Single-cell transcriptome analysis reveals TOX as a promoting factor for T cell exhaustion and a predictor for anti-PD-1 responses in human cancer

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Abstract

Background: T cells exhibit heterogeneous functional states in the tumor microenvironment. Immune checkpoint inhibitors (ICIs) can reinvigorate only the stem cell-like progenitor exhausted T cells, which suggests that inhibiting the exhaustion progress will improve the efficacy of immunotherapy. Thus, regulatory factors promoting T cell exhaustion could serve as potential targets for delaying the process and improving ICI efficacy.

Methods: We analyzed the single-cell transcriptome data derived from human melanoma and non-small cell lung cancer (NSCLC) samples and classified the tumor-infiltrating (TI) CD8+ T cell population based on PDCD1 (PD-1) levels, i.e., PDCD1-high and PDCD1-low cells. Additionally, we identified differentially expressed genes as candidate factors regulating intra-tumoral T cell exhaustion. The co-expression of candidate genes with immune checkpoint (IC) molecules in the TI CD8+ T cells was confirmed by single-cell trajectory and flow cytometry analyses. The loss-of-function effect of the candidate regulator was examined by a cell-based knockdown assay. The clinical effect of the candidate regulator was evaluated based on the overall survival and anti-PD-1 responses.

Results: We retrieved many known factors for regulating T cell exhaustion among the differentially expressed genes between PDCD1-high and PDCD1-low subsets of the TI CD8+ T cells in human melanoma and NSCLC. TOX was the only transcription factor (TF) predicted in both tumor types. TOX levels tend to increase as CD8+ T cells become more exhausted. Flow cytometry analysis revealed a correlation between TOX expression and severity of intra-tumoral T cell exhaustion. TOX knockdown in the human TI CD8+ T cells resulted in downregulation of PD-1, TIM-3, TIGIT, and CTLA-4, which suggests that TOX promotes intra-tumoral T cell exhaustion by upregulating IC proteins in cancer. Finally, the TOX level in the TI T cells was found to be highly predictive of overall survival and anti-PD-1 efficacy in melanoma and NSCLC.

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Background

T cell dysfunction has been reported to be a hallmark of cancers [1]. However, the molecular mechanisms underlying tumor-infiltrating (TI) CD8$^+$ T cell exhaustion, especially the alterations in the transcriptional regulatory networks in cancer, are not completely understood. T cell exhaustion develops progressively during chronic antigen stimulation, which results in a heterogeneous population of exhausted T cells [2]. Recent studies have revealed that the progenitor exhausted and terminally exhausted CD8$^+$ T cells, which are the subsets of exhausted T cells, exhibit differential intrinsic effector functions and reinvigoration potential via programmed cell death 1 (PD-1) inhibition (reviewed in [2]). The progenitor exhausted and terminally exhausted TI CD8$^+$ T cell subsets exhibiting distinct epigenetic and transcriptional characteristics have been detected in the tumor microenvironment [3, 4].

TCF7 (also known as TCF1) is reported to be an essential transcription factor (TF) in the progenitor exhausted CD8$^+$ T cells, which exhibit a stem cell-like phenotype [2]. However, a master regulator that induces the differentiation of progenitor exhausted CD8$^+$ T cells into terminally exhausted CD8$^+$ T cells has not been identified. PD-1 expression is closely correlated with the severity of T cell exhaustion. Thus, several studies have attempted to identify a master regulator that triggers CD8$^+$ T cell exhaustion by focusing on PD-1 expression and the associated regulatory factors. Various regulators, such as eomesodermin (EOMES) and SMAD4, have been reported to be characteristically expressed in the terminally exhausted CD8$^+$ T cells [2, 5]. However, the role of these regulators in the direct regulation of the exhaustion program remains unclear. A detailed understanding of the mechanisms underlying the establishment and maintenance of terminally exhausted CD8$^+$ T cells will aid in the development of novel therapeutic strategies for cancer.

In this study, we demonstrate a strategy for predicting the TFs involved in cellular differentiation based on single-cell transcriptome data analysis. This analysis was used to identify the TFs that promote CD8$^+$ T cell exhaustion in tumors. The single-cell transcriptome data of human melanoma and non-small cell lung cancer (NSCLC) samples were analyzed to systematically predict the regulatory factors involved in T cell exhaustion. This analysis identified that several genes such as thymocyte selection-associated high mobility group box gene (TOX) and immune checkpoint (IC) genes can regulate T cell exhaustion. The analysis of expression dynamics along the pseudo-temporally ordered CD8$^+$ T cells from human tumors revealed that the expression of TOX increases with the exhaustion of CD8$^+$ T cells. Additionally, TOX positively regulated the expression of PD-1, TIM-3, TIGIT, and CTLA-4 in the human TI CD8$^+$ T cells. This suggested that TOX is a key TF that promotes T cell exhaustion by inducing IC molecules in human cancers. Finally, the expression levels of TOX in the TI T cells could predict the overall survival and response to anti-PD-1 therapy in human melanoma and NSCLC. These results suggest that TOX levels can be used for patient stratification during anti-cancer treatment, including immunotherapy, and that TOX can be targeted in the background of immune checkpoint inhibitor (ICI) therapy.

