Undergraduate Research Posters

Structural Insights into Metal-Organic Connectivity by Paramagnetic NMR

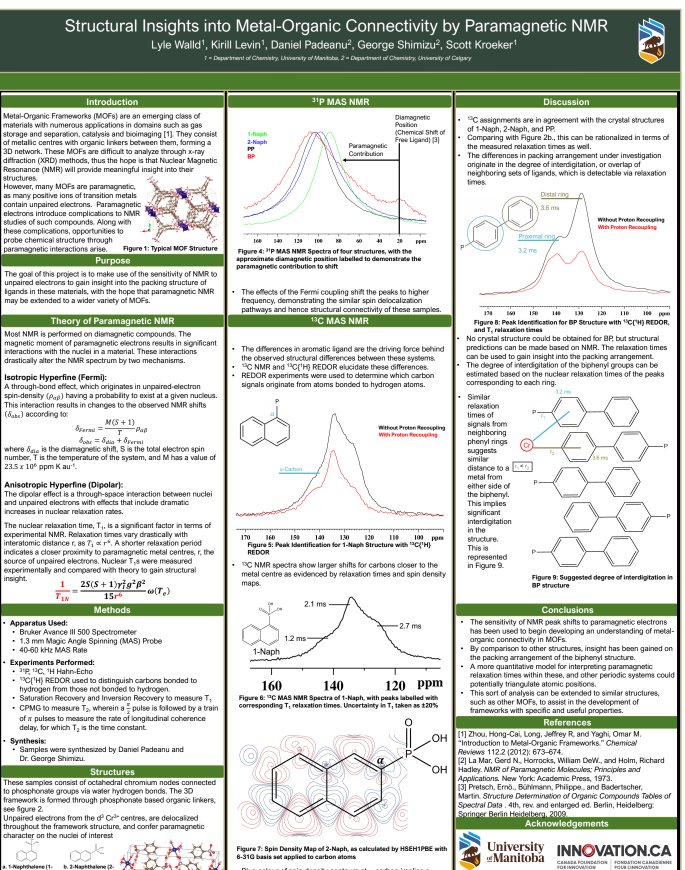
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Abstract

Metal-Organic Frameworks (MOFs) are an emerging class of materials with numerous applications in domains such as gas storage and separation, catalysis and bioimaging [1]. They consist of metallic centres with organic linkers between them, forming a 3D network. These MOFs are difficult to analyze through x-ray diffraction (XRD) methods, thus the hope is that Nuclear Magnetic Resonance (NMR) will provide meaningful insight into their structures. However, many MOFs are paramagnetic, as many positive ions of transition metals contain unpaired electrons. Paramagnetic electrons introduce complications to NMR studies of such compounds. Along with these complications, opportunities to probe chemical structure through paramagnetic interactions arise.





Blue colour of spin density contours at  $\alpha$  carbon implies a positive Fermi shift, agreeing with the findings of <sup>13</sup>C(<sup>1</sup>H} REDOR experiments, where the  $\alpha$  carbon is shifted to higher frequencies.

Figure 3: Phenyl based structu (PP), as determined by singlecrystal X-ray diffraction

Biphenyl (BP) Ligand d. Phenyl (PP) Ligand Figure 2: Ligands present in the

uctures analyzed

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