Difference Hirshfeld fingerprint plots: A tool for studying polymorphs

Damien J. Carter,a Paolo Raiteri,a Keith R. Barnard,b Rhian Gielink,c Mauro Mocerino,c Brian W. Skelton,d Jamila G. Vaughanc, Mark I. Ogden,c,d and Andrew L. Rohlb

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A new tool has been developed to help elucidate the differences in packing between different polymorphs, especially when the differences of interest are small. The technique builds upon the Hirshfeld fingerprint plot pioneered by Spackman and co-workers by subtracting the value at every point in a technique fingerprint plot from the value at every point in another. This is found to reveal differences that are not readily apparent to the eye. By summing the absolute values of these differences, a quantitative measure of the difference between two fingerprint plots can be obtained. The technique was applied to Ni and Cu trans-bis(2-hydroxy-5-methylphenonanoneoximato) complexes determined at two temperatures, with the Ni complex displaying temperature-dependent polymorphism. Difference Hirshfeld fingerprint plots were also generated for calculated structures from DFT simulations that were performed on the experimental structures. These demonstrated that the simulations reproduced the fine detail of the packing.

Introduction

Hirshfeld surfaces have become an invaluable tool for crystallographers and crystal engineers alike. Hirshfeld surfaces enclose molecules within crystals and are defined as the region where the contribution from the electron density within a molecule exceeds that from all other molecules in the crystal. For each point on the Hirshfeld surface, the distance to the nearest atom within the molecule, \( d_e \), can be calculated. Likewise the distance to the nearest atom on the neighbouring molecule, \( d_r \), can also be determined. Plotting these as a 2-D histogram produces the fingerprint plot.

McKinnon and coworkers demonstrated that fingerprint plots are a useful tool to study polymorphism by visually comparing the plots of different polymorphs and noting the differences. This type of visual comparison has been used in a number of studies since in systems ranging from simple molecules such as cis-1,4-cyclohexanediol, pyrazinamide and rubrene to a variety of pharmaceutically relevant compounds to ligands. We have previously used fingerprint plots to elucidate the differences between the five polymorphs of 1,8-dihydroxyanthraquinone, but in addition we also used them to compare experimental and calculated structures of each polymorph. In a similar vein, Schatschneider and Liang used fingerprint plots to monitor structural changes as they simulated compression of crystalline indole. Parkin and coworkers have used cluster analysis and multivariate statistics to calculate similarity indices between different structures and this work has been extended by Collins and coworkers. The latter also examined the effects of temperature on the fingerprint plots of anthracene and noted “The plots are very similar and represent a challenge to any data mining technique”. Latfosińska and coworkers also found small differences between the fingerprint plots of the polymorphs of 3,3′-diindolylmethane determined at several temperatures and defined several measures to quantify the differences.

When the plots are very similar, determining the differences can be a challenge for humans as well as machines. We have been thinking about this challenge for some time but the discovery of temperature-dependent polymorphism in some phenolic oximes pushed our ideas forward.

Phenolic oximes are important reagents in the solvent extraction of copper. The extractants are typically derived from salicylaldehydes or \( \alpha \)-ketophenols, and form square planar complexes stabilized by intramolecular hydrogen bonds (Scheme 1). “Folded” octahedral nickel complexes of these ligands have been reported when isolated in the presence of chelating diamines. The diamines also impact on the solvent extraction behaviour, forming a synergistic system with enhanced nickel extraction. We have previously reported synergistic solvent extraction systems based on aliphatic \( \alpha \)-hydroxyoximes and carboxylate ligands. While attempting to extend this work to phenolic oxime ligands, we crystallised the simple square planar nickel and copper complexes, and observed temperature-dependent polymorphism in the nickel complex.

Here we report the experimental determination of the low temperature Ni oxime structure and the low and high temperature Cu oxime structures using single crystal X-ray diffraction with accompanying DFT calculations investigating the stability of Cu and Ni substituted in the low and high temperature Ni and Cu oximes structures. These yielded a significant number of very similar Hirshfeld fingerprint plots, which prompted a detailed examination of these structures and development of an approach to probe subtle changes in crystal packing.

