

**AhpC of the mycobacterial antioxidant defense system and its interaction with its  
reducing partner Thioredoxin-C**

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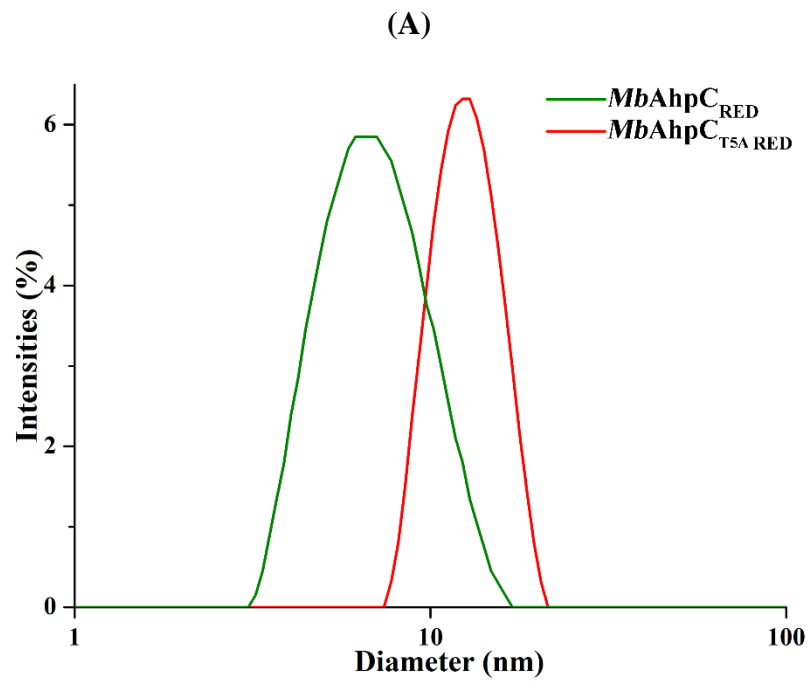
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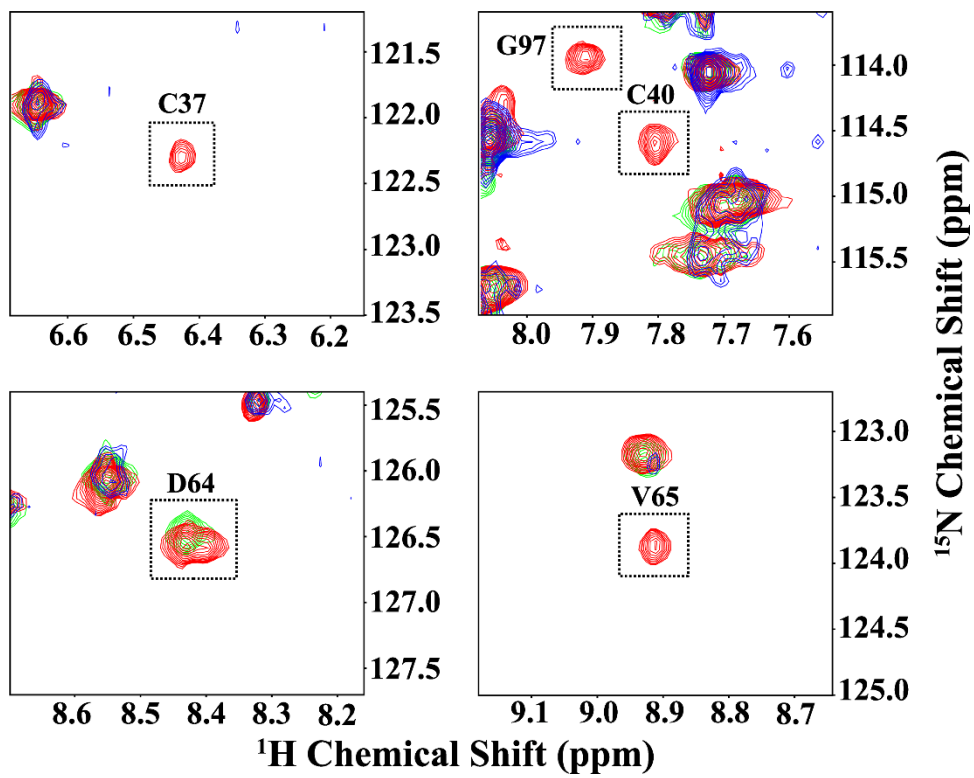
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**Supplementary Figure 1.** DLS analysis of reduced *MbAhpC* (*red*) and the reduced mutant *MbAhpC*<sub>T5A</sub> (*green*) revealed a smaller oligomer in *MbAhpC*<sub>T5A</sub> as compared to wt *MbAhpC*. This further illustrates the importance in T5 in maintaining the dodecamer formation.

**Supplementary Figure 2.** Selected sections of the overlaid <sup>1</sup>H-<sup>15</sup>N-HSQC spectra of free *MbTrxC* (*red*), as well as *MbTrxC-MbAhpC*<sub>T5A/D8A</sub> (*blue*) and *MbTrxC-MbAhpC*<sub>D22N/K25Q/D27N</sub> (*green*) complexes at a molar ratio of 1 (*MbTrxC*) : 2 (*MbAhpC* mutants). The residues undergoing disappearance of resonance or chemical shift perturbation are marked as box and labelled.



Supplementary Figure 1



Supplementary Figure 2