

# Trust, But Verify: On the Importance of Chemical Structure Curation in Cheminformatics and QSAR Modeling Research

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## 1. INTRODUCTION

With the recent advent of high-throughput technologies for both compound synthesis and biological screening, there is no shortage of publicly or commercially available data sets and databases<sup>1</sup> that can be used for computational drug discovery applications (reviewed recently in Williams et al.<sup>2</sup>). Rapid growth of large, publicly available databases (such as PubChem<sup>3</sup> or ChemSpider<sup>4</sup> containing more than 20 million molecular records each) enabled by experimental projects such as NIH's Molecular Libraries and Imaging Initiative<sup>5</sup> provides new opportunities for the development of cheminformatics methodologies and their application to knowledge discovery in molecular databases.

A fundamental assumption of any cheminformatics study is the correctness of the input data generated by experimental scientists and available in various data sets. Nevertheless, a recent study<sup>6</sup> showed that on average there are two errors per each medicinal chemistry publication with an overall error rate for compounds indexed in the WOMBAT database<sup>7</sup> as high as 8%. In another recent study,<sup>8</sup> the authors investigated several public and commercial databases to calculate their error rates: the latter were ranging from 0.1 to 3.4% depending on the database.

How significant is the problem of accurate structure representation (given that the error rates in current databases may appear relatively low) since it concerns exploratory cheminformatics and molecular modeling research? Recent investigations by a large group of collaborators from six laboratories<sup>9,10</sup> have clearly demonstrated that the type of chemical descriptors has much greater influence on the prediction performance of QSAR models than the nature of model optimization techniques. These findings suggest that having erroneous structures represented by erroneous descriptors should have a detrimental effect on model performance. Indeed, a recent seminal publication<sup>8</sup> clearly pointed out the importance of chemical data curation in the context of QSAR modeling. The authors have discussed the error rates in several known databases and evaluated the consequences of both random and systematic errors with respect

to the prediction performances of the derivative QSAR models. They also presented several illustrative examples of incorrect structures generated from either correct or incorrect SMILES. The main conclusions of the study were that small structural errors within a data set could lead to significant losses of predictive ability of QSAR models. The authors further demonstrated that manual curation of structural data leads to substantial increase in the model predictivity. This conclusion becomes especially important in light of the aforementioned study of Oprea et al.<sup>6</sup> that cited a significant error rate in medicinal chemistry literature.

Alarmed by these conclusions, we have examined several popular public databases of bioactive molecules to assess possible error rates in structure representation. For instance, the NCI AIDS Antiviral Screen<sup>11</sup> (the paper describing this screen<sup>12</sup> has been cited 57 times in PubMed Central only) comprises 42687 chemical records with their associated activity. Even quick analysis of this data sets revealed that 4202 (i.e., ca. 10%) compounds should be either treated with care or removed before any cheminformatics investigation: 3350 compounds were mixtures and salts, and we detected 741 pairs of exact duplicates and stereoisomers, possessing different or opposite reported activities. Similar observations can be made for several well-known public databases such as NCI Human Tumor Cell Line and PubChem, as well as for smaller data sets studied and reported in published literature. For instance, in a study already mentioned above, six research teams each specializing in QSAR modeling (including the authors of this paper!) collaborated on the analysis of the *Tetrahymena pyriformis* aquatic toxicity data set<sup>9,10</sup> comprising 1093 compounds. Later this exact data set was used by the organizers of CADASTER toxicity challenge.<sup>13</sup> However, our re-examination of this data set showed the presence of six pairs of duplicates among 1093 compounds because of the presence of different metal cations in salts (with different aquatic toxicities measured by pIGC<sub>50</sub> ranging from ~0.1 to 1 logarithmic unit. Furthermore, in the new external set compiled by the organizers of the CADASTER challenge to evaluate the comparative performance of competing models, eight out of 120 compounds were found to be structurally identical to modeling set compounds but with different toxicity values (~0.3 pIGC<sub>50</sub>).

Such disappointing observations may, at least in part, explain why QSAR models may sometimes fail, which is

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an issue that was brought up in several recent publications<sup>14–16</sup> (but none mentioned structure representation or biological annotation errors as possible sources of poor performance of QSAR models). Obviously, cheminformaticians must only use correct chemical structures and biological activities in their studies. Not surprisingly, any structural error translates into either inability to calculate descriptors for incorrect chemical records or erroneous descriptors. As a consequence, models developed using inaccurate data (either structural or biological) will have insignificant or reduced statistical power (cf., Young et al.<sup>8</sup>) and will be unreliable for prediction. Since the amount of data, the number of models, and the body of cheminformatics publications continue to grow, it becomes increasingly important to address the issue of data quality that inherently affects the quality of models.

Surprisingly, the investigations into how the primary data quality influences the performances of cheminformatics models are almost absent in the published literature. Besides the study by Young et al.<sup>8</sup> mentioned above we also found a paper by Southan et al.,<sup>17</sup> which mentioned briefly some procedures used to determine the number of unique chemical structures in a database. It appears that for the most part cheminformaticians and molecular modelers tend to take published chemical and biological data at their face value and launch calculations without carefully examining the accuracy of the data records. It is indeed difficult to verify the results of biological assays because it is well-known that numerical values of bioactivity for the same compounds measured in the same assays frequently disagree between different laboratories. However, there should be much less disagreement concerning the correct representation of a chemical structure for compounds in the databases except in certain difficult cases, such as chemicals with multiple tautomeric forms.<sup>18</sup> Very often errors in chemical representation are not obvious and are difficult to identify without special tools and protocols.

Both common sense and the recent investigations described above indicate that chemical record curation should be viewed as a separate and critical component of any cheminformatics research. By comparison, the community of protein X-ray crystallographers has long recognized the importance of structural data curation; indeed the Protein Data Bank (PDB) team includes a large group of curators whose major job is to process and validate primary data submitted to the PDB by crystallographers.<sup>19</sup> Furthermore, NIH recently awarded a significant Center grant to a group of scientists from the University of Michigan (<http://csardock.org/>) where one of the major tasks is to curate primary data on protein–ligand complexes deposited to the PDB. Conversely, to the best of our knowledge, even the largest publicly funded cheminformatics project, that is, PubChem, is considered as a data *repository*, that is, no special effort is dedicated to the curation of structural information deposited to PubChem by various contributors. Chemical data curation has been addressed whenever possible within the publicly available ChemSpider project;<sup>4</sup> however, until now, most effort has focused on data collection and database expansion. Thus, it is critical that scientists who build models using data derived from available databases or extracted from publications dedicate their *own* effort to the task of data curation.

Although there are obvious and compelling reasons to believe that chemical data curation should be given a lot of attention, it is also obvious that, for the most part, the basic steps to curate a data set of compounds have been either considered trivial or ignored by experts in the field. For instance, several years ago a group of experts in QSAR modeling developed what is now known as OECD QSAR modeling and validation principles;<sup>16,20</sup> these are a set of guidelines that the researchers should follow to achieve the regulatory acceptance of QSAR models. There are five stated principles that require QSAR models to be associated with (i) defined end point, (ii) unambiguous algorithm, (iii) defined domain of applicability, (iv) appropriate measures of goodness-of-fit, robustness and predictivity, and (v) if possible, mechanistic interpretation. The need to curate primary data used for model development is not even mentioned. Also, in an effort to improve the quality of publications in the QSAR modeling field, the *Journal of Chemical Information and Modeling* published a special editorial highlighting the requirements to QSAR papers that authors should follow to publish their results in the journal.<sup>21</sup> Again, no special attention was given to data curation. Finally, there have been several recent publications addressing common mistakes and criticizing various faulty practices in the QSAR modeling field;<sup>14–16,22–24</sup> however, none of these papers has explicitly discussed the importance of chemical record curation for developing robust QSAR models. There is an obvious trend within the community of QSAR modelers to establish and follow the standardized guidelines for developing statistically robust and externally predictive QSAR models.<sup>25</sup> It appears timely to emphasize the importance of and develop best practices for data preparation prior to initiating the modeling process because it is merely senseless to launch massive cheminformatics or molecular modeling investigations if the underlying chemical structures are not correct.

Arguably, each cheminformatics laboratory may have its own protocol to prepare and curate a compound data set before embarking on a modeling exercise. However, to the best of our knowledge, there is no published compilation of good practices for data set curation that at least beginners if not established researchers are advised to follow. In our opinion, there is a pressing need to amend the five OECD principles by adding a sixth rule that would request careful data curation prior to model development. Thus, this paper presents an attempt to address the issue of chemical data curation in a systematic way by pursuing the following major goals:

- (1) To alert the cheminformatics and molecular modeling community to the fact that a significant fraction of chemical and bioactivity data in the databases used for modeling may be erroneous, which is likely to reduce the quality of derived models.
- (2) To develop a set of data curation procedures integrated into a logical functional workflow that would process the input data and correct structural errors whenever possible (sometimes at the expense of removing incomplete or confusing data records).
- (3) To share organized protocols for data curation with the scientific community by providing sample case studies and explicit pointers to the sources where procedures discussed in this paper are available (with a bias toward data curation

software that is made available free of charge to academic investigators).

- (4) To illustrate, with at least a few examples, that rigorously developed QSAR models using well curated primary data may be employed not only for predicting new structures but also to spot and correct errors in biological data reported in databases used either for model development or model validation.

We wish to emphasize the importance of creating and following a standardized data curation strategy applicable to any ensemble of compounds, which should be really viewed as a community exercise. Although this paper targets mainly the beginners in the cheminformatics field, we believe that it may serve as a general reference for best practices for data set curation that could be useful for all scientists working in the area of cheminformatics, QSAR, and molecular modeling. We point out a general problem in the field that could be responsible for either reducing the quality of published models or preventing the generation of models worthy of publication. Importantly, we do not pretend to provide an ultimate, all-encompassing collection of curation practices covering all types of difficult or ambiguous cases but focus on most frequent and common cases. We hope that other experts will contribute their knowledge and best practices for dealing with both relatively simple as well as complex issues in subsequent publications.