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Methodology

Ligand synthesis

2-Hydroxy-5-methylacetophenoneoxime HL was synthesised following literature methods. A mixture of 2-hydroxy-5-methylacetophenone (19.69 g, 0.131 mol) was combined with hydroxylamine hydrochloride (25.15 g, 0.36 mol), ethanol (220 mL) and pyridine (21 mL) and the reaction mixture gently heated at reflux for 2 h. The ethanol then was removed under reduced pressure and the residue dissolved in ether. The ether solution was washed with 1 M HCl to remove the pyridine. The organic layer was then evaporated leaving white solid oxime HL.

Crystallography

Crystallographic data for the structures were collected at 100(2) K on an Oxford Diffraction Xcalibur diffractometer fitted with Mo Kα radiation. Following analytical absorption corrections and solution by direct methods, the structure was refined against \( F^2 \) with full-matrix least-squares using the program SHExXL-97. All hydrogen atoms were added at calculated positions and refined by use of a riding model with isotropic displacement parameters based on those of the parent atoms. Anisotropic displacement parameters were employed for all the non-hydrogen atoms.

Crystal data and refinement details

NiL₂: T = 100 K. Empirical formula \( \text{C}_{16}\text{H}_{22}\text{NiO}_4 \); formula weight 391.90, triclinic, space group \( P\overline{1}; \alpha = 6.468(13), b = 7.7544(16), c = 8.6026(17) \); \( \alpha = 97.267(17), \beta = 109.253(18), \gamma = 96.531(17) \); \( V = 398.46(14) \); \( Z = 1, D_{\text{c}} = 1.613 \text{ Mg/m}^3, \mu = 1.244 \text{ mm}^{-1} \); crystal size 0.39 × 0.14 × 0.10 mm; \( \theta_{\text{min,max}} = 3.36, 37.22^\circ \), reflections collected = 7581; independent reflections \( = 3960 \ [\text{R}_{\text{int}} = 0.0295], \) data/restraints/parameters 3960/0/118, goodness-of-fit on \( F^2 = 0.941; \) final R indices \( [F>2\sigma(F)] R_{I} = 0.0401, wR_{I} = 0.0809; \) R indices (all data) \( R_{I} = 0.0572, wR_{I} = 0.0841 \). Largest diff. peak and hole = 0.890 and -0.761 e. Å\(^{-3} \).

CCDC = 1519180.

CuL₂: T = 100 K. Empirical formula \( \text{C}_{16}\text{H}_{22}\text{CuO}_7 \); formula weight 391.90, triclinic, space group \( P\overline{1}; \alpha = 6.2388(2), b = 8.1434(2), c = 8.6344(3) \); \( \alpha = 71.541(3), \beta = 79.600(3), \gamma = 85.781(2) \); \( V = 408.92(2) \); \( Z = 1, D_{\text{c}} = 1.591 \text{ Mg/m}^3, \mu = 1.362 \text{ mm}^{-1} \); crystal size 0.27 × 0.09 × 0.07 mm; \( \theta_{\text{min,max}} = 3.03, 40.86^\circ \), reflections collected = 19572; independent reflections \( = 5231 \ [\text{R}_{\text{int}} = 0.0294], \) data/restraints/parameters 5231/0/118, goodness-of-fit on \( F^2 = 0.1065; \) final R indices \( [F>2\sigma(F)] R_{I} = 0.0324, wR_{I} = 0.0841; \) R indices (all data) \( R_{I} = 0.0363, wR_{I} = 0.0865 \). Largest diff. peak and hole = 0.630 and -0.437 e. Å\(^{-3} \).

CCDC = 1519182.

Computations

DFT calculations were performed using the Quantum ESPRESSO code. The project-augmented wave (PAW) method was used with pseudopotentials adopted from the atompaw library. The planewave basis set had a kinetic energy cut-off of 80 Ry and the exchange-hole dipole moment (XDM) method was used in particular B86b-XDM, which combines B86b exchange with PBE correlation, with the XDM parameters \( A1 = 0.337, A2 = 2.488 \). Both Cu and Ni atoms substituted in the low and high temperature Ni and Cu oxime structures were examined. Calculations include spin polarisation and both atomic coordinates and lattice parameters were optimised using the BFGS quasi-newton method with a maximum tolerance of 0.001 Ry/Bohr.
100%. The fingerprint plot can then be viewed using the freely
distributed Gnuplot. A script plot_finger has been written that
uses a scale and colour scheme that exactly reproduces the
fingerprint plots generated by CrystalExplorer. A C program,
\texttt{diff_finger}, has been developed that takes two files generated by
the fingerprint codes and subtracts them. Finally, the script
\texttt{plot_diff_finger} uses Gnuplot to plot the output of \texttt{diff_finger}
using a palette that goes from blue (negative) to white (zero) to
red (positive). All of these programs and scripts are available
online.\textsuperscript{29}

