1	A comparison of nine machine learning mutagenicity models
2	and their application for predicting pyrrolizidine alkaloids
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14	Random forest, support vector machine, logistic regression, neural net-

works and k-nearest neighbor (lazar) algorithms, were applied to a new 15 Salmonella mutagenicity dataset with 8290 unique chemical structures utiliz-16 ing MolPrint2D and Chemistry Development Kit (CDK) descriptors. Cross-17 validation accuracies of all investigated models ranged from 80-85% which 18 is comparable with the interlaboratory variability of the Salmonella muta-19 genicity assay. Pyrrolizidine alkaloid predictions showed a clear distinction 20 between chemical groups, where otonecines had the highest proportion of 21 positive mutagenicity predictions and monoesters the lowest. 22

23 Introduction

The assessment of mutagenicity is an important part in the safety assessment of chemical structures, because mutations may lead to cancer and germ cells damage. The bacterial reverse mutation test (Ames test) is capable to identify substances that cause mutations (e.g., base-pair substitutions, frameshifts, insertions, deletions) and is generally used as the first step in genotoxicity and carcinogenicity assessments.

Computer based (*in silico*) mutagenicity predictions can be used in the early screening of novel compounds (e.g. drug candidates), but they are also gaining regulatory acceptance e.g. for the registration of industrial chemicals within REACH (European Chemicals Agency (ECHA) (2017)) or the assessment of impurities in pharmaceuticals (ICH M7 guideline, Harmonisation of Technical Requirements for Pharmaceuticals for Human Use International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use (ICH) (2017)).

³⁶ Currently, mutagenicity is the toxicological endpoint with the largest amount of public ³⁷ data for almost 10000 structures, whereas datasets for other endpoints contain typically ³⁸ only a few hundred compounds. The Ames test itself is relatively reproducible with an ³⁹ interlaboratory variability of 80-85% (Piegorsch and Zeiger (1991)).

This makes the development of mutagenicity models also interesting from a computational chemistry and machine learning point of view. The relatively large amount of public data reduces the probability of chance effects due to small sample sizes and the reliability of the underlying assay reduces the risk of overfitting experimental errors.

44 Within this study we attempted

- to generate a new public mutagenicity training dataset focusing on Salmonella
 typhimurium, by combining the most comprehensive public datasets
- 47

• to compare the performance of MolPrint2D (MP2D) fingerprints with Chemistry

48 Development Kit (*CDK*) descriptors for mutagenicity predictions

to compare the performance of global QSAR models (random forests (*RF*), support
 vector machines (*SVM*), logistic regression (*LR*), neural nets (*NN*)) with local
 models (lazar)

To demonstrate the application of mutagenicity models to compounds with very limited experimental data and to show their strengths an weaknesses we decided to apply them to Pyrrolizidine alkaloids (PAs).

Pyrrolizidine alkaloids (PAs) are characteristic metabolites of some plant families, mainly: Asteraceae, Boraginaceae, Fabaceae and Orchidaceae (Hartmann and Witte (1995), Langel, Ober, and Pelser (2011)) and form a powerful defence mechanism against herbivores. PAs are heterocyclic ester alkaloids composed of a necine base (two fused five-membered rings joined by a single nitrogen atom) and a necic acid (one or two carboxylic ester arms), occurring principally in two forms, tertiary base PAs and PA N-oxides.

In mammals, PAs are mainly metabolized in the liver. There are three principal
metabolic pathways for 1,2-unsaturated PAs (Chen, Mei, and Fu (2010)):

• Detoxification by hydrolysis of the ester bond on positions C7 and C9 by nonspecific esterases to release necine base and necic acid.

N-oxidation of the necine base to form PA N-oxides, which can be either conjugated
 by phase II enzymes and then excreted or converted back into the corresponding
 parent PA (Wang et al. (2005)). This detoxification pathway is not possible for
 otonecine-type PAs, as they are N-methylated (see Figure 1).

Metabolic activation or toxification by oxidation (for retronecine-type PAs) or
 oxidative N-demethylation (for otonecine-type Pas) by cytochromes P450 isoforms
 CYP2B and 3A (Lin, Cui, and Hawes (1998), Ruan et al. (2014)).

The latter reactions result in the formation of dehydropyrrolizidine (DHP) that is highly reactive and causes damage by building adducts with protein, lipids and DNA (Chen, Mei, and Fu (2010)). On the other hand, open diesters and macrocyclic PAs have a reduced detoxification due to steric hinderance of the respective esterases (Ruan et al. (2014)).

Therefore, the mutagenic probability of PAs is highly dependent on the structure of necine base and necic acid (Hadi et al. (2021); Allemang et al. (2018), Louisse et al. (2019)). However, due to limited availability of pure substances, only a small number of PAs have been investigated experimentally in an Ames test. To overcome this bottleneck, the application of different machine learning models to predict mutagenic probabilities based on structures and properties could provide further insights into the mutagenicity mechanisms of PAs.

