

Correction

Predominant mutated non-canonical tumor-specific antigens identified by proteogenomics demonstrate immunogenicity and tumor suppression in CRC

Haitao Xiang (项海涛), Xiangyu Guan (关翔宇), Yaohua Wei (魏耀华), Shuzhen Luo (罗淑贞), Haibo Zhang (张海波), Fanyu Bu (卜繁宇), Yixin Yan (颜逸鑫), Yunyun Fu (付云芸), Yijian Li (李毅坚), Qumiao Xu (徐曲苗), Penghui Lin (林鹏辉), Dongbing Liu (刘栋兵), Xinlan Zhou (周鑫兰), Feng Gao (高峰), Tai Chen (陈泰), Guangjun Nie (聂广军), Kui Wu (吴逵), Ying Gu (顾颖), Longqi Liu (刘龙奇), Ziqing Ye (叶子青), * Xiaojian Wu (吴小剑), * Ruifang Zhao (赵瑞芳), * Siqi Liu (刘斯奇), * and Xuan Dong (董旋)*

*Correspondence: yzq2229@163.com (Z.Y.), wuxjian@mail.sysu.edu.cn (X.W.), zhaorf@nanoctr.cn (R.Z.), siqiliu@genomics.cn (S.L.), dongxuan@genomcs.cn (X.D.)
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During figure preparation for this article, the authors mistakenly labelled the x axes in Figures 4F and 4G with incorrect values. The figure has been updated with the correct values on the x axes.