Because of the complexity of modern chemical biology, there is a clear separation between scientists who generate data and those who analyze them. For the latter, data analytical studies are impossible without trusting the original data sources; however, it is important, whenever possible, to verify the accuracy of primary data before developing any model. To emphasize this point, the title of this paper in part repeats the famous proverb that was frequently cited by the late president Ronald Reagan during the cold war era and that traces back to the founder of the Russian KGB Felix Dzerzhinsky who supposedly invented it almost 100 years ago as a founding principle of his organization (cf., [http://en.wikipedia.org/wiki/Trust,\\_but\\_verify](http://en.wikipedia.org/wiki/Trust,_but_verify)).

## 2. MAIN STEPS FOR CHEMICAL DATA CURATION

In this section, we discuss the most important steps required to curate a chemical data set (Figure 1). We specifically focus on chemical structure curation procedures and do not cover the highly relevant but special topic of name to structure conversion, which is often used to create chemical databases (cf., section 3.1); several publications have already addressed the latter subject.<sup>7,26–28</sup> Two main issues are emphasized for each curation procedure: first, we discuss the primary reason why a particular operation should be undertaken and then provide practical technical advice as to how to do it efficiently. Our goal here is to create a repository of good practices for chemical structure curation, not a software tutorial. The complete technical details concerning the use of each software package employed in our recommended curation protocols can be found on the respective developers' Web sites and in user manuals. We do not endorse any of the software packages mentioned in this study; however, we are naturally sensitive to the issue of software availability and did tend to select software that is freely available to academic investigators.

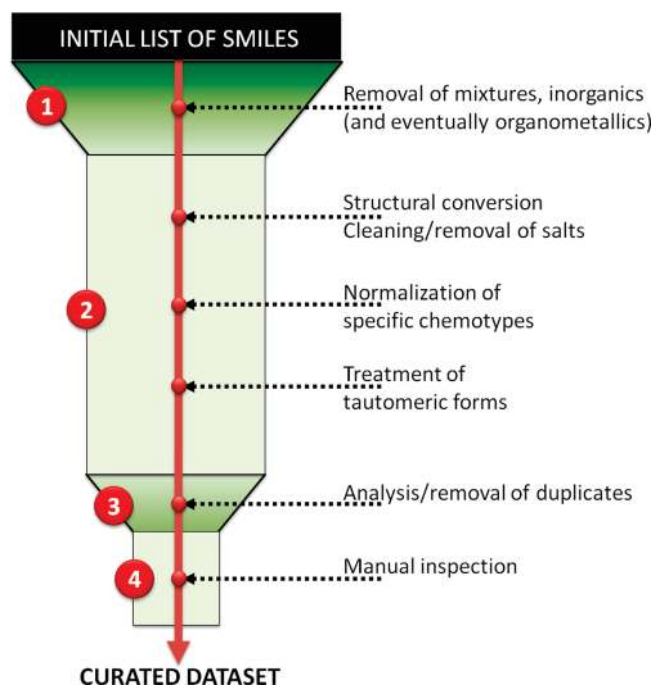


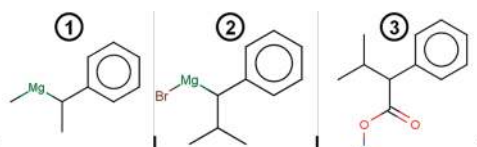
Figure 1. General data set curation workflow.

It is necessary to note that we focus on the 2D level of molecular structure representation. Such limitation assumes that the topological model (or *molecular graph*) implicitly contains most of the essential structural information about a given compound. Thus, the curation procedures described herein lead to cleaned 2D representations of compounds. The methods for efficient conversion of 2D molecular graphs to 3D structures are discussed elsewhere.<sup>29,30</sup>

**2.1. Removal of Inorganics and Mixtures.** Most cheminformatics and QSAR software does not treat inorganic molecules, because the majority of molecular descriptors can be computed for organic compounds only. The inability to model inorganics is an obvious limitation of conventional cheminformatics software. There is a challenging need to develop adequate chemical descriptors for this type of molecules and include them in descriptor calculating software. The fraction of inorganic compounds in most of the available data sets, especially those of relevance to drug discovery is very small. Nevertheless, some data sets generated with the help of automated text-mining approaches extracting data from the literature or electronic sources may contain a significant number of inorganic compounds that are known to have biological effects, for example, toxic effects (cf., section 3.1). At present, all inorganic compounds must be removed before the descriptors are calculated.

Several approaches can be used to rapidly identify and filter out the inorganic compounds. The following protocol is fast and convenient: assuming that SMILES strings for the original data set are available and stored in a single SMI file (each line contains the SMILES string for one compound only), one can calculate their empirical formula (e.g., using the *cxcalc* program included in the ChemAxon JChem package<sup>31</sup>). Then compounds possessing no carbon atoms (i.e., inorganic molecules) can be easily identified and discarded. A simple Perl or Python script can process the entire database in a few seconds, either by analyzing empirical formula or SMILES strings directly. For nonprogrammers representing the majority of cheminformaticians,





DRAGON	REJECTED	REJECTED	REJECTED
MOE	327/333 calculated*	327/333 calculated*	All 2D/3D descriptors calculated
ISIDA	All 2D substructural fragments successfully calculated		
HiT QSAR	All 2D/3D SiRMS descriptors successfully calculated		

\* AM1\_E, AM1\_Eele, AM1\_HF, AM1\_HOMO, AM1\_IP and AM1\_LUMO were not calculated

**Figure 2.** Descriptor calculation for three organometallic compounds using DRAGON, MOE, ISIDA, and HiT QSAR software.

advanced text editors (such as Notepad++<sup>32</sup> for Windows) enable a similar automatic treatment with dedicated tools for substring searching and filtering. Once again, in the end a manual inspection of the SMILES list is recommended.

However, it is much more frequent that a data set includes organic compounds possessing rare elements and organometallics. Dragon molecular descriptors<sup>33</sup> can be calculated only for molecules containing the following 38 atoms: H, B, C, N, O, F, Al, Si, P, S, Cl, Cr, Mn, Fe, Co, Ni, Cu, Zn, Ga, Ge, As, Se, Br, Mo, Ag, Cd, In, Sn, Sb, Te, I, Gd, Pt, Au, Hg, Tl, Pb, Bi. As a result, if some compounds in the data set contain Na, Mg, Ru, etc., they will be rejected by the software (see Figure 2). Meanwhile, such cases could be easily processed by other software, for example, MOE.<sup>34</sup> We shall emphasize that we are not highlighting the relative efficiency of different software to compute descriptors for organometallics but merely point out that cheminformaticians should assess whether their modeling tools can handle such compounds and thus decide if they have to remove or keep them in the data set. There are no available “push-button” solutions for partial data deletion. However, scripts that can identify inorganics by analyzing the compounds’ empirical formula or SMILES strings can also be utilized to detect organometallics and compounds with rare elements.

Similar to inorganics, most current approaches cannot effectively deal with compound mixtures (although a recent study<sup>35</sup> offered an approach to model such systems). Thus, the second important task of this first curation step concerns the identification and the deletion of mixtures. It is of importance to emphasize for beginners that one single SMILES string can code several molecules: for instance the following SMILES string CC(=N[O-])C(=N[O-])C.CC(=N[O-])C(=N[O-])C.[CH2-]I.C1=CC=NC=C1.[Co] contains one cobalt, two molecules of *N,N'*-dioxidobutane-2,3-diimine, one iodomethane, and one pyridine (CID003290398 in the STITCH database<sup>36</sup>). Obviously, it is impossible to calculate descriptors using this SMILES string directly.

Treatment of mixtures is not as simple as it appears. The practice of retaining the component with the highest molecular weight or largest number of atoms is common and widely used, but not necessarily the best solution. The best option is first to delete such records prior to descriptor calculation. However, if there is some reason to believe that the experimentally determined biological activity associated with the record is clearly caused by the largest molecule only and not by mixture itself, it is advisable to use the record for

the largest molecule in the mixture. Such a simple situation is usually possible only for mixtures formed by a relatively large organic molecule and small inorganic molecules, for example, hydrates, hydrochlorides, etc. In such cases, the molecule with the highest molecular weight or the largest number of atoms (preferably) should be retained for the subsequent analysis. Given that descriptors must be calculated for one molecule only, the most relevant compound in the mixture should be determined and selected. Different situations are possible:

- (1) All compounds in the mixture are (or appear to be) identical (e.g., racemic mixtures using 2D representation of molecules). In this case, only one molecule should be kept, and the others should be simply deleted. Certainly, this treatment is only appropriate for 2D QSAR studies when racemic mixture possesses the same activity (property) as corresponding enantiomers.
- (2) The mixture contains one large organic compound and several smaller ones, either organic or inorganic. Generally, it is better to delete the entire record. However, if there are some reasons to believe that the experimentally determined biological activity associated with the record is clearly caused by the largest molecule only and not by the entire mixture, one can keep the record: the compound with the highest molecular weight (or the largest number of atoms) can be kept and the others should be deleted.
- (3) Several similar organic compounds with similar molecular weights are present: these are the most complicated cases, and usually, the deletion of the entire record is recommended (unless the active ingredient is known and can be selected manually) because it is impossible to determine which compound should be retained for modeling using simple rules and automated software. Manual intervention is required for such cases.

For beginners and nonprogrammers, the use of ChemAxon Standardizer<sup>31</sup> is recommended: the treatment of simple cases is fast and simplified by “drag and drop” graphical tools. Experienced users may prefer to use more advanced tools to determine exactly what kind of mixtures (types i, ii, or iii) are present in their data sets.