Materials and Methods

86 Data

87 Mutagenicity training data

An identical training dataset was used for all models. The training dataset was compiled
from the following sources:

- Kazius/Bursi Dataset (4337 compounds, Kazius, McGuire, and Bursi (2005)):
 http://cheminformatics.org/datasets/bursi/cas_4337.zip
- Hansen Dataset (6513 compounds, Hansen et al. (2009)): http://doc.ml.tu-berlin.
 de/toxbenchmark/Mutagenicity_N6512.csv
- EFSA Dataset (695 compounds EFSA (2016)): https://data.europa.eu/euodp/ data/storage/f/2017-0719T142131/GENOTOX%20data%20and%20dictionary.xls

Mutagenicity classifications from Kazius and Hansen datasets were used without further 96 processing. According to these publications, compounds were classified as mutagenic if 97 at least one positive result has been obtained in Salmonella typhimurium strains TA97, 98 TA98, TA100, TA102, TA1535, TA1537 and TA1538 either with or without metabolic 99 activation by S9. E. coli results were not considered in these databases. To achieve 100 consistency with these datasets, EFSA compounds were classified as mutagenic, if at 101 least one positive result was found for the same Salmonella strains either with or without 102 metabolic activation and as non-mutagenic if no positive result was found. The complete 103 dataset contains chemicals from very diverse application areas (e.g. pharmaceuticals, 104 pesticides, industrial chemicals, environmental contaminants). 105

Dataset merges were based on unique SMILES (*Simplified Molecular Input Line Entry Specification*, Weininger, Weininger, and Weininger (1989)) strings of the compound structures. Duplicated experimental data with the same outcome was merged into a single value, because it is likely that it originated from the same experiment. Contradictory results were kept as multiple measurements in the database. The combined training dataset contains 8290 unique structures and 8309 individual measurements. Contradictory results were found for 19 substances.

¹¹³ Source code for all data download, extraction and merge operations is pub-¹¹⁴ licly available from the git repository https://git.in-silico.ch/mutagenicity-paper ¹¹⁵ under a GPL3 License. The new combined dataset can be found at https: ¹¹⁶ //git.in-silico.ch/mutagenicity-paper/tree/mutagenicity/mutagenicity.csv.

¹¹⁷ Pyrrolizidine alkaloid (PA) dataset

The pyrrolizidine alkaloid dataset was created from five independent, necine base substructure searches in PubChem (https://pubchem.ncbi.nlm.nih.gov/) and compared to the PAs listed in EFSA (2011) and the book by Mattocks (1986), to ensure, that all



Figure 1: Structural features of pyrrolizidine alkaloids

- major PAs were included. PAs mentioned in these publications, which were not found
 in the downloaded substances were searched individually in PubChem and, if available,
 downloaded separately. Non-PA substances, duplicates, and isomers were removed from
 the files, but artificial PAs, even if unlikely to occur in nature, were kept. The resulting
 PA dataset comprised a total of 602 different PAs. Further details about the compilation
 of the PA dataset are described in Schöning et al. (2017).
- The PAs in the dataset were classified according to structural features. A total of 9 different structural features were assigned to the necine base, to modifications of the necine base and to the necic acid (Figure 1):
- ¹³⁰ For the necine base, the following structural features were chosen:
- Retronecine-type (1,2-unstaturated necine base, 392 compounds)
- Otonecine-type (1,2-unstaturated necine base, 46 compounds)
- Platynecine-type (1,2-saturated necine base, 140 compounds)

¹³⁴ For the modifications of the necine base, the following structural features were chosen:

- N-oxide-type (84 compounds)
- Dehydropyrrolizidine-type (DHP, pyrrolic ester, 23 compounds)
- Tertiary-type (PAs which were neither from the N-oxide- nor DHP-type, 495 compounds)
- ¹³⁹ For the necic acid, the following structural features were chosen:
- Monoester-type (154 compounds)
- Open-ring diester-type (163 compounds)
- Macrocyclic diester-type (255 compounds)

143 Descriptors

144 MolPrint2D (MP2D) fingerprints

¹⁴⁵ MolPrint2D fingerprints (O'Boyle et al. (2011)) use atom environments as molecular ¹⁴⁶ representation. They determine for each atom in a molecule, the atom types of its ¹⁴⁷ connected atoms to represent their chemical environment. This resembles basically the ¹⁴⁸ chemical concept of functional groups.

In contrast to predefined lists of fragments (e.g. FP3, FP4 or MACCs fingerprints) or descriptors (e.g CDK) they are generated dynamically from chemical structures. This has the advantage that they can capture unknown substructures of toxicological relevance that are not included in other descriptors. In addition, they allow the efficient calculation of chemical similarities (e.g. Tanimoto indices) with simple set operations.

