From transient transcriptome responses to disturbed neurodevelopment: role of histone acetylation and methylation as epigenetic switch between reversible and irreversible drug effects

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Abstract The superordinate principles governing the transcriptome response of differentiating cells exposed to drugs are still unclear. Often, it is assumed that toxicogenomics data reflect the immediate mode of action (MoA) of drugs. Alternatively, transcriptome changes could describe altered differentiation states as indirect consequence of drug exposure. We used here the developmental toxicants valproate and trichostatin A to address this question. Neurally differentiating human embryonic stem cells were treated for 6 days. Histone acetylation (primary MoA) increased quickly and returned to baseline after 48 h. Histone H3 lysine methylation at the promoter of the neurodevelopmental regulators PAX6 or OTX2 was increasingly altered over time. Methylation changes remained persistent and correlated with neurodevelopmental defects and with effects on PAX6 gene expression, also when the drug was washed out after 3–4 days. We hypothesized that drug exposures altering only acetylation would lead to reversible transcriptome changes (indicating MoA), and challenges that altered methylation would lead to irreversible developmental disturbances. Data from pulse-chase experiments corroborated this assumption. Short drug treatment triggered reversible transcriptome changes; longer exposure disrupted neurodevelopment. The disturbed differentiation was reflected by an altered transcriptome pattern, and the observed changes were similar when the drug was washed out during the last 48 h. We conclude that transcriptome data after prolonged chemical stress of differentiating cells mainly reflect the altered developmental stage of the model system and not the drug MoA. We suggest that brief exposures, followed by immediate analysis, are more suitable for information on immediate drug responses and the toxicity MoA.

Keywords Valproic acid · Embryonic stem cell · Histone deacetylase · Developmental toxicity · Histone modification

The authors wish it to be known that, in their opinion, the first two authors should be regarded as joint First Authors.

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Introduction

Toxicogenomics data, systems biology, and the use of human stem cell-based systems are expected to change the ways by which toxicological information will be obtained and interpreted in the future (Hartung et al. 2012; Robinson et al. 2012a; Robinson and Piersma 2013; Waters and Fostel 2004; Wobus and Loser 2011). This is in line with a ‘proposed shift from primarily in vivo animal experimentation to in vitro assays and computational modeling for toxicity assessment,’ as suggested by the lead scientists of US national research agencies (Collins et al. 2008). It also follows the ‘vision for a new toxicology of the twenty-first century’ as promoted by the National Research Council (Andersen and Krewski 2010; NRC 2007). A key assumption for this vision is that it will be possible to define pathways of toxicity, i.e., a drug mode of action (MoA), linking molecular initiating events to a final adverse outcome. This requires an establishment of a ‘systems toxicology’ that models the pathophysiology of the body with computational tools to understand mechanisms of toxicity, similar to systems biology (Hartung et al. 2012).

In the field of developmental toxicology, transcriptome data have been used to infer information on the MoA of chemicals (Colleoni et al. 2011; Hermsen et al. 2013; Vojnits et al. 2012). It is expected that such approaches will lead to major conceptual advances, especially for the use of the emerging technology of differentiating stem cell systems (Crofton et al. 2012; Theunissen et al. 2012; Zimmer et al. 2011). However, more fundamental work is required to understand how experiments need to be designed and interpreted in this field. In contrast to mature tissues or cells, model systems of development do not have a stable baseline, i.e., the transcriptome changes over time, also without toxic stimulus. Moreover, initial exposure to a toxicant may trigger secondary effects even in the absence of the stimulus (Balter et al. 2012). To avoid indirect effects in toxicogenomics measurements, sampling only a few hours after compound exposure has been suggested (Jergil et al. 2011). However, the sensitivity and response to toxicants of a dynamically differentiating system can be different at different times (van Dartel et al. 2009). For instance, in a study of retinoic acid teratogenicity, compound-induced transcriptome changes differed between sampling time points (Robinson et al. 2012b). Such timing effects of toxicants may be due to interference with specific waves of gene regulation. For instance, neurally differentiating mESC showed several of such waves of gene expression, which determined windows of sensitivity to toxicants (Abrances et al. 2009; Zimmer et al. 2011).