**Results & Discussion**

![Scheme 1. General reaction scheme for the formation of a typical ML\textsubscript{2}
complex with 2-hydroxy-5-methylacetophenone oxime, HL.](Image)

Following on from our work with synergistic solvent extraction
systems based on aliphatic \(\alpha\)-hydroxyoximes,\textsuperscript{15-17} copper and
nickel complexes of ketoime HL were crystallised in the
presence of a range of potential co-extractants. The only system
that gave a modified coordination sphere resulted from the
reaction of HL with nickel nitrate where dimethylsulfoxide
molecules were found in the axial positions. In addition, however,
the ligand was nitrated in the ortho position under the reaction
conditions (see SI).\textsuperscript{29} In the course of this work, the CuL\textsubscript{2}, and
NiL\textsubscript{2} complexes were isolated and structurally characterised. A
room temperature structure of NiL\textsubscript{2} has been reported previously
\textsuperscript{30}, but here it was found that the system undergoes a reversible
structural transition at \(\approx 150 \text{ K}\) to give a polymorph with a
substantially changed unit cell (Table 1). The difference in the
structures of the two polymorphs lies in the degree of rotation of
the molecule. The angle between the Ni...Ni(x,y,1+z) vector and
the plane of the molecule increases from \(3.4^\circ\) at room
temperature to \(13.2^\circ\) at 100 K. This is shown in the two cell
projections down the \(c\) axes, Fig. 1a of the room temperature
structure and Fig. 1b for the structure at \(T = 100 \text{ K}\). In contrast,
the CuL\textsubscript{2} structure was found to be isomorphous with the room
temperature Ni structure at both 100 K and 150 K. In both
polymorphs, face-to-face interactions between the planar
molecules dominate. The observation that the nickel L complex
has a structural transition as temperature changes, and the
isomorphous copper complex does not, led us to undertake a
theoretical study to explain why this difference occurs.

**Table 1. Experimental lattice parameters for ML\textsubscript{2}
complexes.**

<table>
<thead>
<tr>
<th>Structure</th>
<th>(a) (Å)</th>
<th>(b) (Å)</th>
<th>(c) (Å)</th>
<th>(α(°))</th>
<th>(β(°))</th>
<th>(γ(°))</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuL\textsubscript{2} @ 100 K</td>
<td>6.243(4)</td>
<td>8.088(6)</td>
<td>8.6219(7)</td>
<td>71.24(1)</td>
<td>79.573(6)</td>
<td>85.968(6)</td>
</tr>
<tr>
<td>CuL\textsubscript{2} @ 150 K</td>
<td>6.238(2)</td>
<td>8.143(2)</td>
<td>8.644(3)</td>
<td>71.42(1)</td>
<td>79.60(3)</td>
<td>85.78(2)</td>
</tr>
<tr>
<td>NiL\textsubscript{2} @ 100 K</td>
<td>6.468(13)</td>
<td>7.7544(16)</td>
<td>8.602(17)</td>
<td>97.26(17)</td>
<td>105.253(18)</td>
<td>96.531(17)</td>
</tr>
<tr>
<td>NiL\textsubscript{2} @ RT</td>
<td>6.290(5)</td>
<td>8.204(7)</td>
<td>8.653(8)</td>
<td>71.92(6)</td>
<td>80.61(7)</td>
<td>83.54(6)</td>
</tr>
</tbody>
</table>

* The experimental Ni\textsubscript{L2} RT structure has been reported by Voutsas, \textit{et al.} \textsuperscript{30}

**Table 2. Calculated lattice parameters of the high and low temperature
CuL\textsubscript{2} and NiL\textsubscript{2} complexes.**