¹⁵⁴ MolPrint2D fingerprints were calculated with the OpenBabel cheminformatics library ¹⁵⁵ (O'Boyle et al. (2011)) for the complete training dataset with 8290 unique structures. ¹⁵⁶ They can be obtained from the following locations:

157 Training data:

158	• sparse representation (https://git.in	-silico.ch/mutagenicity-paper/tree/mutagenicity/
159	mutagenicity-mp2d)	
160	• descriptor matrix (https://git.in-sil	ico.ch/mutagenicity-paper/tree/mutagenicity/
161	mutagenicity-mp2d.csv.gz)	
162	Pyrrolizidine alkaloids:	
163	• sparse representation (https://git.in	-silico.ch/mutagenicity-paper/tree/pyrrolizidine-alkaloids/
164	pa-mp2d)	
165	• descriptor matrix (https://git.in-sili	co.ch/mutagenicity-paper/tree/pyrrolizidine-alkaloids/

166 pa-mp2d.csv)

167 Chemistry Development Kit (CDK) descriptors

Molecular 1D and 2D descriptors were calculated with the PaDEL-Descriptors program (http://www.yapcwsoft.com version 2.21, Yap (2011)). PaDEL uses the Chemistry Development Kit (*CDK*, https://cdk.github.io/index.html) library for descriptor calculations.

As the training dataset contained 8309 instances, it was decided to delete all instances where CDK descriptor calculations failed during pre-processing. Furthermore, 19 substances with contradictory experimental results were removed. The final training dataset contained 1442 descriptors for 8083 compounds.

CDK training data can be obtained from https://git.in-silico.ch/mutagenicity-paper/
tree/mutagenicity/mutagenicity-cdk.csv.

¹⁷⁸ The same procedure was applied for the pyrrolizidine dataset yielding descriptors for

¹⁷⁹ compounds. CDK features for pyrrolizidine alkaloids are available at https://git.in-silico.

¹⁸⁰ ch/mutagenicity-paper/tree/pyrrolizidine-alkaloids/pa-cdk.csv.

181 Algorithms

182 lazar

lazar (*lazy structure activity relationships*) is a modular framework for read-across model
development and validation. It follows the following basic workflow: For a given chemical
structure lazar:

• searches in a database for similar structures (neighbours) with experimental data,

• builds a local QSAR model with these neighbours and

• uses this model to predict the unknown activity of the query compound.

This procedure resembles an automated version of read across predictions in toxicology.
In machine learning terms it would be classified as a k-nearest-neighbour algorithm.

Apart from this basic workflow, lazar is completely modular and allows the researcher to use arbitrary algorithms for similarity searches and local QSAR (*Quantitative structureactivity relationship*) modelling. Algorithms used within this study are described in the following sections.

195 Feature preprocessing

¹⁹⁶ MolPrint2D features were used without preprocessing. Near zero variance and strongly ¹⁹⁷ correlated CDK descriptors were removed and the remaining descriptor values were ¹⁹⁸ centered and scaled. Preprocessing was performed with the R caret preProcess function ¹⁹⁹ using the methods "nzv", "corr", "center" and "scale" with default settings.

200 Neighbour identification

Utilizing this modularity, similarity calculations were based both on MolPrint2D finger prints and on CDK descriptors.

For MolPrint2D fingerprints chemical similarity between two compounds a and b is expressed as the proportion between atom environments common in both structures $A \cap B$ and the total number of atom environments $A \cup B$ (Jaccard/Tanimoto index).

$$sim = \frac{|A \cap B|}{|A \cup B|}$$

For CDK descriptors chemical similarity between two compounds a and b is expressed as the cosine similarity between the descriptor vectors A for a and B for b.

$$sim = \frac{A \cdot B}{|A||B|}$$

Threshold selection is a trade-off between prediction accuracy (high threshold) and the number of predictable compounds (low threshold). As it is in many practical cases desirable to make predictions even in the absence of closely related neighbours, we follow a tiered approach:

- First a similarity threshold of 0.5 (MP2D/Tanimoto) or 0.9 (CDK/Cosine) is used
 to collect neighbours, to create a local QSAR model and to make a prediction for
 the query compound. This are predictions with *high confidence*.
- If any of these steps fails, the procedure is repeated with a similarity threshold of
 0.2 (MP2D/Tanimoto) or 0.7 (CDK/Cosine) and the prediction is flagged with a
 warning that it might be out of the applicability domain of the training data (low
 confidence).
- These similarity thresholds are the default values chosen by software developers and remained unchanged during the course of these experiments.
- Compounds with the same structure as the query structure are automatically eliminated
 from neighbours to obtain unbiased predictions in the presence of duplicates.