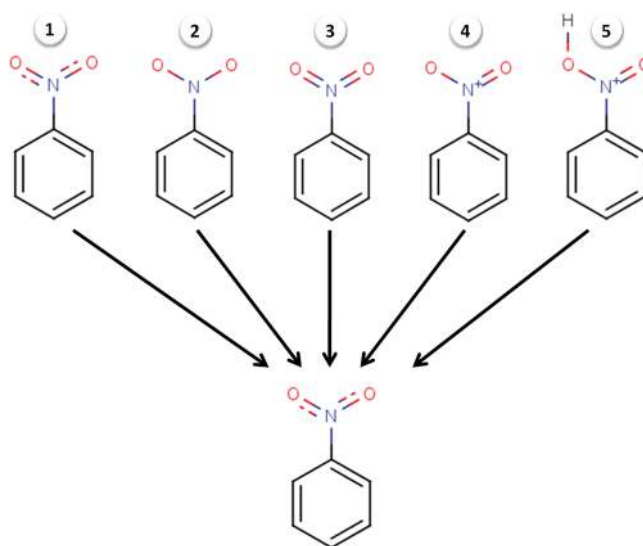
**2.2. Structural Conversion and Cleaning.** The second step of the data set curation entails the conversion of SMILES strings into 2D molecular graphs. Many programs can accomplish this conversion, for example, ChemAxon, MOE,<sup>34</sup> Sybyl,<sup>37</sup> OpenBabel,<sup>38</sup> etc. However, the recent study of Young et al.<sup>8</sup> emphasized the relatively forgotten problem related to the actual reliability of the conversion from SMILES strings to two-dimensional structures. Indeed they used ChemAxon Marvin to convert a library of SMILES and mined the obtained structures for possible errors. Their results showed that very few compounds (4 out of 2118) were converted incorrectly by Marvin; the other errors were related to the presence of wrong initial SMILES strings (because of manual drawing errors or conversion errors from 2D structures into SMILES strings) in the database. This observation suggests that the direct calculation of descriptors from SMILES using any software is much more risky than using alternative formats (e.g., sdf or Mol2) since SMILES do not allow users to visualize, clean, and check chemical structures at 2D level.

Some records in a data set may correspond to salts that are a common form of many drugs. Although properties of

salts can be very different from those of the corresponding neutral molecules<sup>8</sup> and exclusion of salts prior to QSAR analysis is preferred, the removal of metal counterions as well as the neutralization of the remaining carbocations (or carbanions) is still acceptable. Indeed, similar to inorganics, salts are not processed by most of descriptor-generating software, and their presence can generate numerous errors in descriptors' calculation. The neutralization of the charged organic molecules is more disputable: it is quite rare to know precisely the experimental conditions under which the compounds have been tested or the physicochemical environment within cells where the compound is active. In those cases, when the pH of the solution and its exact composition are known, it may be possible to evaluate if a compound should have a charge. Making such prediction still requires the knowledge of  $pK_a$  for each proton donor/acceptor group in the molecule and several available predictors (e.g., from ACDLabs<sup>39</sup> or ChemAxon) could help. When reliable estimates are impossible or if descriptors used for the analysis are insensitive to charge, it is simply recommended to neutralize respective compounds, especially in the case of large data sets containing just a few salts. This task can be successfully realized by MOE,<sup>34</sup> ChemAxon Standardizer, or OpenBabel,<sup>38</sup> which identifies salts, deletes counterions, and then neutralizes the remaining organic compounds. Even zwitterions are successfully treated by most available software. However some difficult cases like compounds where there are covalent bonds between metals and molecules (see compound 3 in Figure 2) can also be encountered. From our experience, such cases are not properly treated by the aforementioned software. Advanced scripts detecting the presence of metals as well as manual curation are thus needed to curate these cases.

Another question concerns the explicit or implicit presence of hydrogen atoms in the structures. In our experience, the use of explicit hydrogens for calculating 2D descriptors leads in most cases to QSAR models with higher prediction performances. However, our experience also suggests that sometimes the use of explicit hydrogens may also introduce noise in the descriptor matrix (especially when using fragment-based descriptors) and, thus, lead to less reliable models. Many software packages claim to have reliable procedures for adding/removing hydrogens. For instance, our experience indicates that removing hydrogens is not well realized in certain cases, for example, when hydrogens are attached to nitrogens in rings or in secondary amines. In those cases, hydrogens are not always removed, which leads to error messages in programs calculating descriptors parametrized for the treatment of hydrogen-depleted graphs, as well as incorrect descriptor values.

**2.3. Normalization of Specific Chemotypes.** Very often the same functional group may be represented by different structural patterns in a given data set. For example, nitro groups have multiple mesomers and, thus, can be represented using two double bonds between nitrogen and oxygens (neutral form), or one single bond linking the nitrogen and the protonated oxygen, or linking both nitrogen and oxygen atoms that are oppositely charged (Figure 3). For cheminformaticians, these situations may lead to serious inconsistency problems, because molecular descriptors calculated for these different representations of the same chemical group would be significantly different. For example, if two identical



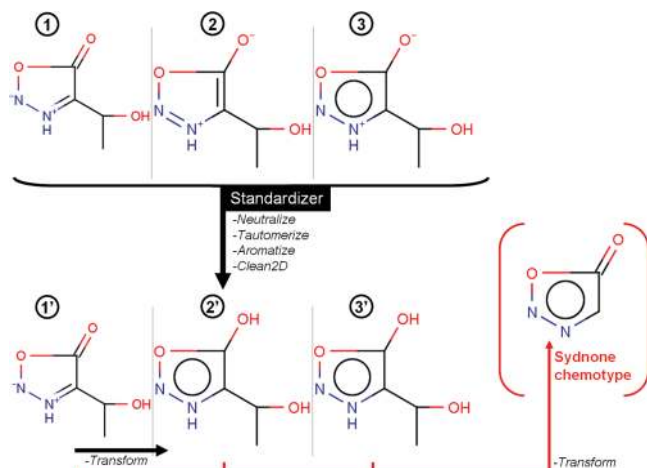
**Figure 3.** Structure normalization: five types of nitro group representations retrieved in the nitroaromatics data set for rats and *T. pyriformis* case studies (see section 3.2 in the text for details).

compounds contain a nitro group represented by two different patterns, they will not be recognized as identical by conventional similarity metrics because some of their computed descriptors will be different.

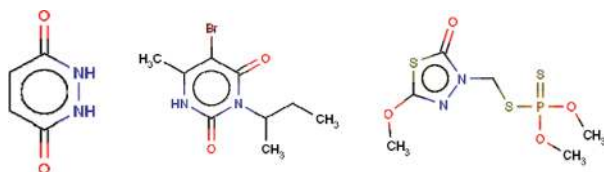
Manual conversion of all functional groups to some standard forms is too time-consuming and could introduce additional human-dependent nonsystematic errors. ChemAxon's Standardizer is probably the most well-known tool to rapidly and efficiently realize chemotype normalizations. Users can manually draw the pattern conversion for several functional groups and store them in a dedicated reusable *xml* rule file. Taking into account the specifics of individual research laboratories (modeling software, descriptors, etc.), the users could build a customized database of functional group conversions, which can be applied to every new data set. Thus, beginners can directly use the library of conversion rules developed by more experienced modelers to treat their data sets in the proper way.

Although ring aromatization and the normalization of carboxyl, nitro, and sulfonyl groups are relatively obvious, more complex cases like anionic heterocycles, polyzwitterions, tautomers, etc., require a deeper analysis and multiple normalization steps. To illustrate this point, we chose three compounds possessing the sydnone chemotype (Figure 4), represented by its different mesomeric forms. The application of the Standardizer with classical settings (neutralize, tautomerize, aromatize, and clean2D) was able to normalize two of the three compounds (2' and 3'), whereas the first compound was not aromatized. A second step using the "transform" function (allowing the conversion of user-defined groups) led to the normalized forms (2' and 3') of the compound 1. Depending on the modeled property, the experimental conditions and the other compounds in the data set, one can transform these compounds into the formal sydnone chemotype (possessing the keto group) like in Figure 4. But, it has to be noted that such notation (aromatic ring and the branched keto group) will be rejected by many descriptor calculating software (Figure 5).

In some cases, compounds may exist in several tautomeric forms,<sup>18</sup> the most common ones being the keto-enolic



**Figure 4.** Use of ChemAxon Standardizer to normalize three compounds possessing the sydnone chemotype (see text for details).



**Figure 5.** Examples of misleading structure representations produced by the “general style” option available in ChemAxon Standardizer, which may serve as a potential source of errors for programs calculating molecular descriptors.

tautomers. Choosing one form instead of another could have a significant impact on the prediction performances of QSAR models built with such data. As suggested in the study by Young et al.,<sup>8</sup> discarding one tautomer form may be realized taking into account the compound’s mechanism of action, if the latter is known for the studied biological activity. For instance, the choice between keto–enolic tautomeric forms may be influenced by the knowledge about the formation of a specific hydrogen bond with the target receptor, or an aromatic ring. Moreover, experimental conditions of the modeled chemical system (especially, pH) are crucial. A remarkable review addressing different problems related to tautomers was recently published by Dr. Yvonne Martin.<sup>18</sup>

**2.4. Removal of Duplicates.** Rigorous statistical analysis of any data set assumes that each compound is unique and thus, structurally different from all other compounds. However, structural duplicates are often present in chemical data sets, especially in large ones. For instance, the same compound could be ordered for a screening campaign from two different sources, given different internal IDs, and thus, two corresponding records would be placed in a database (sometimes, with rather different values of the associated experimentally measured property or bioactivity). QSAR models built for such data sets may have artificially skewed predictivity (see section 3.4) even if a data set contains only a small percentage of duplicates. Duplicates can also affect the observed frequency of a given chemotype in a data set, the distribution of compounds according to their structural similarity, etc. As a consequence, duplicates must be removed prior to any modeling study. The first step of the procedure requires the detection of identical molecular structures within

the set, whereas the second step is dedicated to the comparison of the studied property values for the retrieved duplicates.