Alteration of histone deacetylase (HDAC) activity has been associated with several long-term health consequences, ranging from Alzheimer’s disease (Graff et al. 2012), over toluene poisoning (Sanchez-Serrano et al. 2011) to general teratogenic mechanisms (Menegola et al. 2012). Specific HDAC inhibitor (HDACi) drugs have been particularly well characterized. For instance, the broad-spectrum HDACi valproic acid (VPA), normally used to treat epilepsies, causes the fetal valproate syndrome and has been suspected to trigger autism (Dufour-Rainfray et al. 2010; Jentink et al. 2010; Meador et al. 2009). It triggers gene activation within few hours in several tumor or stem cell lines (Jergil et al. 2009, 2011), and it is well recognized that epigenetic modifications are related to the developmental neurotoxicity of the drug. Most studies addressing the latter mechanism have concentrated on transcription-activating histone modifications such as acetylation of histone 3 at lysine 9 (H3K9Ac) or methylation of histone 3 at lysine 4 (H3K4me) (Hezroni et al. 2011; Marinova et al. 2011; Tung and Winn 2010), and the changes have been found to be reversible after drug withdrawal (Boudadi et al. 2013).

In previous work, we established a model of early neural differentiation of hESC that allowed the identification of developmental toxicants (Balter et al. 2012; Krug et al. 2013). We found that prolonged exposure to the two
HDACi trichostatin A (TSA) or VPA altered the expression of several marker genes in a similar way. However, we also observed that various hESC-based test systems differed strongly in their transcriptome response to VPA (Krug et al. 2013). This latter finding triggered the key question of this study: Are the observations on altered transcriptome patterns in developmental toxicity studies indeed a reflection of a compound’s primary MoA? As alternative hypothesis, we examined whether the data rather reflect an altered cellular phenotype that would result from disturbed differentiation and that would become independent of the continued presence of drug after some time. This was addressed by transcriptome analysis after pulsed drug exposure. A further key question was how a direct, but reversible effect of short
toxicant exposure was switched to a persistent adverse effect, reflected by wrong differentiation after a longer drug exposure. This was addressed by studies of the time dependence of histone modifications. Histone methylations at the promoters of key neurodevelopmental genes were considered as potential persistence detectors responsible for switching a short-term cellular adaptation to permanent toxicity.

Materials and methods

Materials

Gelatine, putrescine, selenium, progesterone, apotransferin, glucose, insulin, valproic acid, and trichostatin A were obtained from Sigma (Steinheim, Germany). Accutase was from PAA (Pasching, Austria). FGF-2 (basic fibroblast growth factor), noggin, and sonic hedgehog were obtained from R&D Systems (Minneapolis, MN, USA). Y-27632, SB-43154, and dorsomorphin dihydrochloride were from Tocris Bioscience (Bristol, UK). MatrigelTM was from BD Biosciences (Massachusetts, USA). All cell culture reagents were from Gibco/Invitrogen (Darmstadt, Germany) unless otherwise specified.

Neuroepithelial differentiation

Human embryonic stem cells (hESC) (H9 from WiCells, Madison, WI, USA) were differentiated as described in detail earlier (Chambers et al. 2009). Briefly, dual SMAD inhibition was used to prevent BMP and TGF signaling and thus to achieve a highly selective neuroectodermal lineage commitment. For handling details, see supplemental methods of Balmer et al. (2012). If not stated otherwise, treatment with trichostatin A (TSA) was done with a concentration of 10 nM and treatment with valproic acid (VPA) was done with a concentration of 600 µM.

Quantitative real-time PCR (qPCR) and microarray analysis

For qPCR analysis, cells were lysed at indicated days of differentiation in TriFast™ (Peqlab, Germany). Total RNA was isolated according to the manufacturer’s instruction, and cDNA was produced using the iScript Kit from BioRad (iScript™ Reverse Transcription Supermix for RT-qPCR, BioRad). Quantitative real-time PCR (qPCR) was performed on a BioRad Light Cycler (Biorad, München, Germany), and transcript levels were quantified as described earlier (Balmer et al. 2012). The sequences of specific primers are given in Fig. S11.