Methods

Preprocessing of single-cell transcriptome data and differential expression analysis

We analyzed the single-cell transcriptome data of tumor samples derived from 17 patients with melanoma (GSE72056) [6] and 14 patients with NSCLC (GSE99254) [7]. The transcriptome data were generated by full-length single-cell RNA sequencing (scRNA-seq) in a single batch. Expression level ($E$) based on the read count data from melanoma samples were normalized as:

$$E_{ij} = \log_2(\text{TPM}/10 + 1) \quad (\text{for the } i\text{th gene of the } j\text{th cell})$$

Cells with fewer than 1700 detected genes (defined by at least 1 mapped read) or exhibiting an average housekeeping expression level (as defined above) of less than 3 were excluded. The read count data from NSCLC samples were normalized by scran [8] method and centered by patient. Low-quality cells were filtered out if the number of expressed genes was smaller than ([median of all cells] − 3 × (median absolute deviation)] or if the mitochondrial gene count proportion of a cell was larger than 10%. Cells were also discarded if the TPM value of...
CD3D was < 3, TPM of CD8A < 3, or CD4 > 30 for fluorescence-activated cell sorting (FACS)-sorted CD8+ T cells and if the TPM of CD4 < 3 or CD8A > 30 for FACS-sorted CD4+ T cells. The preprocessing resulted in 4645 cells from melanoma samples and 11,769 cells from NSCLC samples. We used the normalized expression data as provided by the original studies for both scRNA-seq datasets. To investigate the transition of transcriptional states of CD8+ T cells in the tumor microenvironment, we used the single-cell transcriptome profiles for CD8+ T cell subset of the datasets. For the human melanoma dataset, we first isolated the cells annotated as “T cell” and performed clustering analysis using the Seurat v3 R package. We annotated each cluster based on the marker gene expression for major cell types, and isolated 1072 cells from the cluster annotated as CD8+ T cells based on the expression of CD8, but not of CD4 (CD4−CD8+). For the human NSCLC dataset, we used only 2123 cells annotated as “TTC cell” (tumor cytotoxic T cell) for CD8+ T cells. We divided the CD8+ T cells into 2 subsets based on the expression level of PDCD1 (also known as PD-1) into PDCD1-low (cells with below median $E_{PDCD1}$) and PDCD1-high (cells with above median $E_{PDCD1}$). Next, we analyzed the differential expression of each gene between PDCD1-low and PDCD1-high subsets using the Wilcoxon rank-sum test. The difference was considered statistically significant when the $P$ value was less than 0.05 (*), 0.01 (**), 0.001 (***) and 0.0001 (****). For both tumor scRNA-seq datasets, we selected the differentially expressed genes (DEGs) with $P < 0.001$. We further filtered out candidate genes with the mean of normalized expression value lower than a threshold (1 for melanoma and 2 for NSCLC) in both subsets. This filtration process resulted in 175 and 92 DEGs for melanoma and NSCLC datasets, respectively (Additional file 1: Table S1).

### Dimension reduction and visualization of single-cell transcriptome data

To visualize the relationship among individual cells based on high-dimensional gene expression data, we used t-stochastic neighbor embedding (tSNE) [9], which is one of the most popular methods for dimension reduction. We conducted the tSNE analysis using the Seurat v3 R package with the following parameters: perplexity, 30; number of iterations, 1000. To find the optimal number of PCA dimension, we ran “JackStraw” function of Seurat v3 and chose the largest dimension with $P < 0.05$. We projected the individual cells on the first two tSNE dimensions. Additionally, we used the violin plots to present the density distribution of cells with specific gene expression levels in the $PDCD1$-low and $PDCD1$-high subsets.

### Single-cell trajectory analysis

To investigate the kinetics of gene expression during CD8+ T cell differentiation in the tumor microenvironment, we performed single-cell trajectory analysis using the Monocle 2 software [10]. The scRNA-seq profiles of CD8+ T cells derived from human melanoma (GSE72056) [6] were used to reconstruct the single-cell trajectories for the effector, memory, and exhausted states. We defined the three T cell states of stable endpoint based on the expression of three marker genes [11–13]. The “classifyCells” function and marker expression data were used to classify the T cells into three cellular states: effector state (CD62L+ or CD127+ and $PDCD1^−$), exhausted state (PDCD1+), and memory state (CD62L+ or CD127+). The cells belonging to multiple states and those belonging to none of the three states were assigned as “ambiguous” and “unknown” states, respectively. The group-specific marker genes were selected using the “markerDiffTable” function. Next, we pseudo-temporally ordered the cells using the “reduce-Dimension” and “orderCells” functions. The expression dynamics along the trajectories were visualized using the BEAM analysis tools in the Monocle 2 software. The significance of upregulated expression in the exhausted T cells (or memory T cells) relative to the effector T cells was tested by one-tailed Mann-Whitney $U$ test.

### Clinical sample collection

For the flow cytometric analysis of immune cells, fresh tumor specimens were provided by the Department of Internal Medicine at the Severance Hospital, along with permission to conduct the following study. We enrolled 35 patients with NSCLC and 15 patients with head and neck squamous cell carcinoma (HNSCC) who were treated between 2017 and 2019 in Korea. Detailed information on human subjects has been listed in Additional file 2: Table S2.

### An internal cohort of patients with cancer undergoing anti-PD-1 treatment

To study the correlation between $TOX$ expression level in the TIL T cells and response to anti-PD-1 therapy, we recruited 16 patients with NSCLC from Yonsei Cancer Center, Seoul, Korea. The patients were administered nivolumab or pembrolizumab. Patients exhibiting partial response (PR) or stable disease (SD) for > 6 months were classified as responders, while the patients exhibiting progressive disease (PD) or SD for ≤ 6 months were classified as non-responders based on the Response Evaluation Criteria in Solid Tumors (RECIST) ver. 1.1 [14]. The tumor samples were obtained from patients before immunotherapy. Patient information is shown in Additional file 2: Table S3-4.
**Bulk RNA sequencing data analysis of tumor samples**

Bulk RNA sequencing was performed for 16 samples from patients treated with the PD-1 inhibitor. Of the 16 tumor samples, 11 were fresh samples and 5 were formalin-fixed paraffin-embedded (FFPE) samples. The library was prepared from the samples using the TruSeq RNA Access Library Prep Guide Part # 15049525 Rev. B with the TruSeq RNA Access Library Prep Kit (Illumina). RNA sequencing was performed in HiSeq 2500 (Illumina). The obtained sequencing data were processed as per the manufacturer’s instructions. The read data were aligned with the reference genome (GENCODE, h19 (GRCh37.p13, release 19)) [15] using STAR-2.5.2a [16]. The transcripts were quantified using feature-Counts [17]. The correlation between the read count values of genes between fresh and FFPE samples was evaluated using Pearson’s correlation coefficient. The correlations between intra-fresh sample, intra-FFPE sample, and fresh-FFPE samples as evaluated by Wilcoxon’s rank-sum test were found to be significant.

