**Using machine learning to determine the composition of artifact residues**

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**Abstract** Determining composition of artifact residues is a central problem in metabolomics. Traditionally this is done by comparing using the shared number of mass spectra. While this method is simple and straightforward, its predictions might sometimes be quite poor. In this article, we introduce a novel approach based on ideas from the field of natural language processing to solve this problem. We test our strategy on the [data set set]. The results indicate that our algorithms perform slightly better than the traditional method. We also that our algorithms are automated which makes it easy to reuse and apply to other datasets.

**Keywords:** ancient residue metabolomics, archaeometry, artifact, mass spectral feature, smoking pipe

**Introduction**

Metabolomics is the systematic quantitative and qualitative study of small molecules (or mass spectral features) in biological systems. Brownstein *et al.* (2020) expanded upon this field with their ancient residue metabolomics-based method. Albeit the mass spectral features in this study were not derived from biological systems, they were residues left behind from biological processes; i.e., originating from plants including several *Nicotiana* species that were smoked by indigenous peoples. Before the Brownstein *et al.* (2020) study, ancient residue analysis relied on the biomarker approached. However, the biomarker approach failed to distinguish between related species, leaving open questions about the relationship between plants and people. For ancient residue metabolomics, all compounds are of interest improving the resolution of determining which plants species had been smoked in a particular pipe (Brownstein *et al.*, 2020).

In short, data from hyphenated chromatography instruments (such as gas chromatography- and liquid chromatography-mass spectrometer) are processed and aligned in MZmine 2 (Pluskal *et al.*, 2010), Progenesis QI (Waters Corporation, Milford, MA, USA), or another “omics” software. Afterwards, these data are exported from the software and then processed manually as described in the following passage from Brownstein *et al.* (2020): “[T]he dataset was exported into Microsoft Excel and mass spectral features shared with the blank [samples] were removed from the analysis. The [three solvent] extracts from each sample were combined into a single compound list and compounds with no abundance values were removed. The [ancient samples] were then compared to the [experimental samples using a Venn diagram].” This final component of processing and analyzing the datasets determines which plant species may have been used in an ancient artifact. Because this step requires a manual process, it can introduce errors and is time consuming. Various metabolomics platforms exist including MetaboAnalyst 4.0 (Pang *et al.*, 2021) or XCMS Online (Tautenhahn *et al.*, 2012); however, these platforms are limited in their ability to process datasets from ancient residue studies. Therefore, we introduce a novel, automated method for determining the composition of organic residues in modern smoking pipes. Our approach is inspired by techniques and ideas from the field of Natural Language Processing (we refer readers to the next section for technical discussions of this approach). Additionally, we implement our approach using standard libraries in Python such as Sklearn and Pandas.

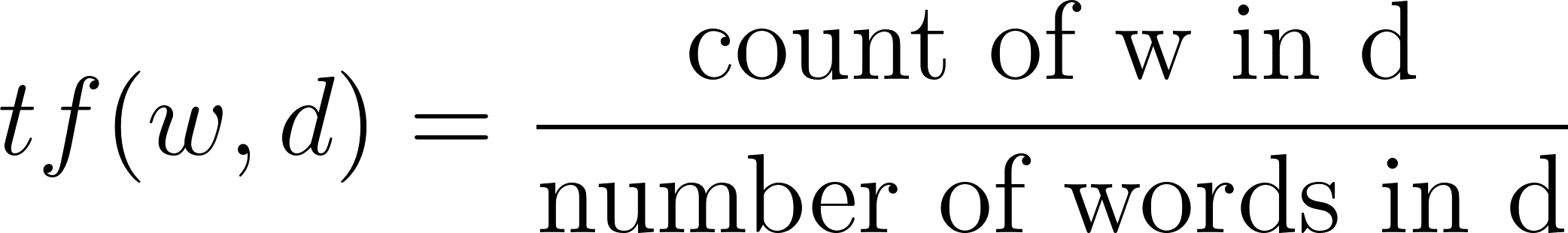
**Materials and Methods**

A novel functionality of our approach is to introduce an entirely new method to compare the mass spectra feature similarities between experimental and ancient artifacts (i.e., sample groups). Our approach is inspired by advances in Natural Language Processing (NLP). Here we use the following analogy:

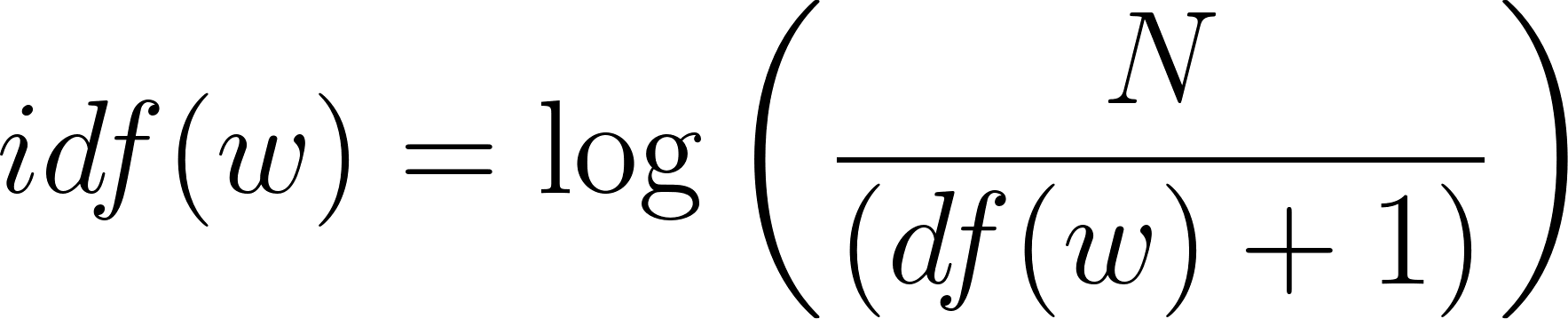
Words 🡨🡪 Mass Spectra Features

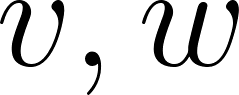
Documents 🡨🡪 Sample Groups

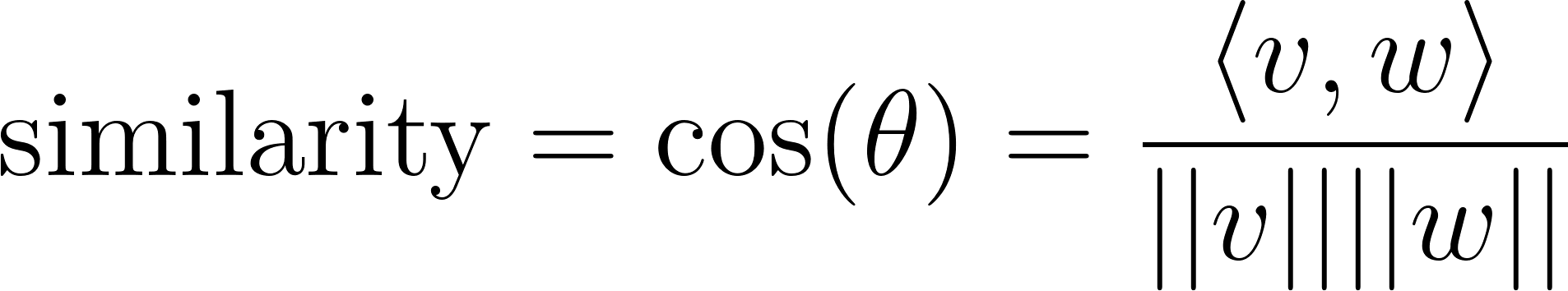
The standard technique in NLP is to first transform the original data into the term frequency- inverse document frequency (TF-IDF) matrix. This transformation helps to resolve the fact that some substances appear more often than others. More precisely, the importance of a term is not solely determined by its frequency in a text (TF) but also how rare this term is in other texts in the corpus (IDF). Let us recall these terminologies mathematically. Term frequency refers to the frequency of a word in a particular document:

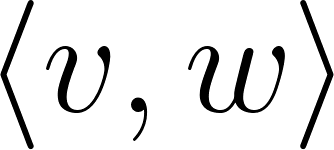
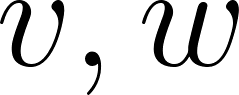
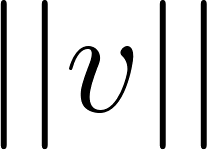
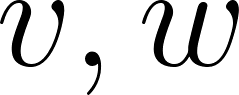
[](https://www.codecogs.com/eqnedit.php?latex=tf(w%2Cd)%3D%5Cfrac%7B%5Ctext%7Bcount%20of%20w%20in%20d%7D%7D%7B%5Ctext%7Bnumber%20of%20words%20in%20d%7D%7D#0)

