Figure S1. Result of the PCA checking gender distribution
The PCA using probes on the X chromosome was performed to examine whether the samples were correctly labeled with the presumed gender. Red dots indicate female samples and black dots indicate male samples. All samples were classified into the presupposed gender groups.
Figure S2. The density plot of each sample’s β values (1/4)
Figure S2. The density plot of each sample’s β values (2/4)
Figure S2. The density plot of each sample’s β values (3/4)
Figure S2. The density plot of each sample’s $\beta$ values (4/4)
**Figure S3.** Correlation between DNAm age and chronological age

DNAm age was predicted using the results of the methylation array. Red and black dots indicate each PD and healthy control subjects respectively. Pearson’s correlation coefficients between DNAm age and chronological age were calculated as 0.77 and 0.90 in each group.
Figure S4. The estimated abundance measures of plasmablasts, CD8^+CD28^-CD45RA^- T cells, naive CD8^+ T cells, and naive CD4^+ T cells. The abundance measures were estimated using the results of the DNA methylation array. Wilcoxon rank-sum tests using the estimated abundance measures were performed between the PD and control subjects. P values are indicated in blue characters. Significance level after the Bonferroni correction was set as $\alpha = 0.005$. 
Figure S5. Filtering procedure of the methylation array data

\( \beta \) values with detection P value \( \leq 0.01 \) were treated as missing values and then probe call rates were determined. Low quality probes were filtered out applying the following criteria; (1) probe call rate \( > 95\% \), (2) probes not on chromosome (Chr.) X and Y, (3) probes not including SNPs with MAF \( \geq 0.05 \), (4) probes not reported as cross-reactive. Finally 376,602 probes remained and were examined in the following analyses.
Figure S6. Q-Q plot of regression analysis
The observed P values are those of regression analysis adjusting the effects of the predicted proportions of leukocyte subsets.
Figure S7. The M value distributions of the significant CpG sites between smokers and non-smokers in PD subjects (1/2)
Figure S7. The M value distributions of the significant CpG sites between smokers and non-smokers in PD subjects (2/2)
Remove β values with detection P value ≥ 0.01

Remove probes with call rate ≤ 95%

476,129 / 485,512 probes

Remove probes other than on chr X

10,832 / 476,129 probes

Remove probes which include SNPs with MAF ≥ 0.05

10,314 / 10,832 probes

Remove probes which are reported to be cross-reactive
(Chen et al., 2013)

9,612 / 10,314 probes

**Figure S8.** Filtering procedure of probes on X chromosome

β values with detection P value < 0.01 were treated as missing values and then probe call rates were determined. Low quality probes were filtered out applying the following criteria; (1) probe call rate > 95%, (2) probes on chromosome (Chr.) X, (3) probes not including SNPs with MAF ≥ 0.05, (4) probes not reported as cross-reactive. Finally 9,612 probes remained and were examined in the following analyses.