Bayesian Genotoxicity

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Abstract

This paper reports an initial exploration relating the genotoxicity of chemicals tested in vivo to the genotoxicity of chemical structures (sets of chemicals with common components). The analysis uses genotoxic results on 568 chemicals to make inference about the genotoxicity of 2893 chemical structures. As a result, there is substantial residual uncertainty. Two simple models are proposed to relate the genotoxicity of a chemical to the genotoxicity of its component chemical structures. Using these models, we find 19 chemical structures that are among the (predicted) most toxic 15 under at least one of the two models. Of these, 11 are already known to be genotoxic; we suggest further in vivo study of the remaining 8. The method used in the paper is Bayesian Markov Chain Monte Carlo. Convergence was rapid for both models. When applied predictively to chemicals, the results are disappointing; nearly every chemical is predicted to be highly toxic. To remedy this, we propose that further progress along this line would depend on the use of biochemically more sophisticated models that focus only on those chemical structures that are bioactive.

Keywords: Bayesian Analysis; Chemical Structures; Markov Chain Monte Carlo; Toxicity

Introduction

Genotoxicity

During cell division in mammals and other animals, DNA packaged into chromosomes divide as two daughter cells are created. If a chemical disturbs this process, chromosomal damage can result. Such damage can lead to cancer, birth defects, fetal deaths and infertility. There are standard methods to test a chemical to see if it induces chromosomal damage [1]. However, such tests are expensive and time-consuming, which has led to interest in using in silico models to predict genotoxicity. The purpose of this paper is to examine the extent to which genotoxicity results can be associated to chemical substructures (i.e. structural features), sets of chemicals that share parts of chemicals with DNA and that might react similarly. To this end, 568 chemicals have been tested for their effects on DNA, and are characterized by 2893 structural features.

Data sources

There were two sets of structural feature sets used to annotate the chemicals, the default set of features and the genetox alerts. The reason there is redundancy in these is the same structural features can occur in the different feature sets. This was done to be comprehensive in annotating the chemical substructures. The in vivo mouse micronucleus results for the chemicals in the Tox21 10K library were derived from the in silico first tool (https://www.leadscope.com/isfcui/app#). An sdf file provided by the US Environmental Protection Agency through their Tox21 web portal (http://www.epa.gov/comptox/dsstox/sdf_tox21s.html) was loaded into the insilico first application and the in vivo mouse micronucleus activity annotations, derived from the Lead scope SAR Genotoxicity Database (http://www.leadscope.com/toxicity_databases/), were mapped to the chemicals. In total 575 of the chemicals in Tox21 library were annotated with in vivo mouse micronucleus (139 positives and 436 negatives).

Chemical structures were retrieved from the Environmental Protection Agency’s (EPA) annotation of the Tox21 10K library (http://www.epa.gov/comptox/dsstox/sdf_tox21s.html). The SDF file provided by the EPA was loaded into the Lead scope software (Version 3.1; Columbus, OH). For the 575 chemicals with micronucleus data a total of 2893 structural features representing medicinal chemistry building blocks were generated. Of the 575 chemicals with micronucleus results 568 contained at least one structural feature. The remaining 7 without a single structural feature were removed from the data set and not considered further. Of the 2893 structural features 2797 are from the default Lead scope structural feature set (identifier F2 -27070). An additional 96 structural features (F27143-F27287) related specifically to DNA damage were also included in order to ensure comprehensive coverage of the structural features most relevant to the biological endpoint under consideration. Some of the DNA damage specific structural features were redundant...
with those contained in the default set of structural features, however if the identical features occurred in both sets they were assigned different identifier numbers and were left in the data set, hence the same structural feature may be associated twice.