The inverse of the document frequency which measures the informativeness/prevalence of term t

[](https://latex-staging.easygenerator.com/eqneditor/editor.php?latex=idf(w)%3D%5Clog%E2%81%A1%5Cleft(%5Cfrac%7BN%7D%7B(df(w)%2B1)%7D%20%5Cright)#0),

where N is the number of documents and df(w) is the number of documents containing w. IDF score depends on the occurrence of terms and not on their numerical frequencies. Once the TF-IDF is computed, we can then use cosine similarity to compare two different groups. Recall that for two vectors [](https://www.codecogs.com/eqnedit.php?latex=v%2Cw#0) their cosine similarity is defined to be cosine of the angle [](https://www.codecogs.com/eqnedit.php?latex=%5Ctheta#0) between them, namely

[](https://www.codecogs.com/eqnedit.php?latex=%5Ctext%7Bsimilarity%7D%3D%5Ccos(%5Ctheta)%3D%5Cfrac%7B%20%5Clangle%20v%2C%20w%20%5Crangle%7D%7B%7C%7Cv%7C%7C%20%7C%7Cw%7C%7C%7D#0)

Here, [](https://www.codecogs.com/eqnedit.php?latex=%5Clangle%20v%2C%20w%20%5Crangle#0) is the inner product of [](https://www.codecogs.com/eqnedit.php?latex=v%2C%20w#0) and [](https://www.codecogs.com/eqnedit.php?latex=%7C%7Cv%7C%7C#0) ,[](https://www.codecogs.com/eqnedit.php?latex=%7C%7Cw%7C%7C#0) is the Euclidean norm of [](https://www.codecogs.com/eqnedit.php?latex=v%2C%20w#0). We note similarity score ranges from -1 meaning exactly opposite to 1 meaning exactly the same, with 0 indicating orthogonality, while in-between values indicate intermediate similarity or dissimilarity.

*Artemisia ludoviciana* Nutt. (*Alu*) leaves, *Arctostaphylos uva-ursi* (L.) Spreng. (*Auv*) leaves, *Cornus sericea* L. (*Cse*) bark, *Gaultheria shallon* Pursh (*Gsh*) leaves, *Lobelia inflata* L. (*Lin*) leaves, *Nicotiana attenuata* Torr. ex S. Watson (*Nat*) leaves, *Nicotiana glauca* Graham (*Ngl*) leaves, *Nicotiana obtusifolia* M. Martens & Galeotti (*Nob*) leaves, *Nicotiana quadrivalvis* Pursh (*Nqu*) leaves, *Nicotiana rustica* L. (*Nru*) leaves, *Nicotiana tabacum* L. (*Nta*) leaves, *Rhus glabra* L. (*Rgl*) autumn leaves, *Salvia sonomensis* Greene (*Sso*) leaves, *Taxus brevifolia* Nutt. (*Tbr*) needles, and *Verbascum thapsus* L. (*Vth*) leaves were collected, freeze-dried for 3 days, and crushed for experimental smoking. American Spirit (AmSp) tobacco (Santa Fe Natural Tobacco Company, Oxford, NC, USA) was purchased from a local grocery store in Pullman, Washington.

The plant materials and AmSp were smoked following the experimental conditions detailed in Brownstein *et al.* (2020). To limit biases, the authors did not know which plant species had been smoked in which clay pipe. After the samples were analyzed by liquid chromatography-mass spectrometry and processed in MZmine 2 (Pluskal *et al.*, 2010) following the parameters described in Brownstein *et al.* (2020), the data were exported into .csv files. Python libraries were then used to apply the TF-IDF computation scores to these datasets.

**Results and Discussion**

We used Python to write the scripts because of the availability of several useful data analysis, machine learning, and deep learning libraries. All the scripts and datasets are available on GitHub: <https://github.com/tungprime/NLP_and_composition_of_artifact_residues>. Our script automates the classical method described in Brownstein *et al.* (2020), as well as utilizes recent advances in machine and deep learning to better predict which plant species had been smoked in a particular artifact (new method). As shown in Table 1, the new method predicts that CP1 was most likely smoked with *Nta* (0.0370). Table 2 summarizes the model predictions of the classical and new methods, and the key provides the expected results. In fact, CP1 was smoked with *Nta*. The new method is capable of accurately predicting the composition of artifact residues \_[can you do the math, I am not sure you want to use relative difference or absolute difference]\_\_\_\_% better than the classical method.

