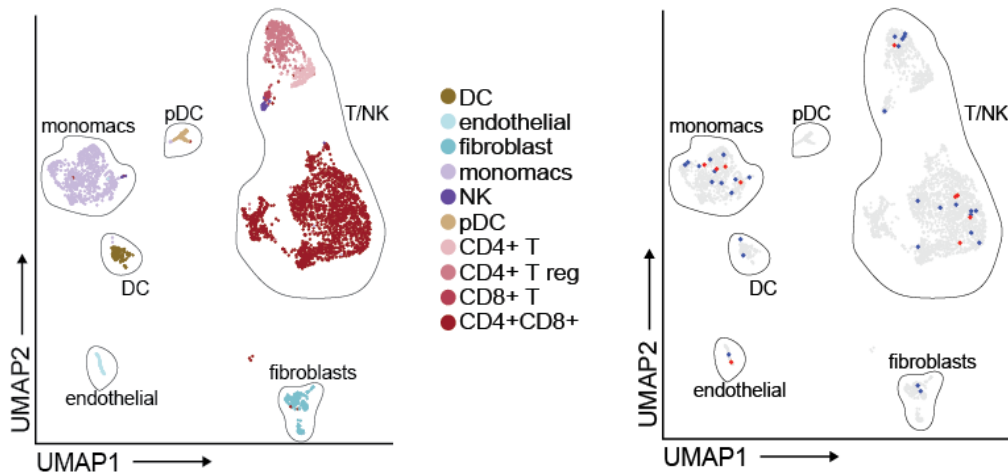


A clonally expanded nodal T-cell population diagnosed as T-cell lymphoma after CAR-T therapy

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Supplementary Figure



Supplementary Fig. 1 Legend: Left UMAP - all cells annotated by major cell type, reproduced from Figure 3A; Right UMAP - cells in which the wild-type (blue) or mutant (red) *TET2* p.776Wfs*4 sequences are identified. Cells in which neither wild type nor mutant sequence is identified are shown in gray.