<table>
<thead>
<tr>
<th>Structure</th>
<th>(a) (Å)</th>
<th>(b) (Å)</th>
<th>(c) (Å)</th>
<th>(α(°))</th>
<th>(β(°))</th>
<th>(γ(°))</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuL\textsubscript{2} @ 100 K</td>
<td>6.24</td>
<td>8.05</td>
<td>8.61</td>
<td>70.85</td>
<td>78.53</td>
<td>86.35</td>
</tr>
<tr>
<td>CuL\textsubscript{2} @ 150 K</td>
<td>6.24</td>
<td>8.05</td>
<td>8.61</td>
<td>70.85</td>
<td>78.52</td>
<td>86.35</td>
</tr>
<tr>
<td>NiL\textsubscript{2} @ 100 K</td>
<td>6.21</td>
<td>8.00</td>
<td>8.58</td>
<td>71.99</td>
<td>79.75</td>
<td>85.61</td>
</tr>
<tr>
<td>NiL\textsubscript{2} @ RT</td>
<td>6.20</td>
<td>8.01</td>
<td>8.58</td>
<td>71.96</td>
<td>79.81</td>
<td>85.59</td>
</tr>
</tbody>
</table>

* The experimental Ni\textsubscript{L2} RT structure has been reported by Voutsas, \textit{et al.} \textsuperscript{30}

Overall, it is found that DFT calculations are able to successfully
reproduce the experimental crystal structures with good accuracy,
with all differences between the calculated and experimental
parameters for the low temperature structures \((\leq 150 \text{ K})\) under
1.5%. At room temperature, the calculated lattice constants are
significantly less than the experimental parameters, as expected.

The XDM method has been reported to perform extremely well
for calculations of molecular crystals, such as the C21 reference
set \textsuperscript{31,32} and a set of monosaccharide structures \textsuperscript{15}. Note that the
two calculated Cu structures starting from the two experimental
Cu structures and the room temperature Ni structure are

![Figure 1. Unit cell contents of the (a) room temperature determination
and (b) low temperature structure of NiL\textsubscript{2}, projected down the
crystallographic c axis. Hydrogen atoms have been omitted for clarity.](Image)
essentially the same (maximum cell length difference of 0.003 Å and cell angle difference of 0.117°). This is also true of the calculated Ni structures (maximum cell length difference of 0.005 Å and cell angle difference of 0.105°). This is expected as all three experimental structures are isomorphous and the DFT calculations are all at 0 K, which gives confidence that the Quantum ESPRESSO optimiser is performing well.

In Table 3, we report the relative total energies for each metal atom in the four L oxime structures.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Relative energy (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CuL₂ @ 100 K</td>
<td>0.37</td>
</tr>
<tr>
<td>CuL₂ @ 150 K</td>
<td>0.38</td>
</tr>
<tr>
<td>NiL₂ @ 100 K</td>
<td>0.00</td>
</tr>
<tr>
<td>NiL₂ @ RT</td>
<td>0.37</td>
</tr>
</tbody>
</table>

The results in Table 3 show that the low temperature Ni structure is the most stable structure for both Ni and Cu complexes. Not surprisingly, the energies of the other three structures are essentially the same for both Cu and Ni as they have all optimised to the same structure as discussed above. The energy differences between the two polymorphs for both Cu and Ni are small, although the difference between the two Ni structures is almost three times higher than for Cu.

This small difference between the enthalpies of polymorphs is not unusual. The experimental difference in energy between the α and β polymorphs of oxalic acid (corrected to ignore thermal and zero-point effects) is only 0.17 kJ/mol, whilst the calculated energy difference between the two forms of aspirin using a similar functional accounting for vDW interactions and excluding thermal and zero-point effects is 0.18 kJ/mol. However, with energy differences this small, the choice of vDW functional can change the calculated relative stabilities. Despite this, the calculated energy differences are consistent with the experimental results as although the 0 K calculations predict that both the Cu and Ni containing complex are most stable in the 100K Ni experimental structure, the energy difference in the case of Cu is only 0.37 kJ/mol which is significantly less than the K_T value of 0.83 kJ/mol at 100 K.