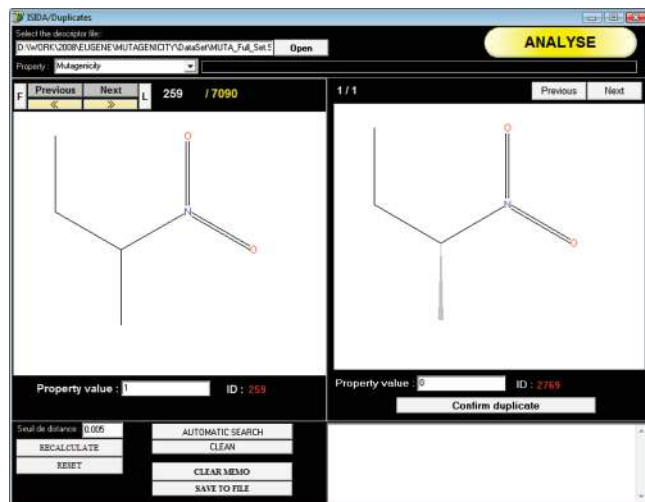
A current practice consists of identifying duplicates from SMILES strings, which is correct if and only if the latter are canonical SMILES. However, the experience shows that most beginners are not aware of this requirement and often use noncanonical SMILES to identify chemical duplicates. Thus, it is of importance to underline that a given compound can be represented by several SMILES strings: for instance, three formally different SMILES strings, that is, O=C(OCC)C, C(=O)(OCC)C, and O(C(C)=O)CC, can code the same compound, that is, ethyl acetate. Without the standardization of these SMILES into the CCOC(C)=O, the canonical form (different from the three previous strings), it is impossible to identify them as duplicates in a particular data set from SMILES strings alone. The calculation of empirical formula from SMILES represents an additional filter to retrieve duplicates.

Once duplicates are identified, the analysis of their properties is mandatory, requiring some manual effort. For a given pair of duplicate structures, if their experimental properties are identical, then one compound should be merely deleted. However, if their experimental properties are numerically different, we shall consider two main scenarios for data curation:

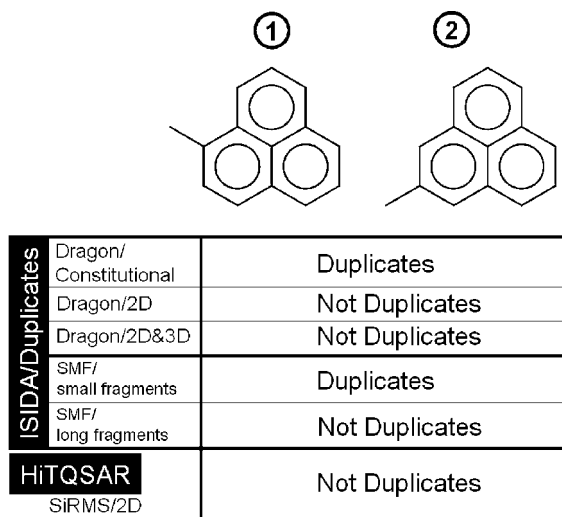
- (1) The property value may be wrong for one compound because of, for example, a human error when the database was built (these types of errors are often manifested in significant outliers when QSAR models are either built or employed for external predictions). Another frequent case is when the data are compiled from literature sources and the same compound was tested in two or more different laboratories under possibly different experimental conditions, variations in the protocol, etc., leading to the difference in (formally the same) measured property. In this case, supplementary investigations must be done to decide if both entries should be deleted from the data set or if only one should be deleted. If the data set is large enough and there is no obvious explanation of such discrepancies, we recommend to place such cases into a special external test set including all suspicious records and try to reveal the most likely true value by comparing experimental records to the results of consensus predictions from statistically significant and externally validated QSAR models.
- (2) Both experimental properties are correct but the previous curation tasks (for example, the removal of counterions in salts) have modified the substance records to create such duplicates. For instance, the two records could correspond to two different salts of the same compound (or a neutral compound and its salt). As previously mentioned, their experimental properties can indeed be very different. If both experimental properties are highly similar, the record can be kept associating the structure with the arithmetic average of properties. If they are significantly different, we recommend eliminating both records.

To successfully and rapidly achieve the removal of duplicates, we currently recommend both ISIDA/Duplicates<sup>40</sup> (see Figure 6) and HiT QSAR;<sup>41,42</sup> these programs are free for academic laboratories and complementary to each other. For





**Figure 6.** Automatic retrieval of structural duplicates using the ISIDA/Duplicates program: example of stereoisomers (Ames mutagenicity data set) with opposite mutagenicity properties.



**Figure 7.** Two structural isomers retrieved as either duplicates or nonduplicates by ISIDA/Duplicates and HiT QSAR according to different pools of chemical descriptors.

each pair of compounds, ISIDA/Duplicates calculates “on the fly” the Euclidean distances between them using the input descriptor matrix of the data set. Then, all pairs with a distance lower than or equal to a user-defined threshold (zero by default) are considered duplicates. Efficiency depends mainly on the type and the number of descriptors used to represent compounds. The definition of duplicates is strongly associated with the types of descriptors employed to characterize chemicals: for example, two isomers of the same structure would be identified as duplicates if the descriptors used are insensitive to branching in the structures, and two stereoisomers will be considered as duplicates in the case where descriptors (e.g., most molecular connectivity indices) do not take into account the chirality. It has to be pointed out that there is no set of descriptors universally recognized to be best for duplicate recognition. An example is given in Figure 7 where two isomers have been processed by ISIDA/Duplicates. If simple descriptors like Dragon/constitutional or ISIDA/small fragments are used, these two compounds are identified as duplicates. One can note that this approach can be useful to rapidly detect pairs of isomers in the database. The use of more complex descriptors such as Dragon/2D and ISIDA/long fragments discriminates the two

isomers which are not identified as duplicates anymore. ISIDA/Duplicates natively calculates long fragment descriptors<sup>43</sup> that take into account molecular branching and atom connectivity properties, and can also import external sets of descriptors (like Dragon, MOE, etc.) to take into account chirality and other three-dimensional properties. The program can automatically identify duplicates and output the resulting list, as well as a curated library of compounds.

In HiT QSAR the search for duplicates is performed by an innovative *one-click* tool implementing the CANON algorithm that employs the canonical numbering using the atom connectivity matrix. Each molecule is represented as a string reflecting the empirical formula/order of connectivity for each atom, for example, benzene would be represented as C<sub>6</sub>H<sub>6</sub>/1\_2a,1\_3a,1\_7s,2\_4a,2\_8s,3\_5a,3\_9s,4\_10s,4\_6a,-5\_11s,5\_6a,6\_12s/, where atoms 1–6 are carbons and 7–12 are hydrogen atoms. If such strings are similar for different records, the respective compounds are reported as duplicates. However, this approach does have certain limitations: for instance, cis–trans and (R–S) isomers, as well as diastereoisomers, are considered as duplicates. For this reason the use of both ISIDA/Duplicates and HiT QSAR programs in concert leads to high retrieval rates of real structural duplicates.

**2.5. Final Manual Checking.** The last step of the curation entails manual inspection of every molecular structure (as much as possible for the large data sets). We recommend inspecting the curated data set carefully to establish the types of chemical scaffolds present in the data set, their relative proportion in the set, etc. Obviously, for large data sets (more than a thousand of compounds), this step is time-consuming and extremely laborious. However, several pieces of advice can be formulated to reduce the amount of effort: for instance, to check only compounds with complex structures or having a large number of atoms. Another apparent solution is to generate a representative sample of the set and then, check it for the presence of potential erroneous structures (rechecking of the whole data set may become unavoidable if significant errors are found).

Common errors identified during the manual cleaning procedure may have different origins:

- (1) The structure is wrong: A rapid check of both IUPAC compound's name (if available) and its structure is essential to identify possible errors concerning the scaffold and positions of substituents (e.g., because of manual errors or program bugs<sup>8</sup> in the conversion of SMILES into 2D structures). Actually, the identification of incorrect structures is the most difficult part of the data curation. The majority of structures are incorrect due to random human errors when the structures have been drawn/converted in an electronic format. Although it is relatively easy for a small data set to check each individual structure and search for perfect agreement between chemical names and the actual structures, it becomes unfeasible for large data sets. For example, it would take a restrictive amount of time to discover that particular chlorine has to be in position 2 and not in position 1 in the 58 653th compound of a studied data set. Therefore, we suggest the following protocol: with the development of numerous freely available chemical databases, it is now relatively simple to mine these databases and retrieve chemical structures from a list of names or CAS numbers. Several entries for each name

**Table 1.** Summary of Major Procedures and Corresponding Relevant Software for Every Step of the Data Curation Process<sup>a</sup>

procedures	software	availability
inorganics removal	ChemAxon/Standardizer	free for academia <sup>31</sup>
structure normalization (fragment removal, structural curation, salt neutralization)	OpenEye/Filter	free for academia <sup>67</sup>
	ChemAxon/Standardizer	free for academia <sup>31</sup>
duplicate removal	OpenBabel	free <sup>38</sup>
	Molecular Networks/CHECK,TAUTOMER	commercial <sup>68</sup>
	ISIDA/Duplicates	free for academia <sup>40</sup>
SDF management/viewer file format converter	HiT QSAR	free for academia <sup>42</sup>
	CCG/MOE	commercial <sup>34</sup>
	ISIDA/EdiSDF	free <sup>40</sup>
	Hyleos/ChemFileBrowser	free <sup>69</sup>
	OpenBabel	free <sup>38</sup>
	ChemAxon/MarwinView	free for academia <sup>31</sup>
	CambridgeSoft/ChemOffice	commercial <sup>70</sup>
	Schrödinger/Canvas	commercial <sup>71</sup>
	ACD/ChemFolder	commercial <sup>39</sup>
	Symyx/Cheminformatics	commercial <sup>72</sup>
CCG/MOE	commercial <sup>34</sup>	
Accelrys/Accord	commercial <sup>73</sup>	
	Triplos/Benchmark Pantheon	commercial <sup>37</sup>