Methods

There are two kinds of data that enter the analysis. The primary data specify, for each chemical, whether it was found to be genotoxic. The secondary data specify, for each chemical \(i\), the structures it contains. The analysis then permits inferences about the genotoxicity of each chemical structure, taking both sources of information into account. Making inferences about the genotoxicity of 2893 chemical structures on the basis of the known genotoxicity of 568 chemicals inevitably involves substantial residual uncertainty. To make the link between the toxicity of a chemical and the toxicity of its constituent chemical structures requires a model. There are two models that are used for the purpose of this research. Let \(S\) denote the set of chemicals sharing chemical structure \(i(i=1,...,2893)\). Suppose chemical structure \(i\) has probability \(p_i\) of not being genotoxic. Then what is the probability \(q_i\) that chemical \(j\) is benign (i.e., not toxic)?

In model I, we suppose that

\[
q_j = \prod_{s \in S_j} p_i
\]

(1)

In words, a chemical is benign if each of the structures to which it belongs, acting independently, is benign. As a consequence, the probability that chemical \(j\) is genotoxic is

\[
1-q_j = 1- \prod_{s \in S_j} p_i.
\]

(2)

In model II, we suppose that the probability that chemical \(j\) is genotoxic is

\[
1-q_j = \max_{j \in S_j} (1-p_i),
\]

the largest of the probabilities that each constituent chemical structure is genotoxic. Then, under model II, the probability that the chemical structure \(j\) is benign is

\[
q_j = 1- \max_{j \in S_j} (1-p_i) = \min_{j \in S_j} p_i.
\]

(4)

It is likely that toxicologists have prior information about the suspected genotoxicity of at least some chemical structures. However, to elicit such information about each of the 2893 chemical structures in question would probably exhaust the patience of both toxicologists and statisticians. Consequently, as an initial computation, we take the prior distribution on \(p = (p_1, ..., p_{2893})\) to be independent and uniform on \((0, 1)\).

Results

Each model was run for 5000 iterations of a Metropolis-Hastings MCMC algorithm [2]. Index plots suggest immediate achievement of equilibrium, so we used all 5000 sampled probabilities. As a check, we ran a second version that differed from the first in the following way: if chemical structure \(j\)'s initial probability in the first run was \(q_j^{(1)}\), the second run started at \(1-q_j^{(1)}\) for each \(j\). Figures 1 & 2 plot the means against each other for these two runs. The results reinforce the conclusion that the MCMC's have reached equilibrium. On average, the 568 chemicals tested have about 30 chemical structures each. Consequently, applying Model I would predict that virtually every chemical is genotoxic. Model II is only slightly better in this predictive sense. Improving these models would require a much more detailed understanding of how chemical structures contribute to the genotoxicity of chemicals than we now have. Nonetheless, and despite the large standard errors in our estimates, we think there is value in identifying suspect chemical structures. That 11 of our 19 had already been found to be genotoxic is a kind of validation. The other eight suggest a possible route of further research. Table 1 records the fifteen most genotoxic chemical structures found under either of the two models. Lines 1-15 are the most toxic (least benign) according to Model I. Lines 1-9, 12, 13, 16-19 are the chemical structures most toxic according to Model II. None of the 19 listed chemical structures were included in chemicals tested as benign.

Table 1: Means and Standard Deviations of \(p\), probability of being benign, under each model, and number of genotoxic chemicals, for each of 19 least benign chemical structures.

| Chemicals                 | Structure ID | Model I Mean | Model I St. Dev. | Model II Mean | Model II St. Dev. |
|---------------------------|--------------|--------------|------------------|--------------|-------------------|
| Carboxydimide             | F6578        | 0.209        | 0.171            | 0.391        | 0.279             |
| Sulfamic acid ester       | F6612        | 0.227        | 0.187            | 0.969        | 0.298             |
| Arsenic                   | F5253        | 0.297        | 0.193            | 0.26         | 0.206             |
| Imidazole 2-methyl        | F13008       | 0.25         | 0.213            | 0.446        | 0.301             |
| Monohaloalkenes           | F27213       | 0.296        | 0.234            | 0.404        | 0.296             |
| Halogenated methanes      | F27151       | 0.296        | 0.23             | 0.429        | 0.293             |
| Methane, 1, 1-dihydroxy   | F25966       | 0.307        | 0.235            | 0.436        | 0.295             |
| Propane, 1-carboxyloxy-3-hydroxy- | F26985     | 0.392        | 0.257            | 0.963        | 0.3               |