We have demonstrated that the energy difference between the two structural types is small. We now turn to Hirshfeld surfaces to characterise the geometric differences between the two structures. As illustrated in Figure 1, the differences appear to be slight, which is why we developed the idea of a difference fingerprint plot outlined in the Methodology section. Before applying difference fingerprint plots to the new structures presented here, it is instructive to utilise them to highlight the differences between two polymorphs whose fingerprint plots have already been discussed in the literature. We chose the system 2-chloro-4-nitrobenzoic acid (2c4n) that has two known polymorphs; both crystallise in P2₁/c and form I has a Z’ of 1 whilst form II has a Z’ of 4. Spackman and Jayatilaka have undertaken an extensive analysis of these two structures. The fingerprint plots of the five symmetry unique molecules generated directly from the Hirshfeld surface calculated in CrystalExplorer using our programs/scripts are reproduced in Figure 2.

Spackman and Jayatilaka concluded that the differences between the two modifications from comparing these fingerprint plots are:

1. The fingerprint plots for each of the four symmetry independent molecules in form II are more similar to each other than they are for the single symmetry independent molecule in form I.
2. All of the symmetry independent molecules in form II have increased intensity around d₁ = d₂ ≈ 2.0 Å, which indicates that they participate in a planar stacking arrangement (in this case π…π stacking), whereas no such arrangement is present in form I.

The two major similarities they identified are:

1. The dominant interaction in both forms is a cyclic hydrogen bond characteristic of carboxylic acids represented by a pair of spikes at the bottom left of the fingerprint plot.
2. All molecules participate in Cl…H hydrogen bonds characterized by a pair of much smaller spikes in the middle of the fingerprint plot at d₁ ≈ 2.0, d₂ ≈ 1.2 and d₁ ≈ 1.2, d₂ ≈ 2.0. These are particularly clear in the fingerprint for molecule C in form II.
Can the use of difference fingerprint plots make these differences clearer and provide additional insights? In Figure 3, the difference between the fingerprint plot of the molecule in form I and each of the molecules in form II is displayed.

Figure 3. Difference fingerprint plots between the single molecule of form I of 2c4n and the four molecules in form II.

Areas in blue are negative i.e. these areas are more intense in the form II fingerprint compared to the form I fingerprint. Thus, the strong blue areas in Figure 3 clearly illustrate that the form II molecules all have greater prevalence of planar stacking motifs compared to form I. Red then highlights areas with more intensity in the form I structure compared to form II. All of the difference plots have red areas at the top right because form I has a feature at these high d values (primarily due to interactions involving Cl) which is not present in the form II fingerprints as clearly shown in Figure 2. The other red feature in all the plots in Figure 3 is the spikes within the cyclic carboxylic acid spikes. These spikes finish at $d_{+} \approx 1.3 \text{ Å}$, $d_{-} \approx 1.0 \text{ Å}$ and $d_{-} \approx 1.0 \text{ Å}$, $d_{+} \approx 1.4 \text{ Å}$. These are due to the presence of shorter H–O–N distances in form I (two less than 2.5 Å) compared to all the molecules in form II (none less than 2.5 Å).

Figure 3 only shows the differences between the molecule in form I with those in form II. However for n molecules there are $n(n-1)/2$ difference fingerprint plots (the denominator of 2 is because for two fingerprint plots A and B, B-A is simply -(A-B) i.e. it is A-B with the colours reversed). For 2c4n with 5 symmetry independent molecules in the two polymorphs, $n = 5$ and thus there are 10 difference fingerprint plots to be considered.

Although a visual analysis of the difference fingerprint plots can provide useful qualitative information about where the molecular environments differ, can these differences be quantified? The simplest measure is to sum the absolute value of each point in the difference plot. This measure is similar to one of the three measures (the Manhattan metric) defined in the work of Latosińska et al.[13], although the magnitudes of our measure are clearly different; we note that the details of their procedure are yet to be published. Table 4 calculates this quantity for the 10 difference plots of 2c4n.

Table 4. Sum of the absolute values of the difference plots of 2c4n.