<sup>a</sup> We invite all interested scientists to enrich this table by adding their preferred procedures and relevant software to the open document available at our web site, <http://mml.unc.edu>.

or CAS ID are likely to be retrieved because of known overlaps between databases; some of them are coded as SMILES strings and some as 2D/3D structures stored in mol, mol2 files, etc. The analysis of these multiple entries for every single compound is then critical: since a given IUPAC name or CAS number is unique for every compound, all structures retrieved using a given query (e.g., 2-chlorobenzylsulfonamide or CAS #89665-79-2) must represent exactly the same compound. On the other hand, one or several (or all) entries of databases may be wrong. The curation task is then to verify whether the actual structure used in modeling corresponds to the structures retrieved from the database mining exercise. There are no freely available tools specifically dedicated to this task, but we believe that challenges of data integration in bio- and cheminformatics will highlight such tools as being critical for efficient merging between different databases. We also consider that the comparison between different structures retrieved for a given query should be realized at two-dimensional level (three-dimensional if the stereochemistry is specified) but not using SMILES only (there is no guaranty that all SMILES strings in all databases are canonical).

- (2) The normalization of bonds is incomplete: common mistakes are related to the presence of different representations of the same functional groups. Despite the normalization procedure, some very specific cases can still be present, and thus, the corresponding chemotypes must be corrected manually.
- (3) Some duplicates may still be present despite the use of automated software to remove them. For instance, some tautomers can still be found. Advanced tools developed internally in private companies or in academic laboratories capable of such fine filtering may exist but they appear to be unavailable in the public domain.
- (4) Other possibilities: wrong charges, presence of explicit hydrogens in a hydrogen depleted structure, incorrect bonds, etc.

**2.6. General Remarks and Disclosure.** We would like to stress that the main purpose of this report is not software comparison; this would be simply impossible since the

majority of such software is likely hidden within industrial laboratories. We are mostly concerned with possible sources of inaccurate structures and adequate procedures implemented in available software that should be followed to correct the erroneous data records. Thus, we have attempted to make Table 1, a repository of available software dedicated to data curation, as complete as possible, concerning cases, procedures and listed selected software that is capable of making the requisite corrections. The only bias that we had was toward software that can be obtained free of charge for academic investigators or with which we had firsthand experience. Our laboratory is not affiliated in any special way with any of the software vendors and therefore any mention of any software should not be regarded as an advertisement.

### 3. EXAMPLES OF APPLICATIONS RELYING ON DATA SET CURATION

**3.1. Cheminformatics Analysis of Compounds Inducing Liver Injuries.** Drug-induced liver injury (DILI) is one of the main causes of drug attrition.<sup>44–46</sup> Elimination of drug candidates likely to cause hepatotoxicity at early stages of drug discovery workflow could significantly increase the rate and reduce the cost of drug development. The ability to predict DILI effects of drug candidates from their chemical structure is critical to help guiding experimental drug discovery projects toward safer medicines. More generally, there is now a great deal of interest both in the U.S. (for instance, with the Toxcast<sup>47</sup> program) and Europe (the REACH regulation<sup>48</sup>) in developing fast and accurate experimental and computational approaches to predict toxic effects of chemicals, for example, hepatotoxicity.

A large amount of published information that could improve our knowledge about DILI mechanisms is available, but the information is spread over a large body of publications using inconsistent terms. Recently, our group in collaboration with the Biowisdom company<sup>49</sup> launched a project concerning the cheminformatics analysis of assertions mined from the biomedical literature that describe DILI effects of chemical compounds. BioWisdom's Sofia platform



**Table 2.** Statistical Parameters of QSAR Models Obtained before and after Curation<sup>a</sup>

ID	name	$R^2$	$Q^2$	$R_{EF}^2$	$S_{ws}$	$S_{cv}$	$S_{EF}$	$R_{EVS}^2$	$R_{EVS(NM)}^2$
1	Rat	0.96	0.84–0.93	0.89–0.92	0.11–0.13	0.16–0.24	0.20–0.26		
2	Rat <sub>(NM)</sub>	0.91–0.97	0.89–0.95	0.45–0.88	0.10–0.18	0.14–0.28	0.28–0.58		
3	TP	0.83		0.76	0.33		0.38	0.54	–0.58
4	TP <sub>(NM)</sub>	0.85		0.54	0.31		0.54	0.49	0.44
5	DILI noncurated	no modeling was possible							
6	DILI <sup>50</sup>	modeling set 5-fold external CV accuracy = 62–68% external sets accuracy = 56–73%							
7 <sup>b</sup>	<sup>62</sup> Ames noncurated	sensitivity <sub>RF</sub> = 83%; sensitivity <sub>SVM</sub> = 87%; specificity <sub>RF</sub> = specificity <sub>SVM</sub> = 75% AUC <sub>GP</sub> = 88%; AUC <sub>SVM</sub> = 89%; AUC <sub>RF</sub> = 83%							
8 <sup>b</sup>	<sup>63</sup> Ames curated	sensitivity <sub>RF</sub> = sensitivity <sub>SVM</sub> = 79%; specificity <sub>RF</sub> = specificity <sub>SVM</sub> = 81% AUC <sub>GP</sub> = 86%; AUC <sub>SVM</sub> = 84%; AUC <sub>RF</sub> = 83%							

<sup>a</sup> Where TP = *Tetrahymena pyriformis* data set, NM = modeling set with various representations of nitro groups,  $R^2$  = determination coefficient,  $Q^2$  = cross validation determination coefficient,  $R_{EF}^2$  = determination coefficient for external folds extracted from the modeling set,  $S_{ws}$  = standard error of a prediction for work set,  $S_{cv}$  = standard error of prediction for work set in cross validation terms,  $S_{is}$  = standard error of a prediction for external folds extracted from the modeling set,  $A$  = number of PLS latent variables,  $D$  = number of descriptors,  $M$  = number of molecules in the work set,  $R_{EVS}^2$  = determination coefficient for external validation set,  $R_{EVS(NM)}^2$  = determination coefficient for external validation set with shuffled nitro groups, AUC = area under curve statistical parameter, RF = random forest, SVM = supporting vector machine, and GP = Gaussian processes. <sup>b</sup> Prediction performances are reported for external validation set.

(<http://www.biowisdom.com/>) was used to generate assertional meta-data, comprising thousands of highly accurate and comprehensive observational statements. These statements are represented in triple constructs: *concept\_relation-ship\_concept*, for example, *Cafestol\_suppresses\_Bile acid biosynthesis*, *Azathioprine\_induces\_Cholestasis*, etc. Each assertion is derived from and evidenced by a variety of electronic data sources. More importantly, the assertional meta-data have been collected across different species, i.e., human, rodent and nonrodent animals.

As is often the case for data sets based on literature sources, the initial data set resulting directly from text mining described compounds by chemical names only. In cases like this, a large number of duplicates is expected because many compounds are described in the literature using many different names; for instance, the ChemSpider<sup>4</sup> search using “aspirin” as a key word indicates that there are more than 35 synonyms that are commonly used for aspirin (or refer to aspirin in literature) and, in total, more than 180 terms (drug names, usual names, etc.) that are related to aspirin. In this recent study,<sup>50</sup> after the identifications and curation of obvious chemical synonyms, a set of 1061 compounds (stored as SMILES strings) was retrieved from the analysis of assertions mined from Medline abstracts using text-mining tools. Unlike most traditional QSAR data sets, the compounds were extremely diverse with almost all possible problematic cases in terms of data set curation: presence of numerous inorganics, mixtures of organics and inorganics, salts, zwitterions and duplicates. No software would be able to compute relevant descriptors for this exotic data set. As a result no possible cheminformatics analysis (especially QSAR modeling) was possible for this data set without thorough data curation (see Table 2). Both automatic and manual procedures have been employed to clean this data set as follows:

(1) Initially, all inorganic compounds have been removed since our data analysis strategy includes the calculation of molecular descriptors for organic compounds only. We should emphasize again that this is an obvious limitation of many cheminformatics approaches since inorganic molecules are definitely known to induce liver injuries; however, the total fraction of inorganics in our data set was relatively small. For example, the following

**Table 3.** Number of Investigated Compounds in the Data Sets before and after Curation

data set	number of compounds	
	original set	curated set
liver toxicants (DILI)	1061	951 (90%)
nitroaromatics (rats)	28	28 (100%)
nitroaromatics ( <i>T. pyriformis</i> )	95	95 (100%)
ToxRefDB	320	292 (91%)
Ames mutagenicity	7090	6542 (92%)
bioavailability (UCSD)	805	734 (91%)

compounds have been removed: activated charcoal, cobalt dichloride, ferrous sulfate, zinc chloride, sulfur, cis-diaminedichloroplatinum, manganese chloride, etc. Moreover, additional compounds were removed because (i) their corresponding SMILES strings could not be identified unequivocally due to the inconsistent name or irrelevant labeling code or (ii) they corresponded to a mixture of compounds (for example, Gramicidin, which involves six antibiotic molecules). Thus, after this step, 993 compounds remained.