**Isolation of Ti lymphocytes from the primary tumor**

Primary tumor tissues were obtained by surgical resection of patient tumors and from tumors developed in mice. The tissues were minced into 1 mm³ pieces and digested with a solution containing 1 mg/mL collagenase type IV (Worthington Biochemical Corp.) and 0.01 mg/mL DNase I (Millipore Sigma Corp.) at 37 °C for 20 min. The dissociated tissues were filtered using a 40-μm cell strainer (Falcon, Corning) into the Roswell Park Memorial Institute (RPMI)-1640 medium (Corning Inc., Corning). Tumor-infiltrating lymphocytes (TILs) were separated using a Percoll gradient (Millipore Sigma Corp.) and suspended in phosphate-buffered saline (PBS) supplemented with 2% fetal bovine serum (FBS; Biowest). The single-cell suspensions were stained with the indicated fluorescent dye-conjugated antibodies.

**Flow cytometric analysis**

Single-cell suspensions were analyzed using the CytoFLEX or CytoFLEX LX flow cytometers (Beckman Coulter) after staining with the following antibodies for murine tissues: CD4-BV785 (RM4-5), CD4-BV510 (RM4-5), CD8-Alexa Fluor 700 (53-6.7), CD8-PerCP-Cy5.5 (53-6.7), PD-1-BV605 (29F.1A12), PD-1-BV421 (29F.1A12), TIM-3-BV605 (RMT3-23), TIM-3-PerCP-Cy5.5 (B8.2C12), LAG-3-BV650 (C9B7W), and T-BET-BV421 (4B10) antibodies (all from BioLegend); TIGIT-BV785 (1G9) antibody (BD Biosciences); 2B4-FITC (eBio244F4) and EOMES-APC (Dan11mag) antibodies (all from Invitrogen); CTLA-4-Alexa Fluor 700 (63828) antibody (R&D Systems); TCF1-Alexa Fluor 488 (C63D9) antibody (Cell Signaling); and TOX-PE (REA473), REA Control (I)-PE (REA293), NR4A1(NUR77)-APC (REA704), and REA Control-APC (REA293) antibodies (all from Miltenyi Biotec). Dead cells were stained using the Live/Dead Fixable Near-IR Dead Cell Stain Kit (Invitrogen). For TF staining, the TI lymphocytes were fixed and permeabilized using the FOXP3 fixation/permeabilization solution (eBioscience). T-BET, EOMES, TCF1, NR4A1, and TOX antibodies or their isotype control antibodies were used for staining after permeabilization.

The following antibodies were used for human sample staining: CD3-BV785 (OKT3), CD8-BV605 (RPA-T8), CD8-BV510 (RPA-T8), CD8-BV650 (RPA-T8), and PD-1-BV421 (EH12.2H7) antibodies (all from BioLegend); T1M-3-BV605 (F38-2E2), LAG-3-FITC (11C3C65), CTLA-4-PE-Cy7 (BN13), 2B4-Alexa Fluor 700 (C1.7), IFN-γ-APC (4S.B3), and TNF-α-PE-Cy7 (MAb11) antibodies; T1M-3-Alexa Fluor 488 (344823) and TIGIT-APC (741182) antibodies (all from R&D Systems); CD4-APC-H7 (L200), TIGIT-BV510 (741182), and T-BET-BV650 (04-46) antibodies (all from BD Biosciences); TOX-PE (REA473), REA Control (I)-PE (REA293), NR4A1-APC (REA704), and REA Control-APC (REA293) antibodies (all from Miltenyi Biotec); TCF1-Alexa Fluor 488 (C63D9) antibodies (Cell Signaling); and EOMES-APC (WD1928), TOX-PE (TXRX10), and Rat IgG2a kappa isotype control-PE (eBR2a) antibodies (Invitrogen). Dead cells were excluded by staining with the Live/Dead™ Fixable Red Dead Cell Stain Kit (Invitrogen). For staining the intra-cellular cytokines and TFs, the cells were fixed and permeabilized with the Foxp3 fixation/permeabilization solution (eBioscience), followed by staining with antibodies against IFN-γ, TNF-α, T-BET, EOMES, TCF1, NR4A1, and TOX, or their isotype controls. Cells were analyzed using the FlowJo software (Tree Star Inc.). The gating strategy used to identify the human TI CD8⁺ T cells is shown in Additional file 2: Figure S1A.

**In vivo tumor models**

Five to 6-week-old female C57BL/6 mice and Balb/c mice were purchased from Charles River Laboratories and The Jackson Laboratory, respectively. MC38 colon cancer cells, TC-1 lung cancer cells, or LLC1 lung cancer cells were injected subcutaneously (10⁶ cells) into the C57BL/6 mice. The CT26 colon cancer cells were injected subcutaneously (10⁶ cells) into the Balb/c mice. The mice were euthanized at day 21 post-tumor cell injection.