<table>
<thead>
<tr>
<th></th>
<th>Form II A</th>
<th>Form II B</th>
<th>Form II C</th>
<th>Form II D</th>
</tr>
</thead>
<tbody>
<tr>
<td>Form I</td>
<td>73.1</td>
<td>75.9</td>
<td>75.0</td>
<td>72.3</td>
</tr>
<tr>
<td>Form II</td>
<td>66.8</td>
<td>63.1</td>
<td>64.6</td>
<td></td>
</tr>
<tr>
<td>Form II</td>
<td>70.2</td>
<td>64.2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Form II</td>
<td>62.7</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The magnitude of these sums is surprisingly large given that comparing two plots where there are no areas in common would lead to a sum of 200 due to the normalisation of the individual fingerprint plots. The results in Table 4 support the observation from the visual comparison of the fingerprint plots by Spackman and Jayatilaka that each of the four symmetry independent molecules in form II are more similar to each other than they are to the single symmetry independent molecule in form I as the values in row I of Table 4 are greater than the values in the other rows. However, there are significant differences in the values obtained for the difference fingerprint plots obtained between the symmetry independent molecules in form II, with the difference fingerprint plot between independent molecules B and C closer to the form I – form II values. Thus, in Figure 4, the difference fingerprint plot of independent molecules B and C in form II are reproduced along with the difference fingerprint plot for form II independent molecules C and D, which is the plot with the lowest sum of absolute values.

Figure 4. Difference fingerprint plots between the independent molecules (a) B and C (b) C and D of form II of 2c4n.

The figure shows that there are significantly more differences in Figure 4(a) than Figure 4(b) and that the former is more similar to the plots in Figure 3. Consequently, the sum of the absolute values of a difference plot is a good measure of the magnitude of the differences between two fingerprint plots and has brought out differences in 2c4n that cannot be discerned simply by looking at the fingerprint plots in Figure 2.

We now turn to the Cu and Ni L complexes. The fingerprint plots corresponding to the experimental structures are given in Figure 5. Simple visual inspection shows that the three isomorphous structures, high temperature NiL$_2$ and the two CuL$_2$ structures, are similar and notably different to the plot for the low temperature NiL$_2$ structure. The dominant feature in the three similar fingerprint plots is at $d_{+} \approx 1.8$, $d_{-} \approx 1.8$ consistent with the
planar stacking arrangement observed.

The Hirshfeld surfaces emphasize that the planar stacking causes a particularly close approach between the phenol C atom and the oxime C atom of the neighboring molecule in all four structures (3.335 Å at 100 K NiL₂, 3.302 Å RT NiL₂, 3.240 Å 100 K CuL₂, 3.258 Å 150 K CuL₂). The unique low temperature Ni structure also shows a close approach between a p-methyl H atom and a neighbouring oxime O atom (C…O, 3.633 Å), which is not observed in the other systems.

In Figure 6, the difference plots between all the experimental structures are displayed and the corresponding sums of absolute values immediately provide some interesting insights. The lowest sum of absolute values by a large margin is for the isostructural CuL₂ and NiL₂ complexes discussed in this paper. In fact, by this measure, the difference between the isostructural CuL₂ at 100 K and NiL₂ at RT is slightly larger than the difference between the non-isostructural CuL₂ at 100 K and NiL₂ at 100 K.

Table 5. Sum of the absolute values of the difference plots of the CuL₂ and NiL₂ complexes discussed in this paper

<table>
<thead>
<tr>
<th></th>
<th>NiL₂ @ RT</th>
<th>CuL₂ @ 100 K</th>
<th>CuL₂ @ 150 K</th>
</tr>
</thead>
<tbody>
<tr>
<td>NiL₂ @ 100 K</td>
<td>60.8</td>
<td>56.8</td>
<td>56.7</td>
</tr>
<tr>
<td>NiL₂ @ RT</td>
<td>59.1</td>
<td>56.8</td>
<td></td>
</tr>
<tr>
<td>CuL₂ @ 100 K</td>
<td></td>
<td></td>
<td>41.1</td>
</tr>
</tbody>
</table>

Although it is not surprising that the highest sum is observed in one of the difference plots between NiL₂ at 100 K and the three other structures that are isostructural with each other, the size of the sum of absolute values of the difference plots between the three isostructural structures themselves are of a similar magnitude. In fact, by this measure, the difference between the isostructural CuL₂ at 100 K and NiL₂ at RT is slightly larger than the difference between the non-isostructural CuL₂ at 100 K and NiL₂ at 100 K.

