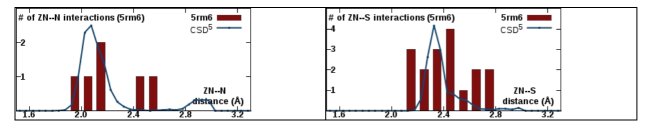
CheckMyMetal(CMM) report for PDB code: 5rm6

PDB title: Pandda analysis group deposition -- crystal structure of sars-cov-2 helicase in complex with z396380540 (2.1Å)

ID	Res.	Metal	Occupancy	B factor (env.) ¹		Valence ²	nVECSUM ³	Geometry ^{1,4}	gRMSD(°) ¹	Vacancy ¹	Bidentate	Alt. metal
A:701	ZN	Zn	1	<u>48.9</u> (60.3)	N_1S_3	<u>1.2</u>	<u>0.15</u>	Tetrahedral	<u>17.9°</u>	0	0	
A:702	ZN	Zn	1	72.4 (78)	N_2S_2	1.9	<u>0.13</u>	Tetrahedral	7.5°	0	0	
A:703	ZN	Zn	1	119.4 (118.9)	S ₄	<u>1.6</u>	0.24	<u>Trigonal</u> <u>Bipyramidal</u>	7°	20%	0	Co, Zn
B:702	ZN	Zn	1	58.7 (61.8)	N_2S_2	1.9	<u>0.17</u>	Tetrahedral	8°	0	0	
B:703	ZN	Zn	1	<u>95.1</u> (80.4)	S_3	1.7	0.39	Tetrahedral	8.8°	<u>25%</u>	0	Cu, Zn, Co
B:704	ZN	Zn	1	103.7 (105.5)	N_1S_3	<u>1.5</u>	0.44	Tetrahedral	<u>14.4°</u>	0	0	Cu, Co
Legend:			Not applicable Outlier <i>Borderline</i> Acceptable									

Column	Description								
	Occupancy of ion under consideration								
1 5	Occupancy of ion under consideration								
B factor	Metal ion B factor, with valence-weighted environmental average B factor in parenthesis								
$(env.)^1$									
Ligands	Elemental composition of the coordination sphere								
1/2lanca4	Summation of bond valence values for an ion binding site. <i>Valence</i> accounts for metal-ligand distances								
	Summation of ligand vectors, weighted by bond valence values and normalized by overall valence. Increase when the coordination sphere is not symmetrical due to incompleteness.								
Geometry ^{1,4}	Arrangement of ligands around the ion, as defined by the NEIGHBORHOOD algorithm								
	R.M.S. Deviation of observed geometry angles (L-M-L angles) compared to ideal geometry, in degrees								
Vacancy ¹	Percentage of unoccupied sites in the coordination sphere for the given geometry								
moeniale	Number of residues that form a bidentate interaction instead of being considered as multiple ligands								
Alt. metal	A list of alternative metal(s) is proposed in descending order of confidency, assuming metal environment is accurately determined. This feature is still experimental. It requires user discrimination and cannot be blindly accepted								

Metal-ligand distance distributions for pdb5rm6.ent in comparison with CSD



- (1) Zheng H, Chordia MD, Cooper DR, Chruszcz M, Müller P, Sheldrick GM, Minor W (2014) *Nature Protocols*, 9(1), 156-70.
- (2) Brown ID (2009) Chem. Rev., 109, 6858-6919.
- (3) Müller P, Köpke S, Sheldrick GM (2003) Acta Crystallogr. D Biol. Crystallogr., 59, 32-37.
- (4) Kuppuraj G, Dudev M, Lim C (2009) J. Phys. Chem. B, 113, 2952-2960.
- (5) CSD: Cambridge Structural Database
- Maintained by: Heping Zheng < dust@iwonka.med.virginia.edu >

Citing CheckMyMetal (CMM):

Validation of metal-binding sites in macromolecular structures with the CheckMyMetal web server. Zheng, H., Chordia, M.D., Cooper, D.R., Chruszcz, M., Müller, P., Sheldrick, G.M., Minor, W. (2014) Nature Protocols, 9(1), 156-70.