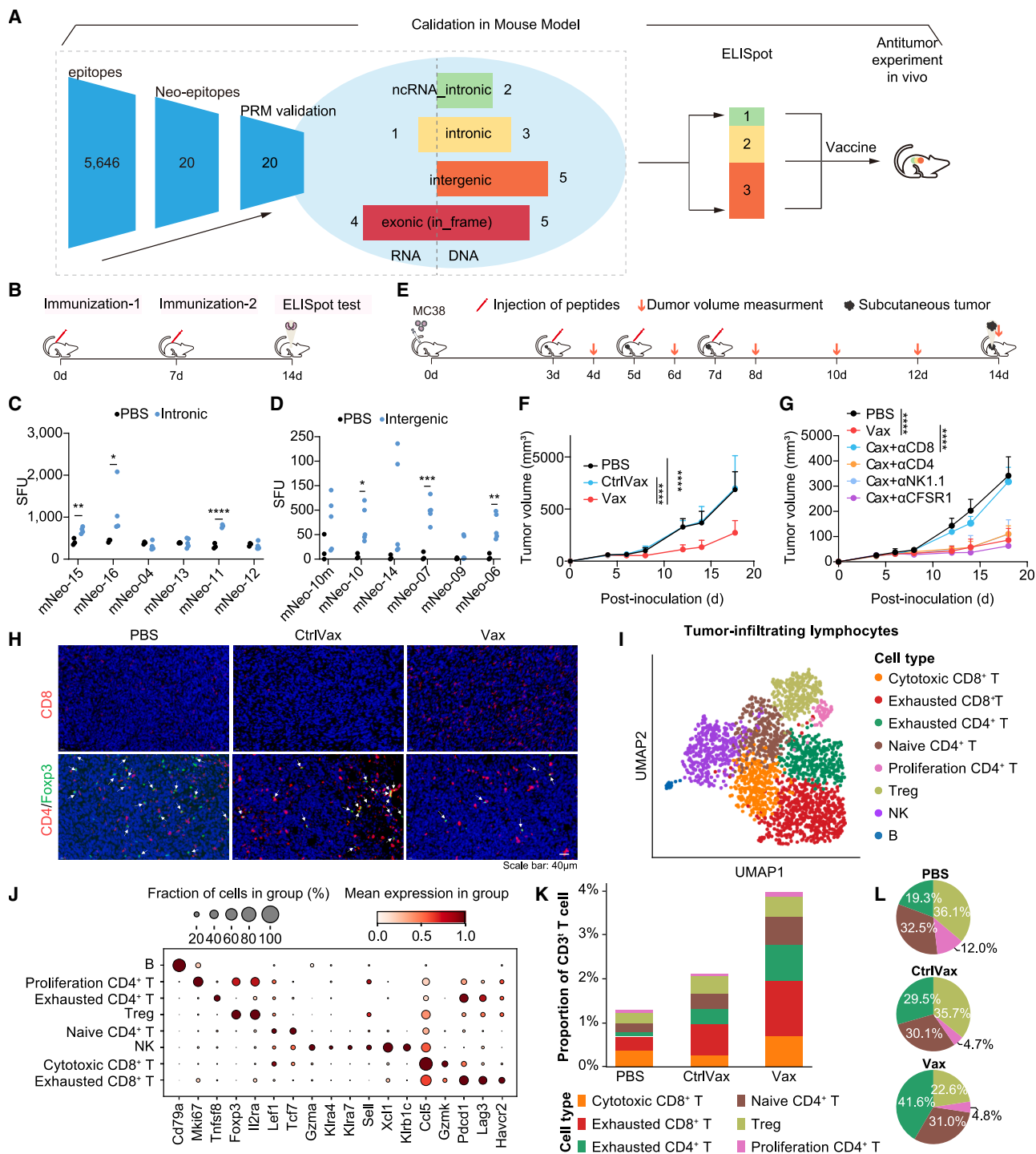


Figure 4. Non-canonical neo-antigens presented by MHC class I effectively activate CD8⁺ T cells and suppress tumor growth in a mouse model (corrected)

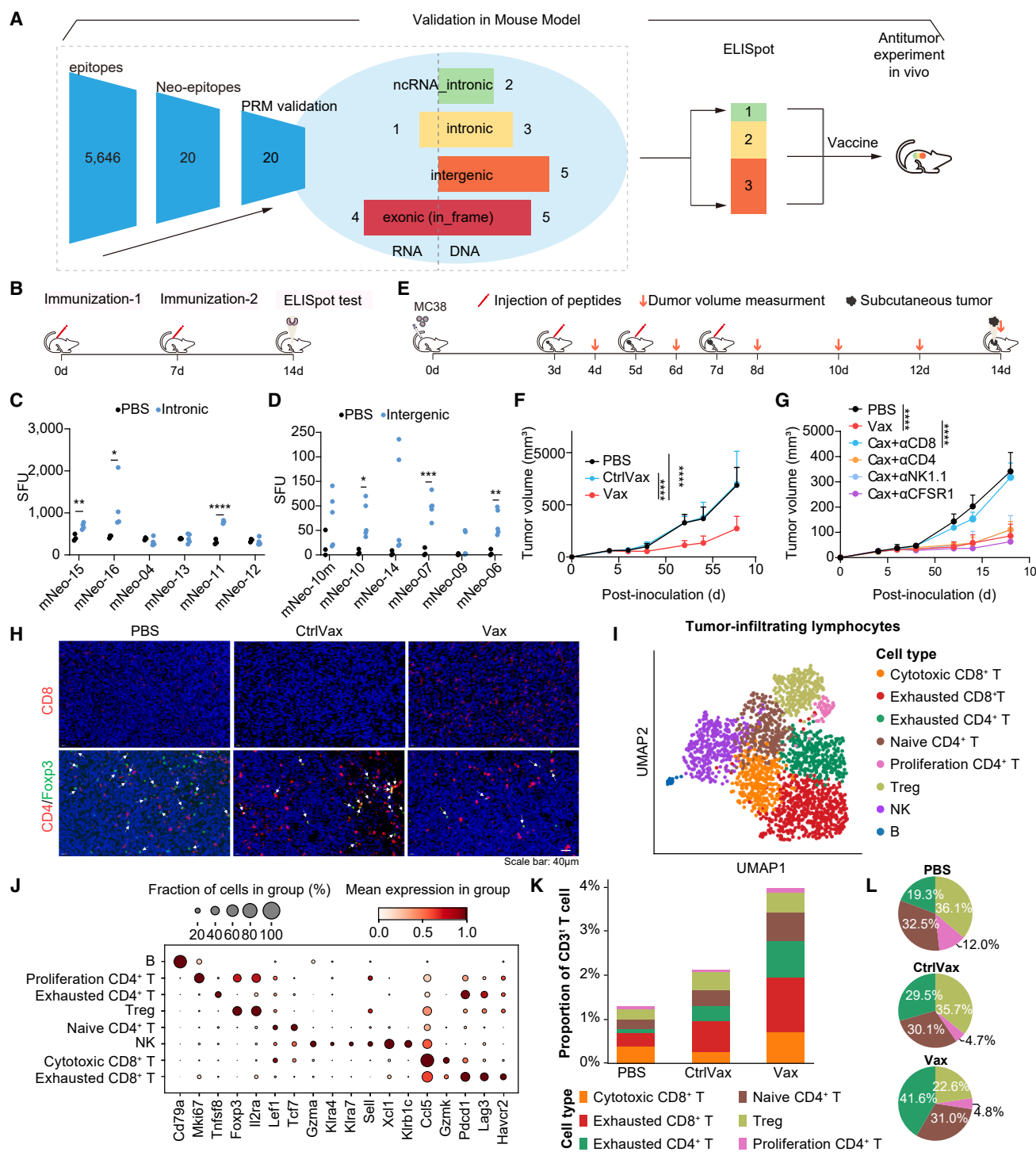


Figure 4. Non-canonical neo-antigens presented by MHC class I effectively activate CD8⁺ T cells and suppress tumor growth in a mouse model (original)