- (2) Then, 2D molecular structures (chemical connectivity maps) have been generated from SMILES strings using the ChemAxon’s JChem 5.1 program under the control of ISIDA to create a unique SDF file containing both structures and DILI profiles. We also used Standardizer to remove all counterions, clean records including multiple compounds, clean the 2D molecular geometries and normalize bonds (aromatic, nitro groups etc.) as described in section 2.
- (3) Finally, duplicate molecular structures were detected automatically using the ISIDA/Duplicates<sup>40</sup> program, followed by careful manual inspection of the entire data set. 951 compounds remained out of the 1061 initial molecules (i.e., as much as ~10% of the data set was eliminated).

The main objective of the study was to demonstrate the usefulness of *classical* cheminformatics approaches to analyze assertions of drug-induced liver effects in different species, and more precisely, to explore the relationships between chemical structures and animal DILI toxicity. Thus, our goal was to extract knowledge about the influence of specific scaffolds and chemotypes on DILI. After the critical

step of chemical data curation, we have explored the issue of concordance of liver effects across species and found that the concordance values between any two species from the three groups studied, i.e., humans, rodents and nonrodent animals, were relatively low (40–45%), which was in agreement with earlier studies reported in the literature.<sup>51–53</sup> The subsequent cluster analysis<sup>54</sup> of the 951 remaining compounds using 2D fragment descriptors<sup>43</sup> allowed us to identify multiple clusters of compounds belonging to structurally congeneric series. Similar liver effect profiles have been observed for most clusters although some compounds appeared as outliers. In several cases of such outliers, additional focused mining of public data sources led to revised assertions that were more in tune with DILI profiles expected on the basis of chemical similarity. Thus, the chemical similarity analysis was helpful in focusing on possible gaps in assertion data for liver effects reported in the literature for different species and correcting the erroneous or missing assertions. In addition, binary QSAR models of liver toxicity were derived and the mean external prediction accuracy in 5-fold external validation study was found to be 65%.

To enable the profile analysis and QSAR modeling using this large data set extracted from literature using automatic text mining tools, it was essential to utilize various data curation procedures described above. Data cleaning and standardizing was critical since no investigation at all would be possible without having consistent structural representations, correct chemical descriptors and the absence of duplicates. We believe that our study<sup>50</sup> presented the first example when rigorous text mining and cheminformatics data analysis were combined toward establishing predictive models of chemical toxicity.

**3.2. QSAR Modeling of Nitroaromatic Toxicants.** There is a public concern and a strong need in evaluating the potential environmental risks associated with the production, storage, and application of explosive compounds, many of which are nitro- and polynitroaromatics. Recently, a data set of nitro-aromatic compounds of military interest was compiled from different sources to investigate the relationships between their chemical structure and toxicity. Each compound was manually inspected in order to create a curated data set. During this process, five different representations of nitro groups were identified (Figure 3). Obviously, the difference in one or two bonds may appear to be insignificant in the context of the entire compound, but in reality, those inconsistencies in the molecular representation of the same functional group could actually lead to different descriptors of the same molecule and, in some cases, to poor QSAR modeling results.

Two sets of nitroaromatic derivatives were compared

- (1) Case Study 1: 28 compounds (2 aromatics and 26 nitroaromatics) tested in rats to evaluate their animal toxicity expressed as  $\log(\text{LD}_{50})$ , mmol/kg.<sup>55</sup>
- (2) Case Study 2: 95 compounds tested against *Tetrahymena pyriformis*, a ciliated freshwater protozoan, to assess their aquatic toxic effects expressed as  $\log(\text{IGC}_{50})$ , mmol/mL.<sup>56</sup>

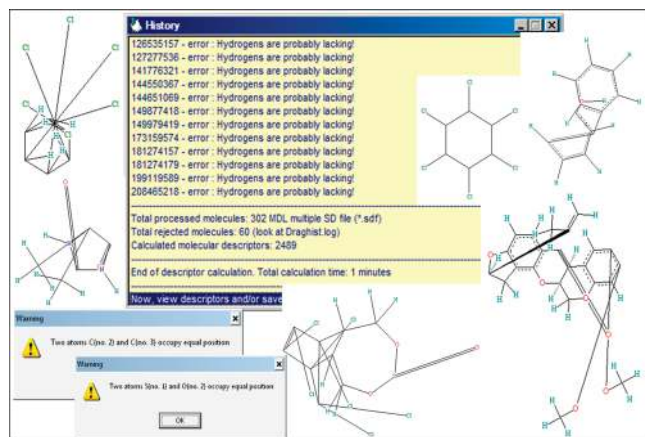
In the first case study, five different representations of nitro groups were equally distributed within the modeling set. Initially three different overlapping validation sets were selected according to.<sup>55,57</sup> Each external set consisted of one aromatic compound and five nitroaromatics with different

types of nitro group representation. The sets with compounds possessing various types of nitro group representations were referred to as “mixed”. 2D simplex descriptors, PLS statistical approach, and principles of external test set formation as described in Kuz'min et al.<sup>55</sup> were used. Three models were developed using the mixed sets (Table 2). One of them had the same predictivity as original models<sup>55</sup> ( $R_{\text{ext}}^2 \approx 0.9$ ), whereas the prediction accuracy for the two others was much lower ( $R_{\text{ext}}^2 = 0.45–0.60$ ). Thus, in one case the differences in the nitro group representation had no effect on model predictivity, whereas in the two others the prediction performance decreased significantly. At the same time the goodness-of-fit and robustness of both groups of models for the training set were equal (Table 2), confirming the well-known Kubinyi paradox<sup>58</sup> and the necessity of external validation.<sup>59</sup> Thus, one can conclude that simultaneous usage of different types of nitro group representation can significantly influence the predictive ability of the models.

In the second case study, 60% of the modeling set compounds possessed nitro groups represented with aromatic bonds (see Figure 3), whereas the remaining four classes of the nitro group patterns were equally represented (10% each). The same proportions were kept for the external validation set consisting of 63 compounds. The statistical metrics resulting from 5-fold external cross-validation of the modeling set and an external validation set were selected as a measure of model predictivity. Similarly to the first case study, the same descriptors and statistical approaches as in the original study<sup>56</sup> were used for model generation for the mixed set. Goodness-of-fit and robustness of the original<sup>56</sup> and shuffled models were equal (Table 2). Overall comparison of model predictivity in the 5-fold external cross-validation experiment shows that the original consensus model is better than the consensus model obtained using the mixed nitro groups. However, their prediction performances estimated using the original external validation set were almost similar. Even more impressive and unexpected results were obtained for the external set when nitro groups were mixed in the same way as for the training set. In the case of a model built on the mixed set, the mixing of the nitro group representation among external validation set compounds did not significantly affect the predictivity ( $R_{\text{ext}}^2 = 0.49$  and  $R_{\text{ext(NM)}}^2 = 0.44$ ), whereas the difference was dramatic for the original model ( $R_{\text{ext}}^2 = 0.54$  and  $R_{\text{ext(NM)}}^2 < 0$ ) (Table 2). Here, the index “NM” refers to the set with the mixed nitro groups.

The results obtained here with two different nitroaromatics data sets validate the necessity of the nitro group standardization. They show that even small differences in structure representation can lead to significant errors, and even robust and inherently predictive models can fail on noncurated external validation sets.

**3.3. ToxRefDB.** At the time of the study, the original version of the ToxRef DataBase (<http://www.epa.gov/NCCT/toxrefdb/>)<sup>60</sup> contained 320 compounds tested for their carcinogenicity in both rats and mice (totally 26 panels of experimental data represented as binary results, that is, toxic or nontoxic). The initial efforts to generate QSAR models for these compounds were unsuccessful: we could not build any statistically significant model based on our standard QSAR modeling workflow.<sup>25</sup> Thus, we have examined the data set for possible errors as follows (and we believe this



**Figure 8.** Real examples of erroneous structure records in chemical databases leading to Dragon error messages.

exercise could be illustrative as an example of a training session on the protocol for data examination prior to model building).

Even a quick examination of the data set (see Figure 8) revealed that a deep cleaning of 2D structures, addition of explicit hydrogens and standardization of problematic nitro and carboxylic groups was necessary. After application of all of the data set curation procedures discussed in section 2, as well as aromatization, mixtures and salts removal, and standardization of nitro and carboxylic groups, only 293 compounds remained. Subsequent search for duplicates using HiT QSAR and ISIDA software revealed the presence of the (S)-isomer of bioallethrin and its racemate. Since stereochemical information was available for one pair of compounds only, these structures were marked as duplicates on 2D level. Thus, 291 from the original 310 structures were accepted for subsequent modeling (see Table 3).

However, further visual inspection of these structures revealed some misleading representations, especially with respect to unexpected assignment of aromatic bond types. This occurred because of the choice of “General style” instead of the “Basic style” option in the ChemAxon Standardizer. Unlike the “Basic style”, the ring of 2-pyridone, for instance, is not aromatized under the “Basic style” option. If the “General style” is effectively correct from the chemistry viewpoint, it has to be stressed that many cheminformatics programs will generate errors or simply reject the compound because a carbon atom is not formally tetravalent under this “General style” representation (see Figure 5). Additional information can be found at the following ChemAxon Web site (<http://www.chemaxon.com/jchem/marvin/help/sci/aromatization-doc.html>). This example demonstrates that there is no simple “push-button” solution for chemical data curation and that data inspection and curation including manual involvement is necessary.