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The standard deviations reported here are calculated as if the MCMC draws are independent. In view of the likely positive serial dependence among them, these calculated standard deviations should be interpreted as lower bounds. Nonetheless, they give a reasonable indication of the uncertainty inherent in the calculated means. While it was not feasible to elicit priors on all 2893 chemical structures being studied here, we report below the literature on the 19 we find most suspect, as listed in Table 1. In doing so, we omit references based on the data we are studying, to avoid double-counting. Eleven of the 19 chemical structures listed in Table 1 are mentioned in the literature as known or suspected genotoxins:

1. Carbodimide (F6578) [3]
2. Arsenic (F5253) [4]
3. Monohaloalkenes (F27213) [5]
4. Halogenated methanes (F27151) [6]
5. Nitrate (F6586) 0.353 0.294 0.926 0.292 1
6. Benzene, 1-alkenyl-3-methoxy (F1930) 0.376 0.273 0.475 0.298 2
7. Benzene, 1-amino(NH2)-9-chloro- (F3891) 0.392 0.277 0.965 0.299 3
8. Alkyl nitrate (F27228) 0.399 0.278 0.961 0.297 1
9. Methane, 1,1-dicarbonyl- (F25859) 0.9 0.268 0.957 0.298 1
10. Pyridine, 2-heteroamino- (F17826) 0.41 0.282 0.979 0.3 2
11. Benzene, 1-amino(NH2)-, 4-chloro- (F3891) 0.98 0.288 0.958 0.298 3
12. Alkyl nitrite (F27228) 0.936 0.281 0.473 0.295 1
13. Carboximido(H2), alkenylacyc- (F5715) 0.972 0.289 0.46 0.295 1
14. Carboximido(H2), alkenyl (F5714) 0.977 0.287 0.46 0.299 3
15. Nitrosamine, including imidacloprid (F27215) 0.493 0.287 0.463 0.3 3
16. Nitrosamine, excluding imidacloprid "alkyl and aryl N-nitroso groups" (F6591) in "a well-established class of chemical carcinogens [10]
17. Sulfonate, 0-alkyl- "alkyl (C< 5) or benzyl ester of sulphonic or phosphonic acid," (F7091)

Our attitude toward the eight chemical structures on our list for which we have not found supporting literature (i.e., lines 2, 4, 7-10, 13 and 14 in Table 1) is as follows: they may be spurious, or they may be genotoxic, and we don’t know which. We would point out, however, that several of them are included in only one
tested chemical, so this finding should be treated with caution. However, there is benefit in reducing the list of suspected chemical structures from the 2893 we started with, to eight. Thus, we propose that chemicals with one or more of these eight structures would be good candidates for further testing. It is also useful to get an overview of the relationship between the results for the two models. Figure 3 displays the means under Models I and II for each chemical structure, plotted against each other.

**Figure 3**: Relationship between the two models: Model I (Product) vs. Model II (Min), reporting the mean probability of chemical structures being benign.

**Conclusion & Discussion**

Strength of this analysis is that we used a standardized endpoint for evaluating genotoxicity, the micronucleus assay. The major limitation of this analysis is the poor predictivity of both models, as nearly all chemicals were predicted to be toxic. This is due to the structure of the models. A typical chemical has about 30 chemical structures. The product of 30 probabilities, as in Model I, leads to a small probability of a chemical being benign. Model II is not quite as severe, but the minimum of 30 probabilities is also likely to be small. Consequently, both models are too pessimistic in predicting the genotoxicity of, as yet untested, chemicals. The issue here is not Bayesian analysis, nor Markov chain Monte Carlo, but rather that Models I and II do not reflect enough toxicological wisdom to be useful predictively. This is not surprising given work by others that shows poor correlation activity relationship rules for predicting carcinogenic potential of unsaturated carbonyl compounds as criteria for genotoxicity and mutagenicity. Environ Health Perspect 88: 99-106.

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