**TOX knockdown in human TI CD8⁺ T cells**

The primary lung cancer cells were dissociated using gentle MACS Dissocator (Miltenyi Biotec) and Human Tumor Dissociation Kit (Miltenyi Biotec), following the manufacturer’s instructions. The TILs were transfected with TOX siRNA—that suppresses TOX
expression—or with control siRNA (Thermo Fisher Scientific) using the Neon transfection system (Invitrogen). The transfected TI lymphocytes were stimulated with the anti-CD3 antibody (1 μg/mL, OKT3, Miltenyi Biotec) coated on the plate for 84 h. For the functional analysis, the cells were re-stimulated for another 6 h with the anti-CD3 antibody in the presence of both GolgiStop and GolgiPlug (BD Biosciences) and stained with antibodies against IFN-γ and TNF-α. Gene knockdown was confirmed by flow cytometry.

**Statistical test for experimental data**

The statistical significance was analyzed using two-tailed unpaired and paired Student’s t tests in the Prism 5.02 software (GraphPad). The data are expressed as mean ± standard error of mean (SEM). The difference was considered statistically significant when the P value was less than 0.05 (*), 0.01 (**), 0.001 (***) and 0.0001 (****).

**Survival analysis and anti-PD-1 response analysis**

We evaluated the T cell-specific TOX gene expression and demonstrated the correlation between TOX expression level in T cells and that in the CD8⁺ T cells using single-cell transcriptome data obtained from human melanoma samples [6]. To evaluate the clinical effect of TOX expression in only TI T cells, we normalized TOX expression to the expression level in the TI T cells using the geometric mean of the expression levels of CD3D, CD3E, and CD3G.

Survival analysis based on the transcriptome and clinical data compiled from The Cancer Genome Atlas (TCGA) for melanoma (SKCM, skin cutaneous melanoma) and NSCLC (LUAD, lung adenocarcinoma, and LUSC, lung squamous cell carcinoma) was performed. The bulk RNA-seq data for tumor samples were downloaded from the Xena database (https://xena.ucsc.edu/), while the clinical data were downloaded from TCGA-CDR [18]. For the survival analysis of patients with NSCLC, we used that data of patients exhibiting top 25% tumor mutation burden. Next, we compared the overall survival rates of patients exhibiting top 30% TOX level (TOX-high) with those of all other patients (TOX-low). The significance of difference between survival times among groups was examined using the log-rank test.

For analyzing the correlation between anti-PD-1 response and TOX expression levels in the TI T cells, we analyzed the bulk RNA-seq data along with clinical information for four independent cohorts of patients who underwent anti-PD-1 immunotherapy, including three published reports [19–21]. Similar to that in the survival analysis, we normalized the TOX expression and generated waterfall plots after segregating the patients based on the mean TOX expression values. For the Hugo et al. [19] dataset, we classified patients annotated as “Partial Response” and “Complete Response” into the responder group, and those annotated as “Progressive Disease” into the non-responder group. For the Jung et al. [21] dataset, we classified the patients annotated as DCB (Durable Clinical Benefit) into the responder group and those annotated as NDB (Non-Durable Benefit) into the non-responder group. For the Riaz et al. [20] dataset, we excluded patients annotated as “Stable Disease” and classified the rest of the patients into two groups in a manner similar to that used for generating the Hugo et al. dataset. To evaluate the predictive power of TOX expression in the TI T cells for governing the responses to anti-PD-1 therapy, we prioritized the patients by sorting the patients exhibiting the lowest TOX level in the TI T cells and performed receiver operating characteristic (ROC) analysis.

**Results**

**Subset analysis of single-cell transcriptome profiles of CD8⁺ T cells for identifying regulators involved in T cell exhaustion**

The distinct cellular states can be often depicted using the expression of single marker genes. Thus, we can identify key genes involved in the progression of T cell exhaustion by analyzing the DEGs between the progenitor exhausted T cells and terminally exhausted T cells. However, the exhausted CD8⁺ T cells in the tumor microenvironment exhibit a continuous spectrum of transcriptional states depending on the exhaustion severity levels [22]. Therefore, we developed a strategy for identifying the genes involved in T cell exhaustion using single-cell transcriptome data (Fig. 1a). The exhausted CD8⁺ T cells exhibiting intermediate PD-1 expression can be reinvigorated by PD-1 inhibition, whereas those exhibiting high PD-1 expression are refractory to this effect [23]. Therefore, we segregated the TI CD8⁺ T cells into two subsets based on the median expression value of PDCD1, i.e., PDCD1-high and PDCD1-low subsets. The localized distribution of PDCD1-high cells in the two-dimensional latent space of the tSNE plot indicated that the PDCD1 marker can aid in distinguishing between the terminally exhausted and progenitor exhausted cells. The DEGs between the two subsets might be the potential factors associated with T cell exhaustion, which can also be confirmed from the similar distribution of DEG-high cells in the same two-dimensional latent space.

This strategy was applied to analyze the single-cell transcriptome profiles of CD8⁺ T cells derived from melanoma [6] (Fig. 1b), which revealed the localized distribution of PDCD1-high cells in the tSNE plot. We identified 175 DEGs between the PDCD1-high and PDCD1-low subsets using the Wilcoxon test (P < 0.001)
Fig. 1 (See legend on next page.)
TOX and IC genes exhibited similar expression dynamics along the single-cell trajectories for TI CD8⁺ T cells

TI T cells that are initially in the effector state (T-eff) soon start becoming dysfunctional and get converted into exhausted T cells (T-exh) as a result of the highly immunosuppressive tumor microenvironment. A subset of persisting T-eff cells differentiates into long-lived and self-renewable memory T cells (T-mem). We hypothesized that if TOX promotes T cell exhaustion, the expression dynamics of TOX during the transition from T-eff to T-exh should differ from that during the transition from T-eff to T-mem. To test this hypothesis, we reconstructed the single-cell trajectories composed of pseudo-temporally ordered CD8⁺ T cells across the three distinct T cell states using the Monocle 2 software [10]. All three single-cell trajectories were significantly enriched for the corresponding cell type assigned based on marker expression (P < 2.2e−16 for exhausted state and memory state, \( P = 7.07e−07 \) for effector state by binomial test) (Fig. 2a), which validates the established trajectories of T cell differentiation in tumor. We found that none of the three branches was severely biased in distribution of cells from each patient, indicating that the observed differential enrichment of three distinct T cell states among trajectories is not based on patient-specific effects (Fig. 2b).