Affymetrix chip-based DNA microarray analysis (Human Genome U133 plus 2.0 arrays) was performed as described earlier (Krug et al. 2013). The data were analyzed for differential expression using the Konstanz Information Miner open source software [KNIME; www.knime.org (Berthold et al. 2007)]. The raw data were preprocessed using robust multiarray analysis (RMA) (Smyth 2005). Background correction, quantile normalization, and summarization were applied to all expression data samples, using the RMA function from the affy package of Bioconductor (Gautier et al. 2004; Gentleman et al. 2004). The limma package (R & Bioconductor) was used to identify differentially expressed genes using indicated groups as control. The moderated t statistics was applied in a pairwise fashion (each treatment was compared to its own control) and was used for assessing the raw significance of differentially expressed genes. Then, final p values were derived using the Benjamini–Hochberg (BH) method to control the false discovery rate (FDR) (Benjamini and Hochberg 1995) due to multiple hypothesis testing. Transcripts with FDR adjusted p value of ≤0.05 and fold change values >1.5 or <2/3 were considered significantly regulated, if not stated otherwise in the figure legend. For Fig. S5, numbers of PS changed during development (D-genes) were calculated relative to hESC. These data were obtained from four independent replicates, and they were considered significant if the Benjamini–Yekutieli (BY)-adjusted p value was <0.01 and the FC was >1.5 or <2/3. For Figs. 3, 4, 5, and 6, numbers of PS changed by the treatment were calculated relative to untreated controls lysed at the same day as the treated samples (T6h to C6h, T4d to C4d and T6d, early pulse (EP), medium pulse (MP), and late pulse (LP) to C6d). Data were obtained from four independent replicates and chosen if the BH-adjusted p value was <0.05 and FC was >1.5 or FC <2/3.

The principal component analysis (PCA) was based on 500 PS with the highest variance.

Western blot

Western blot was performed exactly as previously described (Balmer et al. 2012). For quantification, the signal intensity of H3Ac was normalized to total H3, and acetylated α-tubulin was normalized to total α-tubulin for every time point. These normalized values were then displayed relative to untreated controls at the respective time points. Western blots of PAX6 and OTX2 were quantified using ImageJ. Detailed information on used antibodies is given in Fig. S12.

Chromatin immunoprecipitation (ChIP)

Chromatin immunoprecipitation (ChIP) assays on native chromatin (N-ChIP) (Fig. 1) were performed according to established protocols (Umlauf et al. 2004). Details and...
Fig. 2 Consequences of different drug washout periods for gene expression and histone methylation patterns. For all experiments, hiESC were differentiated to NEP. a Samples for chromatin immunoprecipitation (ChIP) were prepared at the indicated days of differentiation. ChIP was performed with antibodies specific for H3K4me3 or H3K27me3 or control IgG. The enrichment factors of OTX2 and PAX6 promoter sequences are given as % input for H3K4me3 (dark blue) and H3K27me3 (light blue). Data are mean ± SEM of three independent cell preparations. b Differentiating cells were treated with TSA (10 nM) for the indicated time periods, and ChIP was performed with the same antibodies as described in a. The ratio of enrichment factors of H3K4me3 and H3K27me3 was calculated as measure of chromatin opening. Data are given relative to values of untreated control cells at the same time point (n = 3). c Scheme of experimental treatment and sampling for the following experiments. Gray bars indicate the period of drug exposure (e.g., P2d: pulsed drug treatment for 2 days) with 10 nM TSA, and white bars indicate medium without TSA. All samples were analyzed on day 6 of differentiation for each treatment scenario. d Protein levels of PAX6 or OTX2 were determined by Western blot, and relative (vs ctr.) protein levels (n = 3) were quantified. e Transcript levels of pAx6 and OTx2 were determined. They are expressed relative to untreated control on DoD6 (ctr). f ChIP was performed for H3K27me3 (purple) or H3K4me3 (black) on promoter regions of PAX6 and OTX2, and enrichment factors were calculated relative to ChIP with control IgG. Then, these data were normalized to the values obtained for control cells (ctr). For instance, on day 6 of the differentiation, H3K27me3 was 15-fold higher in cells treated for 1 day with TSA and then left in control medium (p1d), compared to cells that were differentiated under control conditions. Data of d–f are mean ± SEM of 3–5 experiments (color figure online).
adaptations were exactly as described previously in detail (Balmer et al. 2012). ChIP assays on cross-linked chromatin (X-ChIP) (Fig. 2) were performed according to Kamieniarz and colleagues and adapted to our differentiating cells (Kamieniarz et al. 2012). Briefly, cells were trypsinized and resuspended in 1% formaldehyde in medium. The cross-link was stopped after 10 min by 125 mM Tris, pH 7.5. Cellular suspensions were centrifuged, washed once in PBS and once in L1 buffer (2 mM EDTA, 0.1% NP-40, 10% glycerol, 25 mM Tris, pH 8), and finally resuspended in L2 buffer (10 mM EDTA pH 8, 1% SDS, 50 mM Tris, pH 8) to a final concentration of 2 × 10^6 cells/ml. Chromatin was sonicated on a Bioruptor® sonifier device (Diagenode, Belgium) by 30 steps of 30/30 s ON/OFF cycles to get a fragment size between 300 and 700 bp, and sonication efficiency was checked on agarose gels. Samples were diluted
1:5 in dilution buffer (0.5 % NP-40, 200 mM NaCl, 50 mM Tris, pH 8) and incubated over night at 4 °C with unspecific control antibody, 2 µl anti-H3K4me3 (17–614 Millipore) or 4 µl anti-H3K27me3 (39535 Active Motif) antibodies. One aliquot, corresponding to 5 % of the input, was stored without antibody treatment. After antibody incubation, the samples were rotated at 4 °C for 3 h with protein A/g Sepharose beads and washed twice in washing buffer (2 mM eDTA, 0.1 % SDS, 0.5 % Np-40, 150 mM NaCl, 20 mM Tris pH 8) and once in final wash buffer (2 mM eDTA, 0.1 % SDS, 0.5 % Np-40, 500 mM NaCl, 20 mM Tris, pH 8). The chromatin was eluted by 2-h incubation and shaking at 65 °C in elution buffer (100 mM NaHcO3, 1 % SDS). The genomic DNA was purified using ChIp DNA clean and Concentrator (Zymo Research) Kit and analyzed by qPCR, to quantify the amount of DNA from the promoter region of selected genes. For data display, the enrichment factor (eF) was calculated from the qPCR threshold cycle values (ct) relative to input according to the formula:

\[ eF (\%) = 100 \times 2^{(ct(5 \% \text{ IP}) - 4.32) - ct(\text{specific antibody})} \]

For Figs. 1e and 2a, we wanted to investigate the effects of TSA or VPA in comparison with untreated controls. We compared the ratios of H3K4me3/H3K27me3 of treated cells to the ratios of H3K4me3/H3K27me3 of untreated cells at the respective days of differentiation. Also for transcript levels, the gene expression was presented relative to untreated controls at the respective days. Therefore, H3K4me3/H3K27me3 ratios or gene expression above 1 indicated that TSA or VPA caused an up-regulation compared with untreated control. H3K4me3/H3K27me3 ratios (methylation ratio) or gene expression below 1 indicated that TSA or VPA caused a down-regulation compared with untreated controls. For detailed information on ChIP, primers and antibodies refer to Fig. S11 and S12.

Statistics and data mining

For statistical analysis of transcript levels and EFs, paired t-tests were performed using log-transformed expression values relative to hESC, if not stated otherwise in the legend. All data are shown, and all statistics performed refer to biological replicates (= independent experiments).
Over-representation of gene ontologies (GOS) was analyzed using g:profiler (Reimand et al. 2011), with p values determined via a hypergeometric distribution. Over-represented GOS were selected, if they belonged to the term domain ‘biological process’ and contained <1,000 genes, and the p value was <0.05. For analyses yielding more than 50 GOS, more stringent selection criteria were used: only GOS that had a p value <0.001 were selected. For production of the GO word clouds scaling of character size was linearly proportional to the negative logarithm of the p value of the respective GO category. GO terms relating to biological processes (bp) were clustered according to their ‘superordinate biological processes’ as described earlier. Example of these larger categories were ‘neuronal differentiation,’ ‘non-neuronal differentiation,’ or ‘migration and adhesion’ (Waldmann et al. 2014).

Venn diagrams were drawn in order to visualize size relations between the compared groups of genes within one diagram. They do not always represent correct ratios, as this would make visualization difficult in case of big size differences. Numbers in Venn diagrams comparing three groups represent the percentage of the (overlapping or unique) part of the diagram relative to samples lysed at DoD4. The corresponding absolute numbers are indicated in the supplementary files. For Venn diagrams with two circles, absolute numbers of pS and their overlap are presented. The numbers that indicate the percentage of the overlap in two group comparisons are relative to the circle that has the same color as the line under the number.