223 Local QSAR models and predictions

Only similar compounds (neighbours) above the threshold are used for local QSAR models. In this investigation, we are using a weighted majority vote from the neighbour's experimental data for mutagenicity classifications. Probabilities for both classes (mutagenic/non-mutagenic) are calculated according to the following formula and the class with the higher probability is used as prediction outcome.

$$p_c = \frac{\sum \sin_{n,c}}{\sum \sin_n}$$

 p_c Probability of class c (e.g. mutagenic or non-mutagenic)

230 $\sum \sin_{n,c}$ Sum of similarities of neighbours with class c

²³¹ $\sum sim_n$ Sum of all neighbours

232 Applicability domain

The applicability domain (AD) of lazar models is determined by the structural diver-233 sity of the training data. If no similar compounds are found in the training data no 234 predictions will be generated. Warnings are issued if the similarity threshold had to be 235 lowered from 0.5 to 0.2 in order to enable predictions. Predictions without warnings 236 can be considered as close to the applicability domain (high confidence) and predictions 237 with warnings as more distant from the applicability domain (low confidence). Quantita-238 tive applicability domain information can be obtained from the similarities of individual 239 neighbours. 240

241 Validation

²⁴² 10-fold cross validation was performed for model evaluation.

243 Pyrrolizidine alkaloid predictions

For the prediction of pyrrolizidine alkaloids models were generated with the MP2D and CDK training datasets. The complete feature set was used for MP2D predictions, for CDK predictions the intersection between training and pyrrolizidine alkaloid features was used.

248 Availability

- Source code for this manuscript (GPL3): https://git.in-silico.ch/lazar/tree/?h= mutagenicity-paper
- Crossvalidation experiments (GPL3): https://git.in-silico.ch/lazar/tree/models/
 ?h=mutagenicity-paper
- Pyrrolizidine alkaloid predictions (GPL3): https://git.in-silico.ch/lazar/tree/
 predictions/?h=mutagenicity-paper
- Public web interface: https://lazar.in-silico.ch

256 **Tensorflow models**

257 Feature Preprocessing

For preprocessing of the CDK features we used a quantile transformation to a uniform distribution. MP2D features were not preprocessed.

260 Random forests (RF)

For the random forest classifier we used the parameters n_estimators=1000 and max_leaf_nodes=200. For the other parameters we used the scikit-learn default values.

²⁶³ Logistic regression (SGD) (*LR-sgd*)

For the logistic regression we used a combination of five trained models. For each model we used a batch size of 64 and trained for 50 epochs. As an optimizer ADAM was chosen. For the other parameters we used the tensorflow default values.

²⁶⁷ Logistic regression (scikit) (*LR-scikit*)

²⁶⁸ For the logistic regression we used as parameters the scikit-learn default values.

269 Neural Nets (NN)

For the neural network we used a combination of five trained models. For each model we used a batch size of 64 and trained for 50 epochs. As an optimizer ADAM was chosen. The neural network had 4 hidden layers with 64 nodes each and a ReLu activation function. For the other parameters we used the tensorflow default values.

274 Support vector machines (SVM)

We used the SVM implemented in scikit-learn. We used the parameters kernel='rbf', gamma='scale'. For the other parameters we used the scikit-learn default values.

277 Validation

²⁷⁸ 10-fold cross-validation was used for all Tensorflow models.

279 Pyrrolizidine alkaloid predictions

For the prediction of pyrrolizidine alkaloids we trained the model described above on the training data. For training and prediction only the features were used that were in the intersection of features from the training data and the pyrrolizidine alkaloids.

283 Availability

	284	Jupyter	notebooks	for t	hese ex	periments	can b	e found	at	the	following	locatio
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285 Crossvalidation:

286	• MolPrint2D fingerprints: https://git.in-silico.ch/mutagenicity-paper/tree/
287	cross validations/tensorflow/prediction-v5-norm.ipynb
288	$\bullet \ {\rm CDK \ descriptors: \ https://git.in-silico.ch/mutagenicity-paper/tree/crossvalidations/}$
289	tensorflow/prediction-v5-ext.ipynb
290	Pyrrolizidine alkaloids:
291	• MolPrint2D fingerprints: https://git.in-silico.ch/mutagenicity-paper/tree/
292	pyrrolizidine-alkaloids/tensorflow/prediction-v5-ext-ext-Padel-2D.ipynb
293	CDK descriptors: https://git.in-silico.ch/mutagenicity-paper/tree/pyrrolizidine-alkaloids/
294	tensorflow/prediction-v5-ext-Padel-2D.ipynb

295 **Results**

²⁹⁶ **10-fold crossvalidations**

- ²⁹⁷ Crossvalidation results are summarized in the following tables: Table 1 shows results
- ²⁹⁸ with MolPrint2D descriptors and Table 2 with CDK descriptors.