Following the chemical record cleaning, the data set appeared ready for the QSAR modeling. However, the curation procedures should not be limited to structure analysis but should also include the evaluation of the quality of the experimental data (as our earlier example with the analysis of DILI data suggests). Thus, we applied ISIDA/Cluster<sup>40</sup> to group similar compounds into clusters. With ISIDA visualization tools, we rapidly identified some suspicious pairs of highly similar compounds (e.g., with only a methyl group in a different position) that nevertheless had large

differences in their toxicity profiles. Some of these cases, for example, ametryn and prometryn could be classical “activity cliffs”, but we also found true and suspicious cases of erroneously annotated compounds, for instance, atrazine. At the time of the study and even in a recent version of the ToxRefDB ([http://www.epa.gov/NCCT/toxrefdb/files/ToxRefDB\\_ChronicCancer\\_2009Apr06.xls](http://www.epa.gov/NCCT/toxrefdb/files/ToxRefDB_ChronicCancer_2009Apr06.xls)), atrazine is annotated as a nontumorigen for both rats and mice. Conversely, two compounds, propazine and simazine, identified by ISIDA/Cluster as structural neighbors of atrazine and different from the latter by presence or absence of only one methyl group, are both annotated as tumorigen agents for rats. After additional investigation we found literature evidence that atrazine has been reported elsewhere as a rat tumorigen.<sup>61</sup> We believe that this example suggest a crucial importance of verifying not only molecular structures but also activity data using cheminformatics tools such as clustering by compound similarity and the analysis of property distributions. The results of QSAR modeling of the curated data set will be published separately.

**3.4. Ames Mutagenicity.** Recently our group initiated a collaborative modeling project involving many international participants to develop most significant models of the Ames mutagenicity; each research group was expected to use different descriptors and machine learning approaches. The data set, kindly provided by Dr. K. Hansen,<sup>62</sup> consists of 7090 compounds classified as mutagenic or nonmutagenic. Briefly, frame-shift mutations or base-pair substitutions can be detected in the Ames test by the exposure of histidine-dependent strains of *Salmonella typhimurium* to a given compound. Herein, mutagenicity is represented in a binary format: a compound is classified as positive (mutagenic) if it significantly induces revertant colony growth at least in one strain. A compound is labeled negative (nonmutagenic) if it does not induce revertant colony growth in any strain tested.

The original data set was curated using both HiT QSAR and ISIDA software: (1) All structural duplicates were removed. If both molecules (according to 2D structures) had the same mutagenicity effect, then one of them was removed, and if both molecules had different mutagenicity effects, then both were deleted. (2) All inorganic compounds were excluded. (3) The remaining structures were cleaned using the ChemAxon Standardizer and HiT QSAR software. (5) The last step before modeling was the repetition of duplicates search and careful manual checking.

(1) One of the most important steps of the curation procedure is the removal of duplicates: 518 pairs of structural duplicates (at 2D level of structure description) were found by both HiT QSAR and ISIDA software (Figure 6). For the original data set the situation was as follows: 80% of compounds were represented by 2D structures without any information about stereochemistry, and for approximately 20% of the compounds, stereochemical information was available. However, in most cases only one of the two enantiomeric forms was characterized. In some cases (about one hundred), the experimental information for both (R)- and (S)-stereoisomers or for two or more diastereoisomers was reported. Moreover, only 7 out of one hundred pairs created by different enantiomers had different mutagenicity. Thus, one can conclude that for the most part



the stereochemistry of investigated compounds does not influence their mutagenic effect. As a result, it appeared logical to represent molecular structures at 2D topological level for further QSAR analysis.

Most of the pairs were formed by *classical* duplicates, that is, identical (topologically or topologically and stereochemically if applicable) structures with the same mutagenicity property; for instance, 2-((4-chlorophenyl)methyl)-oxirane was found twice in the original data set. Some duplicates represented stereoisomers (R–S or diastereomers) with the same mutagenicity (e.g., (*R*)- and (*S*)-penbutolol). In this case, one of the stereoisomers was removed from the data set. The same procedure was applied for diastereomers with identical mutagenicity (e.g., ursodiol and chenodiol).

The situation when regular duplicates, similar or different enantiomers ((*R*)- and (*S*)-2-nitrobutane) or diastereomers had different mutagenicity values was rare (~ 30 pairs including 7 for different enantiomers), and probably, it could be explained by the presence of errors in the interpretation of experimental data. All such suspicious records were excluded from the subsequent analysis. However, in the absence of data curation, we could easily foresee a situation when identical compounds with identical mutagenicity would be distributed between training and test sets. Should this happen we would expect to observe an artificially enhanced predictive accuracy of the training set models. We shall use this example to illustrate how data set curation may help design and/or tune more efficiently the modeling parameters such as descriptor types or the machine learning approach. Thus, after duplicates analysis and removal, 6572 compounds remained.

The group that provided us with the Ames data set has already published the preliminary results of their QSAR studies;<sup>62</sup> however, the statistical parameters of their models using the noncurated data set of 7090 compounds were, probably, overestimated since almost 9% of compounds should have been removed. To assess the consequences of their presence in the data set, we conducted the following study. The data set was randomly divided into two subsets five times following our standard routine for generating training and validation sets for QSAR modeling and then the content of both sets was analyzed in terms of the presence/absence of duplicates. Results showed that 229–255 out of 518 pairs of duplicates were split between the modeling and external validation sets (it corresponds to the probabilistic distribution). This situation could lead to overestimating the model predictivity, despite the usage of three-dimensional structures and the whole collection of Dragon-X 1.2 descriptors (including 3D). We would like to emphasize that the overall quality of QSAR models presented in ref 62 is still high despite the presence of duplicates. Nevertheless, in our opinion, their presence significantly biased almost all steps of modeling, from model building to the selection of best models. After the beginning of our collaborative study, the Ames data set was revised by Dr. Hansen, and only 6512 compounds remained in their study.<sup>63</sup>

- (2) In our study, the remaining 6572 compounds were checked for the presence of inorganic compounds using the ISIDA Software. Thirty inorganic compounds such as ammonia and phosphoric acid were excluded.

- (3) The remaining 6542 structures were cleaned by the ChemAxon Standardizer (addition of explicit hydrogens, benzene ring aromatization, and nitro and carboxylic group standardization) and the HiT QSAR Software (nitro group standardization and connectivity checking).
- (4) The last step before the modeling stage was the repetition of search for duplicates and careful manual checking. This step was obviously time-consuming but necessary, because some erroneous structures (lacking hydrogens, having different tautomeric forms or representations of nitro groups, etc.) may become duplicates after structural cleaning. At the end no additional molecule was removed and 6542 compounds still remained for our international QSAR modeling exercise (see Table 3).

The detailed description of the study and its results obtained on the curated data set will be described in a separate publication because, again, this paper is focusing on the issue of data curation. However, we should mention that in total a group of collaborators has developed as many as 32 predictive QSAR models using different combinations of chemical descriptors and machine learning approaches. It is also worth noting that the results were initially reported by the Hansen group for noncurated data sets<sup>62</sup> and the later modeling of the curated data<sup>63</sup> showed that the predictivity of models developed on not-curated data sets was indeed somewhat overestimated (see Table 2) because of the presence of structural duplicates. Unlike small data sets, we should also emphasize that for such a large data set, the difference in prediction performances of models built before and after curation is statistically significant even when the difference in prediction accuracy is as low as 2%. Moreover, it is clear from Table 2 that models became more balanced (difference between specificity and sensitivity decreased from 8–12% to 2% only). This example illustrates that the use of data curation leads to more predictive and balanced models, along with more objective estimate of their true predictive power.

In addition to statistical aspects of models resulting from the analysis of curated vs noncurated data sets, this study also provides another example of investigation (after the DILI study<sup>50</sup>), where developed QSAR models were successfully used for the correction of erroneously annotated compounds in the data set. A compound was considered suspicious and selected for deeper experimental checking if (1) at least 30 out of 32 models obtained by the different teams failed to predict it accurately (either in modeling or the external validation sets) or (2) two or more structurally similar compounds had different annotations. In total, we have identified 86 suspicious compounds in the external set (51 nonmutagens and 35 mutagens) and 54 compounds in the modeling set (39 nonmutagens and 15 mutagens). Using both manual and automatic literature mining tools, our analysis revealed that 31 compounds (16 from the external set and 15 from the modeling set) were erroneously annotated in the original data set since we have indeed found published evidence that were in agreement with the model predictions for these compounds. Among them, 29 were originally annotated as nonmutagens, predicted as mutagens and were confirmed by at least one publication to be real Ames mutagens. On the other hand, only two mutagens predicted as nonmutagens were confirmed as nonmutagens. These results are in agreement with earlier observations<sup>63</sup> suggest-

ing that the experimental error rate for determining compounds as nonmutagenic is higher than that for mutagens (because compounds tested as negatives in the Ames test in certain bacterial strains may turn out to cause reverse mutations when examined in additional strains). Once again, the identification of 31 mislabeled compounds because of discordance between their stated and predicted properties suggests, perhaps unexpectedly, that predictive QSAR models obtained on carefully curated data sets can be successfully used for experimental biological data curation. Another important impact of experimental data curation applied to these compounds is that, despite an insignificant change in the overall prediction accuracy (see Table 2) for the whole external set (<1%) and the external cross-validation on the modeling set (<0.5%), there is a major improvement in prediction performances for these mislabeled compounds: for the external set, 19% (16 out of 86) of these mislabeled compounds have been predicted correctly (compared to zero percent with the original labels) and 28% (15 out of 54) for the modeling set. Data curation is thus an important criterion for QSAR model improvement in terms of prediction performances and reliability.

**3.5. Bioavailability Competition.** One recent illustration of the famous proverb “*the road to hell is paved with good intentions*” was the recent “QSARworld Modeling Challenge 2008” organized by QSARWorld (<http://www.qsarworld.com>.) Like other similar challenges, its overall objective was to benchmark modeling techniques from different international teams according to their “blind” prediction performances. The participants were asked to build QSAR models to predict human oral bioavailability using a given training set of compounds, for which both chemical structures and experimental biological data were made available. The best model was supposed to be selected based on the prediction accuracy (RMSE) for an external preselected (blind) test set, for which only chemical structures were made available.

