S4 Fig. Association of epigenetic age-predictions with clinical parameters in AML. 

(A) The numbers of mutated genes in AML, as identified in deep sequencing data of TCGA portal, did not correlate with epigenetic age-predictions (99 CpG model). (B) In tendency female AML samples were predicted to be younger than male samples, which is in line with gender-specific deviations in normal blood. (C) The cellular composition in AML samples was estimated by bioinformatics methods described by Houseman et al. (2012; 2014). The predicted percentage of individual cellular subsets in AML samples was not clearly associated with chronological age or epigenetic age-predictions (even though the results suggest moderate increase of CD4+ T-cells in elderly patients).