Table 1: Summary of crossvalidation results with MolPrint2D descriptors (lazar-HC: lazar with high confidence, lazar-all: all lazar predictions, RF: random forests, LR-sgd: logistic regression (stochastic gradient descent), LR-scikit: logistic regression (scikit), NN: neural networks, SVM: support vector machines)

	lazar-HC	lazar-all	RF	LR-sgd	LR-scikit	NN	SVM
Accuracy	84	82	80	84	84	84	84
True positive rate	89	85	78	83	83	82	83

	lazar-HC	lazar-all	RF	LR-sgd	LR-scikit	NN	SVM
True negative rate	78	78	82	84	85	85	86
Positive predictive value	83	80	81	84	84	84	85
Negative predictive value	86	84	80	84	84	83	84
Nr. predictions	5864	7782	8303	8303	8303	8303	8303

Table 2: Summary of crossvalidation results with CDK descriptors (lazar-HC: lazar with high confidence, lazar-all: all lazar predictions, RF: random forests, LR-sgd: logistic regression (stochastic gradient descent), LR-scikit: logistic regression (scikit), NN: neural networks, SVM: support vector machines)

	lazar-HC	lazar-all	RF	LR-sgd	LR-scikit	NN	SVM
Accuracy	85	82	84	79	80	85	82
True positive rate	87	84	81	81	80	85	82
True negative rate	82	80	86	78	80	85	82
Positive predictive value	85	81	85	79	80	85	82
Negative predictive value	85	82	82	80	80	85	82
Nr. predictions	4872	7353	8077	8077	8077	8077	8077

Figure 2 depicts the position of all crossvalidation results in receiver operating characteristic (ROC) space.

- 301 Confusion matrices for all models are available from the git repository https://git.in-
- 302 silico.ch/mutagenicity-paper/tree/crossvalidations/confusion-matrices/, individual pre-

dictions can be found in https://git.in-silico.ch/mutagenicity-paper/tree/crossvalidations/predictions/.

- $_{304}$ All investigated algorithm/descriptor combinations give accuracies between (80 and 85%)
- ³⁰⁵ which is equivalent to the experimental variability of the Salmonella typhimurium mu-
- ³⁰⁶ tagenicity bioassay (80-85%, Piegorsch and Zeiger (1991)). Sensitivities and specificities



Figure 2: ROC plot of crossvalidation results (lazar-HC: lazar with high confidence, lazar-all: all lazar predictions, RF: random forests, LR-sgd: logistic regression (stochastic gradient descent), LR-scikit: logistic regression (scikit), NN: neural networks, SVM: support vector machines).

307 are balanced in all of these models.

³⁰⁸ Pyrrolizidine alkaloid mutagenicity predictions

Mutagenicity predictions of 602 pyrrolizidine alkaloids (PAs) from all investigated models can be downloaded from https://git.in-silico.ch/mutagenicity-paper/tree/ pyrrolizidine-alkaloids/pa-predictions.csv. A visual representation of all PA predictions can be found at https://git.in-silico.ch/mutagenicity-paper/tree/pyrrolizidine-alkaloids/ pa-predictions.pdf.

For the visualisation of the position of pyrrolizidine alkaloids in respect to the train-314 ing data set we have applied t-distributed stochastic neighbor embedding (t-SNE, 315 Maaten and Hinton (2008)) for MolPrint2D and CDK descriptors. t-SNE maps 316 each high-dimensional object (chemical) to a two-dimensional point, maintaining the 317 high-dimensional distances of the objects. Similar objects are represented by nearby 318 points and dissimilar objects are represented by distant points. t-SNE coordinates were 319 calculated with the R Rtsne package using the default settings (perplexity = 30, theta 320 $= 0.5, \max$ iter = 1000). 321

Figure 3 shows the t-SNE of pyrrolizidine alkaloids (PA) and the mutagenicity training data in MP2D space (Tanimoto/Jaccard similarity), which resembles basically the structural diversity of the investigated compounds.

Figure 4 shows the t-SNE of pyrrolizidine alkaloids (PA) and the mutagenicity training data in CDK space (Euclidean similarity), which resembles basically the physicalchemical properties of the investigated compounds.

Figure 5 and Figure 6 depict two example pyrrolizidine alkaloid mutagenicity predictions in the context of training data. t-SNE visualisations of all investigated models can be downloaded from https://git.in-silico.ch/mutagenicity-paper/figures.



Figure 3: t-SNE visualisation of mutagenicity training data and pyrrolizidine alkaloids (PA) in MP2D space



Figure 4: t-SNE visualisation of mutagenicity training data and pyrrolizidine alkaloids (PA) in CDK space



Figure 5: t-SNE visualisation of MP2D random forest predictions



Figure 6: t-SNE visualisation of all CDK lazar predictions

Table 3 summarises the outcome of pyrrolizidine alkaloid predictions from all models with MolPrint2D and CDK descriptors.