Our group (and we believe many others) welcomed this challenge as an excellent opportunity to share the knowledge and experience between QSAR specialists throughout the world. We fully expected to participate in the challenge that was bound to enable the building of collective QSAR wisdom. Unfortunately, serious concerns regarding the modeling set, as well as the method of performance evaluation of QSAR models, were rapidly identified. In fact, the concerns were so strong that we concluded that the entire exercise was pointless and withdrew from the participation.

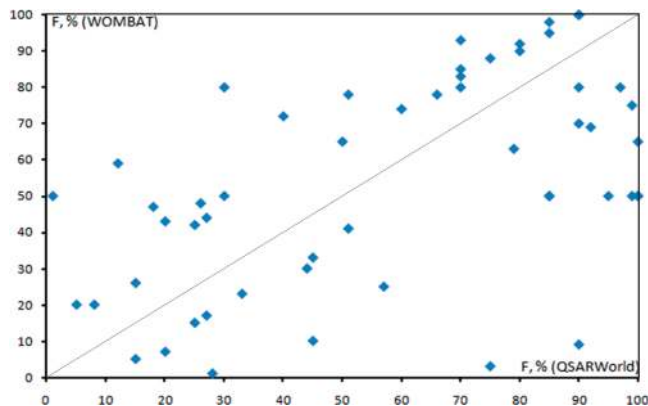
Our decision was based on the following specific reasons. The oral bioavailability data for all test set compounds were supposed to be completely unavailable so that the independent modeling effort could not be compromised in any way (as was indeed the case for another recent Solubility Challenge<sup>64,65</sup>). Nevertheless, these “external” data were in fact publicly available through the link placed at the QSARWorld Challenge Web site ([http://modem.ucsd.edu/adme/databases/databases\\_extend.htm](http://modem.ucsd.edu/adme/databases/databases_extend.htm)) at the same time when the competition was announced. In fact, the entire data set (i.e., both training and test sets) was compiled and kindly provided to us upon request by Dr. Tingjun Hou (Dept. of Chemistry and Biochemistry, University of California in San Diego). In addition, the SDF files provided by QSARWorld contained compound IDs from the Dr. Hou’s database.<sup>66</sup> As a result the blind competitive aspect of the Challenge was

obviously compromised because all experimental values for the external data set were known.

Moreover, it appeared that the quality of the data was low despite the absence of obvious errors in the structure representations. There were only few accidental mistakes, for example, lack of aromaticity in trapidil or a single (C–O) instead of a double (C=O) bond in ticarcillin. Most likely, some curation procedures were applied by creators of this data set which is appropriate before launching this kind of challenge. However, the search for duplicates using the HiT QSAR software<sup>41,42</sup> revealed 11 pairs of similar structures. Three of them were different in terms of stereochemical configuration (quinidine and quinine, betamethasone and dexamethasone, and levofloxacin and ofloxacin), and nevertheless, different isomers were reported to be associated with identical biological data. The structures in the remaining eight pairs were completely identical. The only difference was in the chemical names of duplicates, for example, dronabinol and tetrahydrocannabinol or metaprotenerol and orciprenaline. Thus, while the search for duplicates on the basis of names or CAS numbers can be done very rapidly, it is an inefficient tool to discover all chemically identical pairs of compounds. Furthermore, the experimental bioavailability values for similar compounds found in these four pairs differed by 8–43%. Such discrepancies are common for data sets compiled from the literature, when experimental values of investigated property are usually averaged. Actually, as we suggested above, activity values with small variations from one source to another could be averaged, but not in the cases such as quetiapine with bioavailability equal to 90% as reported in one source<sup>66</sup> and 9% as reported elsewhere.<sup>7</sup> Most likely, this difference is caused by simple human error, and “9” was intended to be reported as “90”. In this case, the removal of a compound with large deviation between experimental values reported in different sources from the training set is highly appropriate. It is also desirable to create a special test set containing such suspicious compounds to help reveal the true value by the means of consensus predictions using QSAR models built on “clean” data sets (cf. our description of both DILI and Ames studies above demonstrating the power of rigorously built QSAR models for correcting false biological data).

In case of any doubt concerning the consistency of the data, the search for analogous information in available sources is also a good solution. Specifically, it was found that some experimental data on human oral bioavailability provided by the Challenge organizers were not always measured on humans. Moreover, the simple comparison of the bioavailability data provided by the Challenge project with those found in WOMBAT-PK<sup>7</sup> for a large number of identical compounds revealed a large discrepancy between the two sources for many compounds. Thus, among 220 identical compounds found in both the Challenge and Wombat data sets, 54 compounds showed significant ( $\geq 10\%$ ) differences:  $\geq 20\%$  for 25 compounds;  $\geq 30\%$  for 15 compounds, and  $\geq 50\%$  for 5 compounds (Figure 9). Thus, after the application of the curation procedures to the original data set, consisting of 805 compounds, only 734 structures remained (see Table 3).

Obviously, we strongly support the idea of organizing QSAR Challenges in general. However, each Challenge should be designed very carefully. We believe that the



**Figure 9.** Experimental bioavailability values (%) from QSAR-World competition (X-axis) versus WOMBAT (Y-axis) for 55 overlapping compounds.

following simple recommendations may be useful when organizing competitive QSAR studies:

- (1) Biological data for the test set should be truly unavailable until the completion of the competition (a good example is the recent 2009 CADASTER Toxicity Challenge<sup>13</sup>).
- (2) Chemical structures should be thoroughly curated, for example, using procedures outlined in this paper.
- (3) Biological data should be of high quality, curated and consistent.

Results of this challenge, which are already posted at the QSARWorld Web site (<http://www.qsarworld.com/modleingcompetition08results1.php>), also demonstrate the importance of data curation. The best-model is characterized by RMSE of 30%, which is completely unacceptable given that the entire range of data values was 0–100%. Moreover, no additional statistical characteristics of the model quality were reported. In our opinion, the use of RMSE values only to estimate any model performance is insufficient; we always also report  $R_{\text{test}}^2$  (the coefficient of determination is more informative than RMSE in revealing whether or not predicted values reproduce quantitatively the experimental ones) and expect its value to exceed 0.6 to claim that the model is acceptable (as we mentioned the  $R_{\text{test}}^2$  value was not reported but it is highly unlikely that it exceeds 0.6 given the very high RMSE value). This opinion was also validated by the citation of Dr D. Krstajic, the winner of the bioavailability challenge, posted on the same Web site: “*Unfortunately, for the supplied data set, we were not able to find any good model.*” This case study reconfirms that robust and predictive QSAR models cannot be obtained using low-quality data.

#### 4. CONCLUSIONS

Molecular modelers and cheminformaticians typically analyze experimental data generated by other scientists. Consequently, when it comes to data accuracy, cheminformaticians are always at the mercy of data providers who may inadvertently publish (partially) erroneous data. Thus, dataset curation is crucial for any cheminformatics analysis such as similarity searching, clustering, QSAR modeling, virtual screening, etc, especially nowadays when the availability of chemical datasets in public domain has skyrocketed in recent years. Despite the obvious importance of this preliminary step in the computational analysis of any dataset, there appears to be no commonly accepted guidance or set of procedures for chemical data curation. The main objective of this paper was to emphasize the need for a standardized

chemical data curation strategy that should be followed at the onset of any molecular modeling investigation.

In this study, the most important steps of data set cleaning have been described including the removal of inorganics, organometallics, counterions, and mixtures; structural cleaning, ring aromatization, normalization of specific chemotypes, curation of tautomeric forms; deletion of duplicates; and manual checking of the structures and biological activities. Some general rules following the discussed applications were also formulated:

- (1) It is risky to calculate chemical descriptors directly from SMILES. It is preferable to compute descriptors (integral, fragments, etc.) from curated 2D (or 3D if necessary) chemical structures, where all chemotypes are strictly normalized.
- (2) Structural comparison across available databases may facilitate the detection of incorrect structures.
- (3) Even small differences in functional group representations could cause significant errors in models.
- (4) Searching for structure-based duplicates and their removal is one of the mandatory steps in QSAR analysis. Such searches based on chemical name or CAS number only are both incomplete and inefficient.
- (5) Because of the large number of experimental data sources, the search for additional information about investigated property in all available sources is desired to extract valuable knowledge and to compare the data to detect activity cliffs and identify diverse sources of errors.
- (6) Nothing can replace hands-on participation in the process because some errors obvious to a human are still not obvious for computers. After finishing all the mentioned steps, structures and activities should be checked manually once again.

Table 1 summarizes all proposed procedures (cf., also Figure 1) and provides a list of available software for every step of the curation process. This list is not exhaustive, and we invite readers to enrich it via a special link on our Web site (<http://chembench.mml.unc.edu/>) by adding references and links to other software that could be of interest in the context of data curation.

To emphasize the importance of data curation as a mandatory step in data analysis, we have discussed several case studies where chemical curation of the original “raw” database enabled the successful modeling study (specifically, QSAR analysis) or resulted in a significant improvement of model’s prediction accuracy. We also demonstrated that in some cases rigorously developed QSAR models could be even used to correct erroneous biological data associated with chemical compounds. We consider our paper complementary to the study by Young et al.<sup>8</sup> that provided several examples to illustrate how poor data quality could have detrimental influence on QSAR models.

In conclusion, we believe that there is a clear need for an additional principle that should be added to the five OECD principles for QSAR model development and validation,<sup>16,20</sup> and this principle should address the need for data curation before the model development is initiated. We suggest that this additional principle could be formulated as follows: “To ensure the consideration of (Q)SAR models for regulatory purposes, the models must be trained and validated on chemical data sets that have been thoroughly curated with respect to both chemical structure and associated target



property values.” We believe that good practices for curation of chemical records outlined in this paper will be of value to all scientists working in the fields of molecular modeling, cheminformatics, and QSAR modeling.

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