


# Scikit-learn

- <http://scikit-learn.org>
- <http://www.innoarchitech.com/machine-learning-an-in-depth-non-technical-guide/>

 **Fix Me!** : ajouter des exemples avec analyse de textes contenant des termes scientifiques, des noms de substances chimiques,...

- [Scikit-chem](#), simple cheminformatics for Python
- [MolMiner](#), for extracting compounds from scientific literature
- [ChemDataExtractor](#), automatically extract chemical information from scientific documents. Cf. [ChemDataExtractor: A Toolkit for Automated Extraction of Chemical Information from the Scientific Literature](#) Matthew C. Swain and Jacqueline M. Cole J. Chem. Inf. Model., 2016, 56 (10), pp 1894–1904 DOI: 10.1021/acs.jcim.6b00207
  - voir aussi <https://jcheminf.springeropen.com/articles/10.1186/1758-2946-7-S1-S2>

## References

- <https://mubaris.com/2017-10-01/kmeans-clustering-in-python>
- <https://www.codementor.io/garethdwyer/introduction-to-machine-learning-with-python-s-scikit-learn-czha398p1>

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