Table 5: Summary of pyrrolizidine alkaloid predictions								
Model	MP2D Mutagenic	Nr. predictions	CDK Mutagenic	Nr. predictions				
lazar-all	20% (111)	93%~(560)	39%~(193)	83%~(500)				
lazar-HC	25%~(76)	50%~(301)	45% (111)	41% (246)				
RF	5%~(28)	100%~(602)	2% (10)	100% (602)				
LR-sgd	21%~(127)	100%~(602)	$16\% \ (97)$	100% (602)				
LR-scikit	20%~(118)	100%~(602)	15%~(88)	100% (602)				
NN	21%~(124)	100%~(602)	25%~(150)	100% (602)				
SVM	14% (82)	100%~(602)	3%~(19)	$100\% \ (602)$				

Table 3: Summary of pyrrolizidine alkaloid predictions

Figure 7 displays the proportion of positive mutagenicity predictions from all models for the different pyrrolizidine alkaloid groups. Tensorflow models predicted all 602 pyrrolizidine alkaloids, lazar MP2D models predicted 560 compounds (301 with high confidence) and lazar CDK models 500 compounds (246 with high confidence).

For the lazar-HC model, only 50/41% of the PA dataset were within the stricter similarity thresholds of 0.5/0.9 (MP2D/CDK). Reduction of the similarity threshold to 0.2/0.5 in the lazar-all model increased the amount of predictable PAs to 93/83%. As the other ML models do not consider applicability domains, all PAs were predicted.

Although most of the models show similar accuracies, sensitivities and specificities in crossvalidation experiments some of the models (MPD-RF, CDK-RF and CDK-SVM) predict a lower number of mutagens (2-5%) than the majority of the models (14-25%, Table 3, Figure 7).

345 Over all models, the mean value of mutagenic predicted PAs was highest for otonecines



Figure 7: Summary of pyrrolizidine alkaloid predictions

(65%, 407/623), followed by macrocyclic diesters (31%, 1042/3356), dehydropyrrolizidines (27%, 74/268), tertiary PAs (19%, 1201/6307) and retronecines (15%, 762/5054).

When excluding the aforementioned three deviating models, the rank order stays the same, but the percentage of mutagenic PAs is higher.

The following rank order for mutagenic probability can be deduced from the results of all models taken together:

- 353 Necine base: Platynecine < Retronecine « Otonecine
- 354 Necic acid: Monoester < Diester « Macrocyclic diester
- 355 Modification of necine base: N-oxide < Tertiary PA < Dehydropyrrolizidine

356 Discussion

357 Data

A new training dataset for *Salmonella* mutagenicity was created from three different sources (Kazius, McGuire, and Bursi (2005), Hansen et al. (2009), EFSA (2016)). It contains 8290 unique chemical structures, which is according to our knowledge the largest public mutagenicity dataset presently available. The new training data can be downloaded from https://git.in-silico.ch/mutagenicity-paper/tree/mutagenicity/ mutagenicity.csv.

364 Algorithms

1azar is formally a k-nearest-neighbor algorithm that searches for similar structures for a given compound and calculates the prediction based on the experimental data for these structures. The QSAR literature calls such models frequently *local models*, because models are generated specifically for each query compound. The investigated tensorflow models are in contrast *global models*, i.e. a single model is used to make predictions for all compounds. It has been postulated in the past, that local models are more accurate, because they can account better for mechanisms that affect only a subset of the training data.

Table 1, Table 2 and Figure 2 show that the crossvalidation accuracies of all models are 373 comparable to the experimental variability of the Salmonella typhimurium mutagenicity 374 bioassay (80-85% according to Piegorsch and Zeiger (1991)). All of these models have 375 balanced sensitivity (true positive rate) and specificity (true negative rate) and provide 376 highly significant concordance with experimental data (as determined by McNemar's 377 Test). This is a clear indication that *in silico* predictions can be as reliable as the 378 bioassays. Given that the variability of experimental data is similar to model variability 379 it is impossible to decide which model gives the most accurate predictions, as models 380 with higher accuracies might just approximate experimental errors better than more 381 robust models. 382

Our results do not support the assumption that local models are superior to global models for classification purposes. For regression models (lowest observed effect level) we have found however that local models may outperform global models (Helma et al. (2018)) with accuracies similar to experimental variability.