The expression of IC genes, such as CTLA4, HAVCR2, LAG3, PDCD1, and TIGIT, was upregulated along the pseudotime of CD8⁺ T cell exhaustion (Fig. 2c, d). Compared to the effector state, the exhaustion state (but not the memory state) was associated with significantly enhanced expression of IC genes (P values are one-tailed Mann-Whitney U test) (Fig. 2d). Notably, the changes in the expression of TOX followed an identical trend along the pseudotime trajectories (Fig. 2c, d). As the expression level of IC molecules is correlated with the severity of CD8⁺ T cell exhaustion, these results indicate that TOX expression is correlated with the severity of CD8⁺ T cell exhaustion in tumors.

Further, we examined the expression dynamics of other TFs reported to be involved in T cell exhaustion. The expression of NRA1, a TF that induces T cell exhaustion [27, 28], was upregulated at later stages of the exhaustion state. However, the upregulated expression of IC genes, such as HAVCR2, additional file 1: Table S1A). Notably, the expression levels of IC genes, such as HAVCR2 (also known as Tim-3), CTLA4, and TIGIT in the PDCD1-high subset were higher than those in the PDCD1-low subset. Additionally, the distribution patterns of the IC genes in the DEG-high and those in the PDCD1-high cells were similar in the tSNE plot. A subset analysis of single-cell transcriptome profiles of CD8⁺ T cells derived from NSCLC [7] (Fig. 1c) revealed 92 DEGs (Additional file 1: Table S1B). The single-cell transcriptome profile analysis of NSCLC samples revealed that the IC genes in the PDCD1-high subset were upregulated. The correlation between PDCD1 expression and other IC genes validated the effectiveness of subset analysis of single-cell transcriptome data for identifying the genes involved in T cell exhaustion. We found that neither PDCD1-high nor PDCD1-low subset were severely biased in distribution of cells from each patient in both single-cell transcriptome datasets reanalyzed, indicating that the observed differential expression is not based on patient-specific effect (Additional file 2: Figure S2).

To identify the key trans-acting regulators involved in the regulation of intra-tumoral T cell exhaustion, we focused on 13 and 5 TFs (annotated by Ravasi et al. [24]) among the DEGs identified from melanoma and NSCLC samples, respectively. We successfully retrieved several TFs previously reported to be involved in T cell exhaustion, such as BATF [25], NFATC1 [25], and PRDM1 [26]. This further highlighted the effectiveness of predicting regulatory factors using single-cell transcriptome data. We observed that the TFs are differentially expressed in the PDCD1-high and PDCD1-low subsets among the melanoma or NSCLC samples (Fig. 1d). Some regulators may be specifically involved in particular cancer types. The tumor specificity identified based on the statistical analysis must be evaluated by follow-up functional analysis. Notably, TOX was the only candidate TF identified in both melanoma and NSCLC samples. The distribution patterns of TOX-high cells and PDCD1-high cells were similar in the latent space of the tSNE plot for both melanoma and NSCLC, which were similar to the distribution patterns of IC genes (see Fig. 1b, c).
during the entire exhaustion state was not significant (P = 0.1209). Recently, NFAT1 (also known as NFATC2) and NFAT2 (also known as NFATC1) were reported to be the TFs that promote T cell exhaustion [29–31]. The expression of NFATC1 (P = 7.637e-05), but not that of NFATC2 (P = 0.8299), in the exhausted state was significantly higher than that in the effector state, which was in agreement with the results of a previous study [31]. Interestingly, the same study also demonstrated that TOX expression is induced by NFAT2 (NFATC1). TOX2 was also reported to be involved in CD8+ T cell exhaustion [28, 30, 32]. We could not detect the upregulated expression of TOX2 in the TI T cells as the expression was low. However, the expression level of TOX2 in the exhausted state was significantly higher than that in the effector state (P = 1.915e-05). TCF7 (also known as TCF1) is a key regulator of progenitor exhausted T cells [33–37]. The expression of TCF7 in the memory state was significantly higher than that in the effector state (P = 7.01e-08). This result is consistent with that of a previous study that reported the essential roles of Tcf-1 in establishing the CD8+ T cell memory [38] and the memory-like cell functions of Tcf-1+CD8+ T cell in chronic infections [36] in the mouse model. Consequently, the overall consistency of the observed expression dynamics of known TFs for T cell exhaustion in single-cell trajectories in conjunction with the previous studies validates our reconstructed single-cell trajectories of T cell differentiation in tumors.