![Figure 5](image)  
Figure 5. Fingerprint plots for the CuL₂ and NiL₂ complexes discussed in this paper.
The difference fingerprint plots of NiL₂ at RT with the two Cu structures (Figure 6(d,e)), all of which are isostructural, show that the Cu fingerprints have significantly more intensity centred around \(d_i \approx 1.7\), \(d_i \approx 1.7\) whereas the NiL₂ at RT has higher intensity centred around \(d_i \approx 1.9\), \(d_i \approx 1.9\), again consistent with the short phenol C atom to oxime C atom intermolecular distances.

For the plots for difference between the calculated Ni and Cu structures in the (c) NiL₂ 100 K structure and (d) NiL₂ RT structure, the difference plots between the calculated Ni and Cu structures in the (c) NiL₂ 100 K structure and (d) NiL₂ RT structure.

In Figure 7, the difference plots between the four calculated structures are displayed. The absolute sums of the differences of each metal in the two structural types in Figure 7(a,b) are 55.6 and 54.3 respectively and the difference plots look very similar. These difference plots are most similar to Figure 6(a), the difference plot for the experimental Ni structures, although it must be noted that the calculated structures are both at 0 K, whereas the experimental structures are at 100 K and RT respectively, leading to the greater magnitude differences observed in Figure 6(a). For the plots for difference between the Cu and Ni in a given structural type in Figure 7(c,d), the absolute sums of the differences are 47.9 and 47.3 respectively. This immediately shows that the differences due to the different structural types are much larger than the differences associated with changing the coordinating metal ion. Although this was also evident in the differences in experimental structures, the effect is much less marked (a difference in absolute values of \(-4\) in the experimental structures vs \(-9\) in the calculated structures), presumably because of the different temperatures of the crystal structure determinations. There is no experimental equivalent to Figure 7(c) as there is no Cu equivalent of the NiL₂ 100 K structure. However, for Figure 7(d), the closest experimental equivalent (albeit the experimental structures being determined at 150 K and RT) is Figure 6(e) and the two plots are similar in that there is a blue area at \(d_i \approx 1.7\), \(d_i \approx 1.7\), with a concentration of red immediately above the blue area.

The similarities between the difference plots derived from calculated structures compared to those derived from experimental structures suggest that the simulations are reproducing the fine detail of the packing in the structural types. A more direct way to assess the accuracy of the simulations is to calculate difference fingerprint plots between a calculated and experimental structure. The lower the temperature of the experimental structure, the better the anticipated agreement. Thus, in Figure 8, the difference plots between the calculated structures and the 100 K experimental structures are reproduced. The absolute sums of the differences are 47.5 for the NiL₂ 100 K structure and 45.8 for the CuL₂ 100 K structure (which is isostructural with the other two structures). These are the two lowest absolute sums of differences after the difference between the two Cu structures (which are isostructural at 100 K and 150 K) in Figure 6(f). The two plots in Figure 8 have little structure, again in common with Figure 6(f), illustrating that the differences between the calculated and experimental structures is small and are of the same order as differences between the experimental determinations at 100 K and 150 K. This assessment of the quality of the simulated structures would not be possible without the use of difference fingerprint plots.

Conclusions

The observation of temperature-dependent polymorphism in NiL₂, but not CuL₂, is consistent with the calculated relative energies of the structures, albeit with small energy differences. To examine structural differences in these systems in detail, we were driven to develop the software needed to generate difference fingerprint plots derived from Hirshfeld surfaces. Difference fingerprint plots are simply the result of subtracting one fingerprint plot from another. The resulting plots were found to reveal differences between structures that are not apparent upon visual inspection of the individual fingerprint plots. In addition, summing the absolute values of a difference fingerprint plot provides a quantitative measure of the difference between two structures. This summed value complements the qualitative conclusions drawn from visual inspection of the plots. This
quantitative measure may be particularly useful for providing a consistent means of comparing calculated and experimental structures, as well as classifying structures produced in polymorph prediction.

