

APPLICATION NOTES

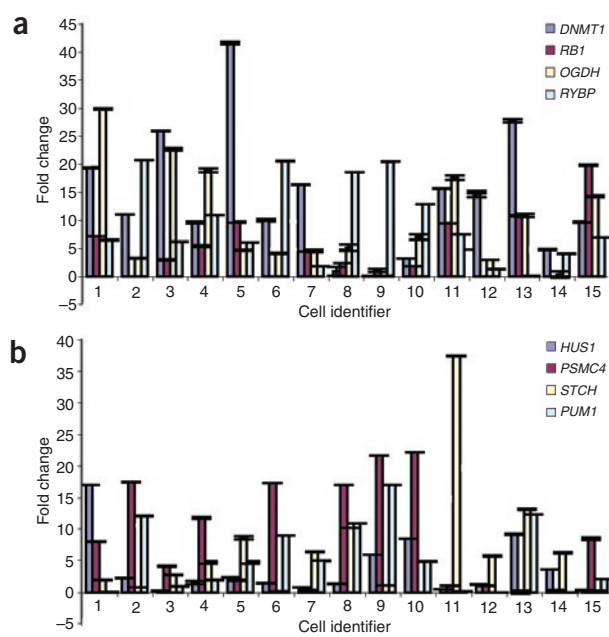


Figure 2 | Fold changes in expression of individual genes. **(a,b)** Δ Ct values were calculated for different sets of genes using the *APC* gene as a reference, and those values were then converted to fold change values for each of the 15 cell replicates. Error bars, standard deviations based on the Ct standard deviation of the three technical replicates converted to fold change.

to the gene of interest and a fluorescent resonance energy transfer (FRET) hydrolysis probe located between the two primers) were loaded into a separate set of 48 inlets.

The workflow is a simple, five-step procedure: prime, transfer, load, run and analyze. You begin by priming the dynamic array to close the interface valves, which prevents premature mixing of samples and assays. This is as simple as choosing a menu item on the touch-screen of the loader. After the chip has been 'primed', then you pipette your samples—premixed with master mix—into separate sample inlets of the dynamic array. The primer/probe sets are pipetted into separate inlets on the chip. You place the dynamic array on the IFC Controller and the software directs the pressure-loading of the assay components into the reaction chambers. The assay components are automatically combined on-chip. You then place the dynamic array on the Fluidigm Real-Time PCR System for thermal cycling and fluorescence detection. Once the thermal cycling is complete, the real-time qPCR Analysis software provides amplification curves, color-coded heat maps (**Fig. 1**) and cycle threshold (Ct) data for each run.

Our results (**Fig. 2**) showed that there was meaningful variation (defined as more than tenfold) in gene expression among the 15 individual cells from the same stage of early embryonic development.

Conclusion

Using Fluidigm's Dynamic Array for single-cell gene expression analyses is inexpensive and easy to use, giving researchers the ability to test a large number of cells and genes at the same time with data quality rivaling benchmark real-time qPCR results.

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