### TOX protein level correlated with the severity of intra-tumoral CD8+ T cell exhaustion in human cancers

The correlation between TOX protein expression and severity of intra-tumoral T cell exhaustion was evaluated by flow cytometric analysis of TI lymphocytes isolated from human primary tumor specimens from patients with NSCLC or HNSCC who underwent surgical resection at the Severance Hospital (Additional file 2: Table S2). TOX expression was positively correlated with the expression of various IC molecules, such as PD-1, TIM-3, and TIGIT—at protein level—in the TI CD8+ T cells isolated from both NSCLC and HNSCC tumor tissues (Fig. 3a). Additionally, the proportion of TOX+ TI CD8+ T cells was significantly associated with the expression of IC molecules (Fig. 3b). The TOX+ TI CD8+ T cells were significantly enriched in the population expressing other IC molecules, such as CTLA-4, LAG-3, and 2B4 (Additional file 2: Figure S1B). The PD-1+TIM-3+ CD8+ T cells exhibit the terminally exhausted phenotype, while the PD-1+TIM-3+ CD8+ T cells exhibit the progenitor exhausted phenotype in chronic viral infection as well as tumors [39, 40]. Thus, we sub-divided the population into PD-1+TIM-3-, PD-1+TIM-3+, and PD-1+TIM-3- cells and compared the TOX levels in these three subsets. Among the subsets, the terminally exhausted TI CD8+ T cells exhibited significantly high TOX level. Each population can be arranged in the following order of decreasing TOX expression: PD-1+TIM-3+ > PD-1+TIM-3+ > PD-1+TIM-3+ (Fig. 3c). Flow cytometric analysis of mouse TI CD8+ cells isolated from various cancer models, including MC38 colon cancer, CT26 colon cancer, TC-1 lung cancer, and LLC1 lung cancer, revealed a similar correlation between TOX expression and TI CD8+ T cell exhaustion severity (Additional file 2: Figure S3 and S4A). These results strongly suggest that TOX expression is closely associated with the severity of TI CD8+ T cell exhaustion.

We also investigated the correlation between other TFs and TI CD8+ T cell exhaustion severity. The expression patterns of NR4A1, T-BET, EOMES, and TCF1—which are reported to regulate T cell exhaustion—were examined in the human NSCLC and mouse tumors, including MC38, CT26, TC-1, and LLC1. The expression levels of other TFs did not correlate with PD-1 expression in the TI CD8+ T cells from human NSCLC tumors (Additional file 2: Figure S1C). Similarly, the expression levels of TFs, such as NR4A1 and T-BET, were not correlated with PD-1 expression in the TI CD8+ T cells from various mouse tumors (Additional file 2: Figure S4B). The results of the flow cytometric analysis agreed with those of the single-cell trajectory analysis. The analysis revealed that the expression of NR4A1 and TCF7 in the exhausted T cells was not significantly upregulated.
when compared to that in the effector T cells (see Fig. 2d). Notably, in some mouse tumors, the expression levels of EOMES and TCF1 were negatively and positively correlated with PD-1 expression, respectively. These results indicate that among the TFs reported to be involved in CD8\(^+\) T cell exhaustion, only the expression level of TOX was positively correlated with PD-1 expression.

**TOX knockdown disrupts expression of checkpoint molecules in human TI CD8\(^+\) T cells and restores their anti-tumor function**

As TOX expression was positively correlated with the severity of CD8\(^+\) T cell exhaustion, we hypothesized that TOX is a positive regulator of the exhaustion process in human cancers. Therefore, we evaluated the effect of TOX loss of function on IC molecules in the human TI

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Fig. 3 Correlation of TOX expression with the severity of CD8\(^+\) T cell exhaustion in human tumors. **a**–**c** Flow cytometric analysis of the tumor-infiltrating (TI) CD8\(^+\) T cells isolated from human non-small cell lung cancer (NSCLC) (n = 20) and head and neck squamous cell carcinoma (HNSC C) (n = 15). **a** Representative plots showing the co-expression of TOX and immune checkpoint (IC) molecules (PD-1, TIM-3, and TIGIT) in the TI CD8\(^+\) T cells. **b** Percentage of TOX\(^+\) cells in the two subpopulations of the TI CD8\(^+\) T cells (expressing or not expressing a specific IC molecule). Each line in the graph indicates the data derived from the same tumor tissue of each individual patient. **c** TOX protein levels in the three subsets of TI CD8\(^+\) T cells with different severities of exhaustion, i.e., PD-1\(^-\)TIM-3\(^-\) (orange), PD-1\(^+\)TIM-3\(^-\) (blue), and PD-1\(^+\)TIM-3\(^+\) (red). Histogram represents TOX expression level in each subset of the TI CD8\(^+\) T cells. Distribution of TOX-expressing subsets of TI CD8\(^+\) T cells across patients is summarized in grouped scattered plots. ns, not significant; *P < 0.05; **P < 0.01; ***P < 0.001. All statistical analyses were performed using unpaired Student's t test.
CD8+ T cells. The TI lymphocytes derived from tumor tissue of patients with NSCLC were transfected with TOX siRNA. Interestingly, the knockdown of TOX in the TI CD8+ T cells resulted in a significant reduction of cells expressing IC molecules, such as PD-1, TIM-3, TIGIT, and CTLA-4. On the other hand, TOX knockdown did not lead to the statistically consistent pattern of change in the percentage of LAG-3- or 2B4-expressing TI CD8+ T cells (Fig. 4a). These results indicate that TOX positively regulates the expression of various IC molecules to promote CD8+ T cell exhaustion in human cancer. Additionally, we observed that the frequency of TI CD8+ T cells that secrete effector cytokines (IFN-γ and TNF-α) significantly increased upon TOX knockdown, suggesting that the anti-tumor function of TI CD8+ T cells could be restored upon TOX knockdown (Fig. 4b). Taken together, these observations suggest that TOX is a key regulator of terminally exhausted CD8+ T cell differentiation in human cancer.

