
Sypplementary Materials: Deregulated miRNAs Contribute to Silencing of B-cell Specific Transcription Factors and Activation of NF- κ B in Classical Hodgkin Lymphoma

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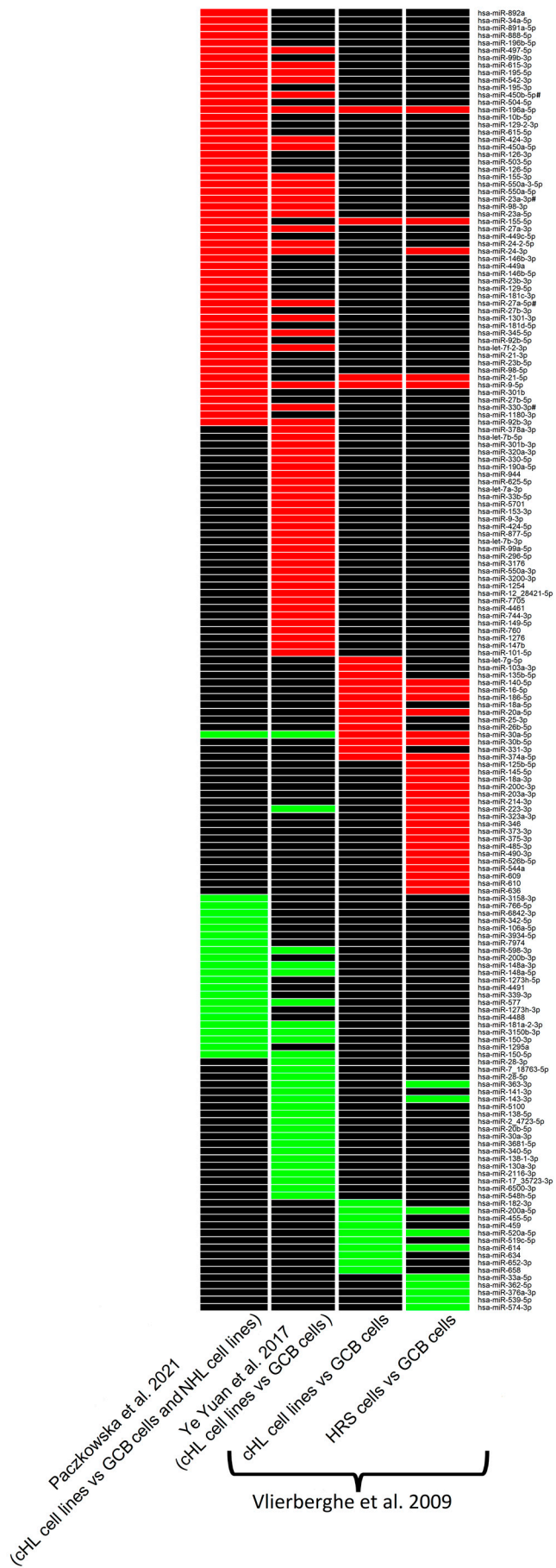
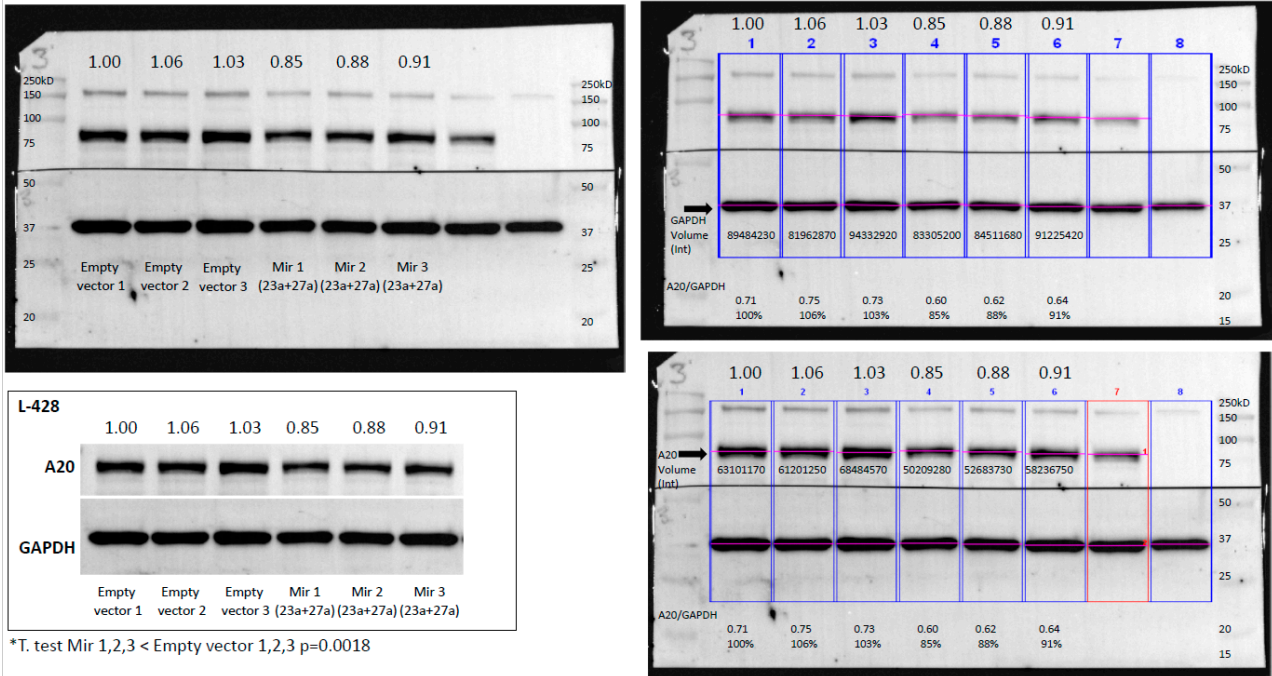


