related scenarios. The applicability of the framework is demonstrated through a real-world case study of a multi-tier aerospace manufacturing supply chain affected by the risk of delayed deliveries. Results of experiments conducted within the case study show that the framework can achieve

good performance across a variety of metrics using both black box and interpretable machine learning techniques. Prioritising interpretability over performance requires a compromise that is minor in terms of recall (5% decrease in prediction performance) but much higher in terms of average precision (37% decrease).

Future research directions on the application of AI techniques in SCRM include: (a) exploring a more feature-rich dataset and a larger set of machine learning techniques, including, for instance, neural networks and deep learning, and their effects on interpretability and performance; (b) extracting knowledge through a combination of data-driven and knowledge-based AI techniques [37,38] and using it to derive managerial insights and influence supply chain decision-making processes; and (c) investigating whether similar approaches can be applied in other phases of the SCRM process, such as risk assessment and response.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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