**TOX expression level in TI T cells is predictive for overall survival and anti-PD-1 efficacy in human cancers**

As T cells play a major role in eliminating the cancer cells from tumors, their functional state affects the prognosis and therapeutic efficacy. In this study, the single-cell transcriptome profiles of melanoma patients...
indicated that TOX exhibited a highly specific expression pattern in the T cells [6] (Fig. 5a). As there was a correlation between TOX expression and TI CD8⁺ T cell exhaustion, we hypothesized that TOX expression in the TI T cells may be used as a clinical indicator during cancer treatment. To test this hypothesis, we analyzed the TCGA survival data with respect to the corrected TOX expression level to evaluate the effect of T cell levels within each sample by classifying the TOX expression based on the geometric mean of CD3 gene expression levels (CD3D, CD3E, and CD3G) as described in a previous study [41]. We examined the TOX expression level not only in TI CD8⁺ T cells but also in CD4⁺ T cells. Some portion of the normalized TOX expression in the T cells originates from CD4⁺ T cells, which cannot be negated by a marker-based normalization approach. However, if the TOX expression level in the T cells is positively correlated with that in the CD8⁺ T cells, the TOX level in all T cells would be proportional to that of CD8⁺ T cells. The analysis of single-cell transcriptome data from patients with melanoma [6] revealed a strong positive correlation between TOX expression level in all T cells and in CD8⁺ T cells (Fig. 5b). This enabled the evaluation of differences in the overall survival between cancer patients exhibiting varying TOX expression levels in the TI T cells. The low TOX expression level in the T cells was associated with enhanced overall survival rate (P = 0.0022, log-rank test) for TCGA melanoma cohort (SKCM, skin cutaneous melanoma) (Fig. 5c). Similarly, the survival analysis of TCGA NSCLC cohort (LUAD, lung adenocarcinoma, and LUSC, lung squamous cell carcinoma) revealed that a high overall survival rate was associated with low TOX expression level in the T cells (P = 0.0393, log-rank test) (Fig. 5d). A similar analysis was applied to other cancer types in TCGA cohort, which revealed that the survival rate of patients with bladder cancer (BLCA, bladder urothelial carcinoma), head and neck cancer (HNSC, head and neck squamous cell carcinoma), sarcoma (SARC, sarcoma), and uterine cancer (UCEC, uterine corpus endometrial carcinoma)
was associated with low TOX expression level in the TI T cells (Additional file 2: Figure S5). Notably, anti-PD-1 therapy has been approved by the US FDA (Food and Drug Administration) for treating bladder cancer and head and neck cancer. These results suggest that TOX expression levels in the TI T cells can be used to predict the overall survival of patients with cancer.

Next, we evaluated the power of TOX expression level in the TI T cells to predict the response to anti-PD-1 immunotherapy by analyzing the bulk RNA-seq data and clinical information in previously published reports regarding independent patient cohorts who underwent anti-PD-1 therapy. Interestingly, the TOX expression level in the TI T cells was inversely correlated with the anti-PD-1 immunotherapy response in two published cohorts of patients with cancer, i.e., melanoma cohorts [19] (Fig. 5e) and NSCLC cohorts [21] (Fig. 5f). A similar analysis was performed on our internal cohort of patients with NSCLC recruited from Yonsei Cancer Center in Korea, which revealed a similar inverse correlation between TOX expression level in the TI T cells and anti-PD-1 responses (Fig. 5g). In the two NSCLC cohorts, the TOX expression level in the TILs of responders was significantly lower than that in the TILs of non-responders ($P = 0.016$ and $0.002$, two-tailed Mann-Whitney $U$ test). Furthermore, we prioritized the patients exhibiting the lowest TOX expression level in the TI T cells. This stratification revealed high area under the receiver operating characteristic curve (AUROC) scores, which indicated that low TOX expression was a predictive marker for anti-PD-1 therapy response (Fig. 5h). The prediction of the anti-PD-1 response based on the TOX expression level in the TI T cells was not effective in another published cohort of patients with melanoma [20]. The relative abundance of TI T cells in this cohort was significantly lower than that in the three predictive cohorts (Additional file 2: Figure S6), which explains the poor predictive power of TOX expression in T cells for anti-PD-1 response. Therefore, we suggest that TOX expression level in the TI T cells can be a useful predictor for anti-PD-1 efficacy in human cancer. In summary, these results suggest that TOX expression level in the TI T cells can be used for patient stratification in cancer treatment, including anti-PD-1 immunotherapy.

**Discussion**

In this study, we identified the regulatory factors involved in TI CD8$^+$ T cell exhaustion by analyzing the scRNA-seq data using a method that mimics the subset analysis of flow or mass cytometry data. In contrast to conventional cytometry that can quantify a maximum of 50 pre-selected proteins, the scRNA-seq enables genome-wide expression analysis at the transcript level. The unbiased search for genes that are co-expressed with PDCD1, which is a major marker gene for T cell exhaustion, revealed not only IC genes but also TFs involved in T cell exhaustion, such as BATF, NFATC1, and PRDM1. The novel candidate genes reported in this study would be useful sources for future investigations on the molecular process underlying T cell exhaustion. A limitation of our prediction approach is that some genes that are not co-expressed with PDCD1, but are involved in the regulation of T cell exhaustion, may be missed. This limitation could be partially overcome by using additional marker genes for T cell exhaustion, such as TIM-3 and LAG-3.

Our results demonstrated the effectiveness of subset analysis using the scRNA-seq data to identify the regulatory molecules mediating cellular transitions across a continuous spectrum of transcriptional states. Recently, we published a web server, VirtualCytometry [42], which enables interactive subset analysis of public scRNA-seq datasets for human and mouse immune cells derived from various tissues, including tumors. Users can reproduce the gene prioritization for T cell exhaustion using the web server. The proposed method may be applied for studying cellular differentiation of various immune cells with appropriate marker genes.