As all investigated algorithms give similar accuracies the selection will depend more on practical considerations than on intrinsic properties. Nearest neighbor algorithms like lazar have the practical advantage that the rationales for individual predictions can be presented in a straightforward manner that is understandable without a background in statistics or machine learning (a screenshot of the mutagenicity prediction for 12,21-Dihydroxy-4-methyl-4,8-secosenecinonan-8,11,16-trione is depicted in Figure 8). This allows a critical examination of individual predictions and prevents blind trust in models ³⁹⁴ that are intransparent to users with a toxicological background.

395 **Descriptors**

³⁹⁶ This study uses two types of descriptors for the characterisation of chemical structures:

MolPrint2D fingerprints (MP2D, Bender et al. (2004)) use atom environments (i.e. connected atom types for all atoms in a molecule) as molecular representation, which resembles basically the chemical concept of functional groups. MP2D descriptors are used to determine chemical similarities in the default **lazar** settings, and previous experiments have shown, that they give more accurate results than predefined fingerprints (e.g. MACCS, FP2-4).

Chemistry Development Kit (CDK, Willighagen, Mayfield, and Alvarsson (2017)) descriptors
tors were calculated with the PaDEL graphical interface (Yap (2011)). They include 1D
and 2D topological descriptors as well as physical-chemical properties.

All investigated algorithms obtained models within the experimental variability for both
types of descriptors (Table 1, Table 2, Figure 2).

Given that similar predictive accuracies are obtainable from both types of descriptors
the choice depends once more on practical considerations:

MolPrint2D fragments can be calculated very efficiently for every well defined chemical structure with OpenBabel (O'Boyle et al. (2011)). CDK descriptor calculations are in contrast much more resource intensive and may fail for a significant number of compounds (from 8290).

⁴¹⁴ MolPrint2D fragments are generated dynamically from chemical structures and can be
⁴¹⁵ used to determine if a compound contains structural features that are absent in training
⁴¹⁶ data. This feature can be used to determine applicability domains. CDK descriptors
⁴¹⁷ contain in contrast a predefined set of descriptors with unknown toxicological relevance.



Figure 8: lazar screenshot of 12,21-Dihydroxy-4-methyl-4,8-secosenecinonan-8,11,16-trione mutagenicity prediction

MolPrint2D fingerprints can be represented very efficiently as sets of features that are present in a given compound which makes similarity calculations very efficient. Due to the large number of substructures present in training compounds, they lead however to large and sparsely populated datasets, if they have to be expanded to a binary matrix (e.g. as input for tensorflow models). CDK descriptors contain in contrast in every case matrices with 1442 columns which can cause substantial computational overhead.

424 Pyrrolizidine alkaloid mutagenicity predictions

425 Algorithms and descriptors

Figure 7 shows a clear differentiation between the different pyrrolizidine alkaloid groups.
Nevertheless differences between predictions from different algorithms and descriptors
(Table 3) were not expected based on crossvalidation results.

In order to investigate, if any of the investigated models show systematic errors in the vicinity of pyrrolizidine-alkaloids we have performed a detailled t-SNE analysis of all models (see Figure 5 and Figure 6 for two examples, all visualisations can be found at https://git.in-silico.ch/mutagenicity-paper/tree/figures).

⁴³³ None of the models showed obvious deviations from their expected behaviour, so the
⁴³⁴ reason for the disagreement between some of the models remains unclear at the moment.
⁴³⁵ It is however possible that some systematic errors are covered up by converting high
⁴³⁶ dimensional spaces to two coordinates and are thus invisible in t-SNE visualisations.

Only two compounds from the PA dataset (Senecivernine and Retronecine) are part of
the training set. Both are non-mutagenic and were predicted as non-mutagenic by all
models (instances have been removed from the training set for unbiased predictions).
Despite the exact concordance, we cannot draw any general conclusions about model
performance based on two examples with a single outcome.

442 Necic acid

The rank order of the necic acid is comparable in all models. PAs from the monoester 443 type had the lowest genotoxic probability, followed by PAs from the open-ring diester 444 type. PAs with macrocyclic diesters had the highest genotoxic probability. The result 445 fits well with current state of knowledge: in general, PAs, which have a macrocyclic 446 diesters as necic acid, are considered to be more mutagenic than those with an open-ring 447 diester or monoester (EFSA (2011), Fu et al. (2004)). As pointed out above, open 448 diesters and macrocyclic PAs have a reduced detoxification due to steric hinderance of 449 the respective esterases (Ruan et al. (2014)). This was also confirmed by more recent 450 studies, confirming that macrocyclic- and open-diesters are more genotoxic in vitro than 451 monoesters (Hadi et al. (2021); Allemang et al. (2018), Louisse et al. (2019)). 452

453 Necine base

In the rank order of necine base PAs, platynecine is the least mutagenic, followed by retronecine, and otonecine. Saturated PAs of the platynecine-type are generally accepted to be less or non-mutagenic and have been shown in *in vitro* experiments to form no DNA-adducts (Xia et al. (2013)). In literature, otonecine-type PAs were shown to be more mutagenic than those of the retronecine-type (Li et al. (2013)).

