Supplementary Figures S1-S14

Figure S1. PCA plots of the HSPC dataset A) no preprocessing, B) after application of netSmooth, C), using scImpute, and D) after application of MAGIC.
**Figure S2.** t-SNE plots of the HSPC dataset A) no preprocessing, B) after application of netSmooth, C) using scImpute, and D) after application of MAGIC.
Figure S3. Single cells from the embryonic development dataset were clustered using the robust clustering procedure, and the log-transformed expression values of the 500 most differentially expressed genes (by edgeR-QLF test adjusted P value) in any of the discovered clusters are shown in a heatmap, as well as cluster assignments and cell types. A) raw (no imputation), B) after application of netSmooth, C) missing values imputed using scImpute D) after application of MAGIC.
Figure S4. t-SNE plots of the embryonic development dataset A) no preprocessing, B) after application of *netSmooth*, C), using scImpute, and D) after application of MAGIC.
Figure S5. The proportion of genes with 0 counts is a proxy for technical dropouts. A) no preprocessing, B) after application of netSmooth, C), using scImpute, and D) after application of MAGIC.
Figure S6. Single cells from the glioblastoma dataset were clustered using the robust clustering procedure, and the log-transformed expression values of the 500 most differentially expressed genes (by edgeR-QLF test adjusted P value) in any of the discovered clusters are shown in a heatmap, as well as cluster assignments and cell types. A) raw (no imputation), B) after application of netSmooth, C) missing values imputed using scImpute, D) after application of MAGIC.
Figure S7. PCA plots of the glioblastoma dataset A) no preprocessing, B) after application of netSmooth, C), using scImpute, and D) after application of MAGIC.

Figure S8. Cluster purity by smoothing parameter. A) for the hematopoiesis dataset with a directional (signed) graph, where inhibitory interactions have a negative edge weight. B) For the hematopoiesis dataset using a gene network with only genes that have a cell-type specific expression in any cell type. C) In the glioblastoma dataset using a gene network from HumanNet.
Figure S9. Cluster purity including CIDR. Same as Figure 3 with CIDR included.

Figure S10. Adjusted mutual information including CIDR. Same as Figure 3 with CIDR included.

Figure S11. Cluster purity including CIDR. Same as Figure 5 with CIDR included.
Figure S12. Adjusted mutual information including CIDR. Same as Figure 5, with CIDR included.

Figure S13. Cluster purity including CIDR. Same as Figure 7, with CIDR included.

Figure S14. Adjusted mutual information including CIDR. Same as Figure 7, with CIDR included.