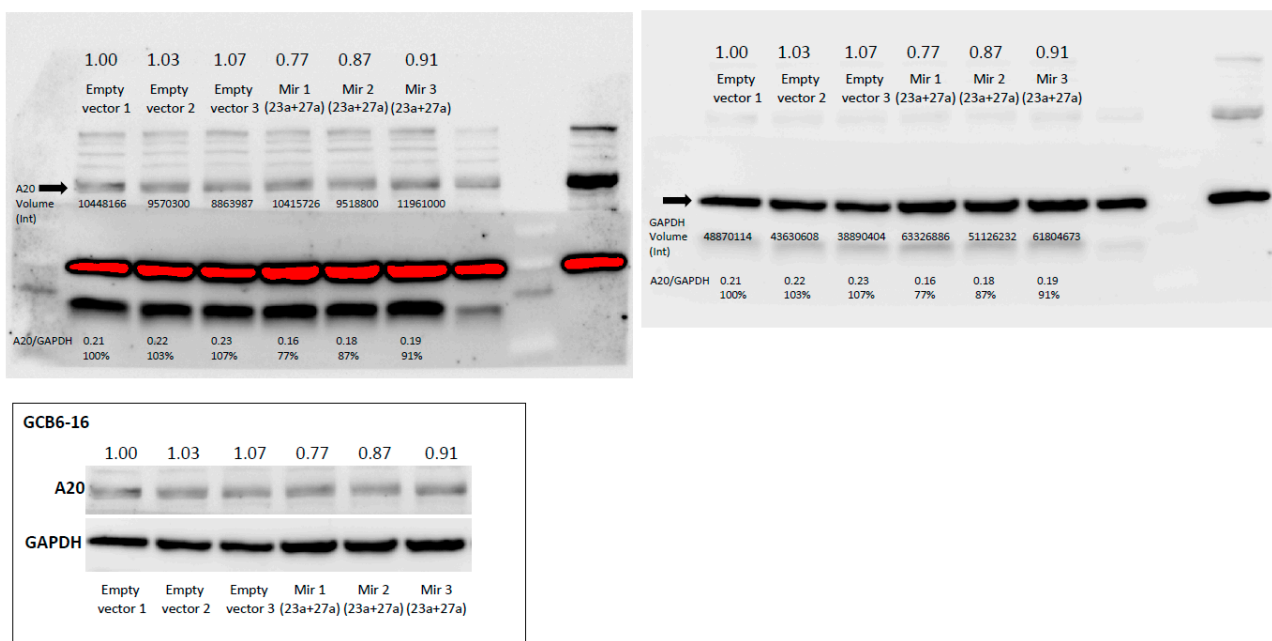
Figure S1. The overlap of deregulated miRNAs in cHL cell lines and primary HRS cells compared to NHL cell lines and GCB cells in previously published studies and this study. Columns represent miRNA expression profiles from the particular studies. Red and green colors indicate overexpressed and downregulated miRNAs, respectively, black color represents no significant change in miRNA expression in the respective study.

Figure 3. L-428



*T. test Mir 1,2,3 < Empty vector 1,2,3 p=0.0018

Figure 3. GCB6-16



*T. test Mir 1,2,3 < Empty vector 1,2,3 p= 0.0083

Figure S2. Uncropped Figure 3.