459 Modifications of necine base

The group-specific results reflect the expected relationship between the groups: the low mutagenic probability of *N*-oxides and the high probability of dehydropyrrolizidines (DHP) (Chen, Mei, and Fu (2010)). However, *N*-oxides may be *in vivo* converted back to their parent mutagenic/tumorigenic parent PA (Yan et al. (2008)), on the other hand they are highly water soluble and generally considered as detoxification products, which are *in vivo* quickly renally eliminated (Chen, Mei, and Fu (2010)).

DHP are regarded as the toxic principle in the metabolism of PAs, and are known to 466 produce protein- and DNA-adducts (Chen, Mei, and Fu (2010)). None of our investigated 467 models did meet this expectation and all of them predicted the majority of DHP as non-468 mutagenic. However, the following issues need to be considered. On the one hand, all 469 DHP were outside of the stricter applicability domain of MP2D lazar. This indicates 470 that they are structurally very different than the training data and might be out of the 471 applicability domain of all models based on this training set. In addition, DHP has two 472 unsaturated double bounds in its necine base, making it highly reactive. DHP and other 473 comparable molecules have a very short lifespan in vivo, and usually cannot be used in 474 in vitro experiments. 475

Overall the low number of positive mutagenicity predictions was unexpected. PAs are 476 generally considered to be genotoxic, and the mode of action is also known. Therefore, 477 the fact that some models predict the majority of PAs as not mutagenic seems contradic-478 tory. To understand this result, the experimental basis of the training dataset has to be 479 considered. The training dataset is based on the Salmonella typhimurium mutagenicity 480 bioassay (Ames test). There are some studies, which show mutagenicity of PAs in the 481 Ames test (Chen, Mei, and Fu (2010)). Also, Rubiolo et al. (1992) examined several 482 different PAs and several different extracts of PA-containing plants in the Ames test. 483 They found that the Ames test was indeed able to detect mutagenicity of PAs, but in 484 general, appeared to have a low sensitivity. The pre-incubation phase for metabolic 485 activation of PAs by microsomal enzymes was the sensitivity-limiting step. This could 486 very well mean that the low sensitivity of the Ames test for PAs is also reflected in the 487 investigated models. 488

In summary, we found marked differences in the predicted genotoxic probability between the PA groups: most mutagenic appeared the otonecines and macrocyclic diesters, least mutagenic the platynecines and the mono- and diesters. These results are comparable with *in vitro* measurements in hepatic HepaRG cells (Louisse et al. (2019)), where relative potencies (RP) were determined: for otonecines and cyclic diesters RP = 1, for open diesters RP = 0.1 and for monoesters RP = 0.01.

⁴⁹⁵ Due to a lack of differential data, European authorities based their risk assessment in ⁴⁹⁶ a worst-case approach on lasiocarpine, for which sufficient data on genotoxicity and ⁴⁹⁷ carcinogenicity were available (HMPC (2014), EMA (2020)). Our data further support ⁴⁹⁸ a tiered risk assessment based on *in silico* and experimental data on the relative potency ⁴⁹⁹ of individual PAs as already suggested by other authors (Merz and Schrenk (2016), Rutz ⁵⁰⁰ et al. (2020), Louisse et al. (2019)).

The practical question how to choose model predictions in the absence of experimental data remains open. Tensorflow predictions do not include applicability domain estimations and the rationales for predictions cannot be traced by toxicologists. Transparent models like **lazar** may have an advantage in this context, because they present rationales for predictions (similar compounds with experimental data) which can be accepted or rejected by toxicologists and provide validated applicability domain estimations.

507 Conclusions

A new public *Salmonella* mutagenicity training dataset with 8309 experimental results was created and used to train lazar and Tensorflow models with MolPrint2D and CDK descriptors. All investigated algorithm and descriptor combinations showed accuracies comparable to the interlaboratory variability of the Ames test.

⁵¹² Pyrrolizidine alkaloid predictions showed a clear separation between different classes of ⁵¹³ PAs which were generally in accordance with the current toxicological knowledge about ⁵¹⁴ these compounds. Some of the models showed however a substantially lower number of ⁵¹⁵ mutagenicity predictions, despite similar crossvalidation results and we were unable to ⁵¹⁶ identify the reasons for this discrepancy within this investigation.

⁵¹⁷ Our data show that large difference exist with regard to mutagenic probabilities between ⁵¹⁸ different pyrrolizidine subgroups. To adjust risk assessment of pyrrolizidine contami-⁵¹⁹ nation, our data supports a tiered risk assessment based on *in silico* predictions and ⁵²⁰ experimental data of individual pyrrolizidine alkaloids.

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