Acknowledgments

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Notes and references

Synthesis and structure determination of [Ni(L-NO₂)₂(dmso)₂]

**Synthesis**
Ni(NO₃)₂·6H₂O (0.0321 g, 0.11 mmol) was dissolved in ethyl acetate (10 mL) with dibutylphosphate (40 μL, 0.20 mmol). The yellow-green solution was examined using UV-Vis spectroscopy. Ketoxime LH (0.7452 g, 4.5 mmol) was dissolved in ethyl acetate (25 mL). The ketoxime solution 2.5 mL 0.18 M 0.45 mmol) was added to the metal/acid solution and mixed and the resulting green solution was re-examined via UV-Vis spectroscopy. After 24 hours the solution had darkened to dark yellow and a green jelly-like precipitate had formed. The precipitate was isolated and was found to be insoluble in dichloromethane, methanol, water, chloroform, acetone and ethyl acetate. The green precipitate was eventually dissolved in DMSO to give a yellow solution which was left to sit for several days and afforded red diamond shaped crystals.

**Crystallography**
The crystal data for [Ni(L-NO₂)₂(dmso)₂] are summarized in Table S1 with the structure depicted in Fig. S1 where ellipsoids have been drawn at the 50% probability level. Hydrogen bonding geometries are given in Table S2. Crystallographic data for the structures were collected at 100(2) K on an Oxford Diffraction Xcalibur diffractometer using Mo Kα radiation. Following analytical absorption corrections and solution by direct methods, the structure was refined against F² with full-matrix least-squares using the program SHELXL-97. The hydroxyl hydrogen atom was clearly observed in later difference maps and was refined without restraints. All remaining hydrogen atoms were added at calculated positions and refined by use of a riding model. The isotropic displacement parameters were based on the equivalent parameter of the parent atom. Anisotropic displacement parameters were employed for the non-hydrogen atoms.
Results

The results of the structure determination were consistent with the formulation \([\text{Ni}(\text{L-NO}_2)_2(\text{dmso})_2]\). The ketoxime ligand \(\text{HL}\) is presumed to have been nitrated as a result of strong heating in DMSO in the presence of nickel nitrate, converting the ligand to 2-hydroxy-5-methyl-3-nitroacetophenoneoxime (\(\text{HL-NO}_2\)). The molecule is situated on a crystallographic inversion centre. The oxime hydroxyl hydrogen atom is involved in bifurcated hydrogen bonds to the coordinated phenolic oxygen atom and to one of the oxygen atoms of the nitro group of the other ligand.

\textbf{Fig. S1} Structure of the molecule of \([\text{Ni}(\text{L-NO}_2)_2(\text{dmso})_2]\) showing the numbering scheme and the intramolecular hydrogen bonding.
**Table S1.** Crystal data and structure refinement for [Ni(L-NO$_2$)$_2$(dmos)$_2$] CCDC = 1526949.

<table>
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<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Empirical formula</td>
<td>C$<em>{22}$H$</em>{30}$N$<em>4$NiO$</em>{10}$S$_2$</td>
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<tr>
<td>Formula weight</td>
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<tr>
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<td>100(2) K</td>
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<tr>
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<tr>
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</tr>
<tr>
<td></td>
<td>$b = 7.7396(3)$ Å</td>
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<tr>
<td></td>
<td>$c = 15.3660(6)$ Å</td>
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<tr>
<td></td>
<td>$\beta = 95.439(5)^\circ$</td>
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<tr>
<td>Volume</td>
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<tr>
<td>Z</td>
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<tr>
<td>Density (calculated)</td>
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<tr>
<td>Absorption coefficient</td>
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<td>F(000)</td>
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<tr>
<td>Crystal size</td>
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<td>Independent reflections</td>
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<td>Full-matrix least-squares on $F^2$</td>
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<tr>
<td>R indices (all data)</td>
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<tr>
<td>Largest diff. peak and hole</td>
<td>0.538 and -0.346 e.Å$^{-3}$</td>
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Table S2. Hydrogen bonds for mojv9 [Å and °].

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<tr>
<th>D-H...A</th>
<th>d(D-H)</th>
<th>d(H...A)</th>
<th>d(D...A)</th>
<th>&lt;(DHA)</th>
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<tr>
<td>O(21)-H(21)...O(1)</td>
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<td>2.03(2)</td>
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<td>O(21)-H(21)...O(61)</td>
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<td>2.22(2)</td>
<td>2.8438(13)</td>
<td>137(2)</td>
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</tbody>
</table>

Symmetry transformations used to generate equivalent atoms:

* 1-x,1-y,1-z

References