Applying the subset analysis described in this study to the single-cell transcriptome profiles of CD8$^+$ T cells from human melanoma and NSCLC, which are currently the two most prevalent cancer types treated by anti-PD-1 therapy in the clinic, we could identify TOX as the top candidate TF for both cancer types. Previously, TOX was reported to regulate the development of CD4$^+$ T cells, natural killer cells, and lymphoid tissue inducer cells [43]. A series of recent studies reported that TOX promotes CD8$^+$ T cell exhaustion in chronic viral infection and cancer [30–32, 44–47]. Using the scRNA-seq data generated from human tumor samples, we also independently demonstrated that TOX promotes CD8$^+$ T cell exhaustion in human cancer. The analysis of expression dynamics in single-cell trajectories, and the flow cytometry analysis of expression correlation demonstrated that TOX is a more influential regulator of CD8$^+$ T cell exhaustion than other known factors in human cancer. Therefore, inhibition of TOX may potentially inhibit the cellular differentiation program in the generation of terminally exhausted T cells, thereby consequently enhancing the reinvigoration potential of progenitor exhausted T cells. To the best of our knowledge, this is the first study to demonstrate the feasibility of patient stratification for evaluating the anti-PD-1 therapy response based on the TOX expression level in TI T cells for multiple types of cancer (melanoma and lung cancer). Therefore, development of TOX-based biomarkers for prognosis of anti-cancer treatment and anti-PD-1 response can help in the improvement of cancer immunotherapy.
Previous studies have demonstrated that NFAT induces the expression of TOX, TOX2, and NR4A1 [29, 30]. The flow cytometric analysis revealed that there was a significant difference in the proportion of TOX⁺ cells among the TI CD8⁺ T cells exhibiting and not exhibiting PD-1 expression, which was not observed for NR4A1⁺ cells (see Additional file 2: Figure S1C). Consistently, there was no significant difference in NR4A1 levels between the effector cells and exhausted cells in the single-cell trajectory. However, the expression of NR4A1 was upregulated at later stages of the exhaustion process (see Fig. 2d). The cross-regulation between TOX and NR4A1 is unknown. TOX knockdown in the TI CD8⁺ T cells significantly downregulated the expression of IC molecules, such as PD-1, but not those of NR4A1 (Additional file 2: Figure S7). This indicated that TOX does not regulate NR4A1. Additionally, TOX and NR4A1 may independently contribute to T cell exhaustion. However, the possibility of NR4A1 regulating TOX as an upstream regulator cannot be ruled out. It would be interesting to investigate if TOX and NR4A1 can regulate each other at the transcriptional level, and to determine the more effective factor among the two with respect to T cell exhaustion.

One of the most common applications of single-cell transcriptome data is pseudotime analysis, which reconstructs the biological processes by exploiting the heterogeneity of cell population. Using highly heterogeneous CD8⁺ T cells in the tumor, we could successfully reconstruct the trajectories from effector state into either memory or exhausted state by pseudo-temporal ordering of cells in the low-dimensional latent space based on the transcriptome profiles of individual cells. Next, we divided the cells into effector, memory, and exhausted cells based on the branching point of the trajectories to confirm the expression dynamics of many known regulators of T cell exhaustion. The trajectories of the cells also enabled us to perform differential expression analysis between short pseudo-temporal stages, such as between late effector state and early exhausted state to identify regulators that upregulate or maintain TOX. A detailed analysis of the single-cell transcriptome data will facilitate the unraveling of the regulatory circuit involved in CD8⁺ T cell exhaustion.

Conclusions
To identify regulatory factors promoting T cell exhaustion in the tumor microenvironment, we prioritized TFs by differential expression between two subsets of TI CD8⁺ T cells divided by different levels of PDCD1 expression. We found that TOX levels positively correlate with those of IC molecules in human melanoma and NSCLC. The expression dynamics along the pseudo-temporally ordered CD8⁺ T cells from human tumors revealed that the expression of TOX increases with the exhaustion of CD8⁺ T cells. In addition, TOX knockdown in TI CD8⁺ T cells from human tumor reduced the number of cells expressing PD-1, TIM-3, TIGIT, and CTLA-4. These results indicate that TOX promotes T cell exhaustion via positive regulation of IC molecules. We also observe that TOX level in the TI T cells inversely correlates with overall survival and anti-PD-1 response in human melanoma and NSCLC, which suggest that TOX expression in the TI T cells can be used for patient stratification before anti-tumor immunotherapy, such as anti-PD-1 treatment. Taken all these results together, we propose TOX as a positive regulator for T cell exhaustion and a predictor for anti-PD-1 responses in human cancer. TOX inhibition may potentiallypede T cell exhaustion and improve ICI efficacy.

Supplementary information
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Authors’ contributions
KK, SJH, and IL designed the study. KK performed the bioinformatics analysis of single-cell and bulk omics data. SP performed the experimental analysis. SYP, DHK, YMP, and YWK provided the clinical samples and clinical data. GK supported the bioinformatics analysis. HRK provided the bulk RNA sequencing data and advice for this project. SJH and IL supervised the project. KK, SP, SJH, and IL wrote the manuscript. The authors read and approved the final manuscript.

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Availability of data and materials
Single-cell transcriptome data of tumor samples for melanoma and NSCLC are available from Gene Expression Omnibus (GEO) database, GSE72056 [6] and GSE39925 [7], respectively. The bulk RNA sequencing data for the cohort of 16 patients with NSCLC who received anti-PD-1 therapy are also available from the GEO database (GSE126044).

Ethics approval and consent to participate
The studies were approved by the Institutional Review Board of Yonsei University Severance Hospital with IRB no. 4-2016-0678 for the 16 NSCLC patients undergoing anti-PD-1 treatment and IRB no. 4-2016-0788 for the 35 patients with NSCLC and 15 patients with HNSCC. All patients who participated in these studies provided written informed consent prior to enrolment and sample collection at Yonsei University Severance Hospital. The research conformed to the principles of the Helsinki Declaration.

The mice were maintained in accordance with the Laboratory Animals Welfare Act, the Guide for the Care and Use of Laboratory Animals, and the Guidelines and Policies for Rodent Experiments proposed by the Institutional Animal Care and Use Committee (no. IACUC-A021809-78-01) of the Yonsei University Health System.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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