Results

Time courses of histone acetylation and altered marker gene expression triggered by HDACi during early neurogenesis

A pure population of neuroepithelial cells (NEP) can be generated from human embryonic stem cells (hESC) within 6 days (Balmer et al. 2012; Chambers et al. 2009; Krug et al. 2013). This model system is characterized by up-regulation of the transcription factors PAX6 and OTX2 (Fig. 1a). We found earlier that continued exposure to TSA and VPA affects the expression of these two genes. In addition, we observed that short drug exposure (for 24 h) showed the expected biochemical effect [increased histone acetylation on day-of-differentiation 1 (DoD1)]. However, when differentiating hESC, treated for the first 24 h
with HDACi, were examined on day-of-differentiation 6 (DoD6), there was no effect on the expression of the neuroepithelial markers (p Ax6 and OTx2) (Balmer et al. 2012; Chambers et al. 2009; Krug et al. 2013). To better understand the relationship of biochemical changes and the expression of differentiation markers, we started this study by examining the time course of key events including gene expression and various histone modifications.

Analysis of mRNA expression of p Ax6 during undisturbed differentiation showed that this gene is very little regulated during the first 3 days, while significant OTx2 up-regulation was already detectable after 2 days. Both genes reached high levels (compared to hESC) on DoD6. Exposure to HDACi (equipotent concentrations of 600 µM VPA or 10 nM TSA) during the entire differentiation process led to a relative down-regulation of both NEP marker genes. Consistent with the time course of developmental up-regulation of the marker genes, the drugs affected OTX2 already from early time points on, while PAX6 levels were only reduced at DoD4–6 (Fig. 1c).

To examine protein acetylation triggered by drug treatment, we used Western blotting. Histone H3 and α-tubulin were selected as abundant and well-characterized target proteins of HDACs. Exposure to TSA or VPA for the first 6 h of differentiation triggered strong acetylation of histone H3 on the whole-cell level. TSA also increased acetylation of α-tubulin (p < 0.05), as expected from the HDAC inhibition profile. VPA effects on this target were not significant, which is consistent with the fact that VPA inhibits specifically HDAC class I enzymes and not the tubulin acetylating class II HDACs (Fass et al. 2010; Gottlicher et al. 2001; Khan et al. 2008). Already after 24 h, the extent of protein acetylation was strongly reduced compared with 6-h drug treatment, and after 48 h, the effect vanished (despite the continued presence of the drugs) (Fig. 1d).

This was most likely due to cellular counter-regulations, as HDACs and histone acetyl transferases are dynamically regulated during differentiation (Weng et al. 2012) and adaptive effects have been described for HDACi drug treatment (Kataoka et al. 2013; Tung and Winn 2010).
Nevertheless, we also considered that global analysis of protein acetylation may not be sensitive enough. Therefore, we studied histone H3 acetylation of the lysine-27 residue (H3K27Ac) at four promoter sites of interest (transcription start site of PAX6, OTX2, OCT4, Nanog) using chromatin immunoprecipitation (ChIP). No drug-induced changes in the acetylation levels were observed on DoD1 or any of the following days (Fig. S1A). Various control experiments showed that histone acetylation was technically measurable in our cells and that the H3K27 residue can be affected by HDACi (at higher concentrations) at the chosen promoter sites (Fig. S1B). These data suggest that the low, human-relevant developmental toxicant drug concentrations used here triggered at best very weak, non-measurable promoter acetylation. A more persistent effect on gene regulation and differentiation may thus require another histone modification, such as methylation.

Time course of histone methylation changes triggered by HDACi during early neurogenesis

We had found earlier that a 6-day continued treatment with TSA resulted in alterations of histone methylation, but it remained unclear when such changes happened and how persistent they were (15). As a 4-day treatment with subsequent washout of the drug was sufficient to alter the expression of PAX6 (Balmer et al. 2012; Chambers et al. 2009; Krug et al. 2013), we investigated now in a first pilot experiment, whether the histone methylation pattern was already changed after 4 days and whether such changes would persist throughout a drug washout period. ChIP was performed with antibodies specific for the open/active chromatin mark trimethylated lysine 4 of histone 3 (H3K4me3) and with antibodies specific for the closed/inactive chromatin mark trimethylated lysine 27 of histone 3 (H3K27me3) (Fig. S2). Ratios of H3K4me3 to H3K27me3 enrichment were calculated as a simple measure of the chromatin state [high ratio: open promoter and low ratio: rather silenced promoter (Burney et al. 2013)] (Fig. 1e). As expected, we found the methylation ratio of GAPDH, an actively described gene affected neither by TSA nor by differentiation, to be very high and to remain unchanged. We also confirmed our earlier finding that PAX6 and OTX2 have a high methylation ratio in normal NEP (DoD6) and that TSA strongly reduced this ratio ($p < 0.001$). