Supplementary Figure 6 WaferGen SmartChIP QRTPCR verification of the most divergent genes identified as DEG by RNA-Seq. Official gene symbols are indicated in upper left corner with the number of RNA-Seq aligned reads in parentheses ( ) and number of samples with C\textsubscript{T} values lower than background in brackets [ ] for vehicle control samples. Bars represent mean fold-change determined by WaferGen technology (±SEM), the red line represents RNA-Seq fold-change, and the green line represents Agilent fold change. Significant differences within WaferGen data were determined by one-way ANOVA followed by Dunnett’s post-hoc test and indicated by an asterisk (*). Red (RNA-Seq) and green (Agilent) dots represent P1(f) values with size indicating level of significance (small ~0.8, large ~1). Labels on the X-axis indicate the dose of TCDD (µg/kg), number of aligned RNA-Seq reads, and number of samples with C\textsubscript{T} values lower than background. Dashed lines indicate 1.5 and 2.0 [fold-change] thresholds to identify DEGs.