**Table 1.** Similarity scores of clay pipe 1 (CP1) smoked with an unknown plant sample. Sixteen (16) different experimental pipes smoked with only one of the plant species or AmSp was compared to CP1. The top five scores were only included in the table. *Nta* is the most likely candidate smoked in CP1.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | *Nta* | *Ngl* | *Nat* | AmSp | *Nob* |
| Similarity scores | 0.0370 | 0.0287 | 0.0232 | 0.0185 | 0.0170 |

*Nat*, *Nicotiana attenuata*; *Ngl*, *Nicotiana glauca*; *Nob*, *Nicotiana obtusifolia*; *Nta*, *Nicotiana tabacum*; and AmSp, American Spirit.

Contamination is a major concern for ancient residue metabolomics (Damitio *et al.*, 2021; Zimmermann *et al.*, 2021). For instance, residues from commercial tobacco smoke may contaminate the surface of artifacts at excavation sites or on display at a museum. Thus, we included AmSp in our study as a contaminate control. With the contaminate control (0.0185), we were still able to accurately determine the composition of CP1 (Table 1) and the other clay pipes. Utilizing contaminate controls will improve confidence in whether or not a particular artifact had been smoked with an endemic tobacco. Furthermore, our new method will enable researchers to confidently determine if the caffeine present in/on an artifact resulted from ancient cacao or holly brewing practices instead of modern contaminates from caffeinated beverages such as coffee (King *et al.*, 2017; Washburn *et al.*, 2014).

It was also revealed that neither the new nor classical methods could accurately predict that CP8 had a mixture of *Nta* and *Auv*. However, the experimental pipes compared to CP8 had only been smoked with *one* plant species. It is possible that training the new method with experimental pipes smoked with complex mixtures may improve the likelihood of predicting pipes smoked with more than one plant material.

**Table 2.** Predicted plant species in each clay pipe (CP). Sixteen (16) different experimental pipes smoked with only one of the plant species or AmSp was compared to each CP. The expected results are the plant species listed under key.

|  |  |  |  |
| --- | --- | --- | --- |
| Clay pipe (CP) | Classical method (number of mass spectra features shared with the experimental pipe) | New method (similarity score) | Key |
| CP1 | *Auv* (3) | *Nta* ( | *Nta* |
| CP2 | *Nat* (6) and *Nqu* (6) | *Nqu* ( | *Nqu* |
| CP3 | *Nob* (10) | *Nob* ( | *Nob* |
| CP4 | *Nat* (4) | *Nta* ( | *Alu* |
| CP5 | *Ngl* (6) | *Ngl* ( | *Ngl* |
| CP6 | *Lin* (10) | *Ngl* ( | *Lin* |
| CP7 | *Auv* (13) | *Auv* ( | *Auv* |
| CP8 | *Auv* (9) | *Cse* ( | *Nta* mixed with *Auv* |

*Alu*, *Artemisia ludoviciana*; *Auv*, *Arctostaphylos uva-ursi*; *Cse*, *Cornus sericea*; *Lin*, *Lobelia inflata*; *Nat*, *Nicotiana attenuata*; *Ngl*, *Nicotiana glauca*; *Nob*, *Nicotiana obtusifolia*; *Nqu*, *Nicotiana quadrivalvis*; and *Nta*, *Nicotiana tabacum*.

We see that while the old method can only predict 4/8 (50%) samples correctly, our new method is slightly better; it classifies 5/8 (62%) samples correctly.

**Conclusion**

Machine and deep learning have been vital tools for solving problems in biology where traditional methods seem inadequate or are time-consuming. Combining conventional and machine learning-based methods to process and analyze data helps researchers gain a more in-depth analysis of their data. The new method was used to predict which plant species had been smoked in modern clay pipes, and we believe this method can be applied to ancient smoking pipes, brewing vessels, and other artifacts.

**Data availability statement:** All scripts and datasets used in this study are in the GitHub repository: <https://github.com/tungprime/NLP_and_composition_of_artifact_residues>

**Author contributions:** T.T.N. and K.J.B. designed the project. T.T.N. developed and wrote the scripts. T.T.N. and K.J.B. wrote the paper. All authors approved of the final version of the paper.

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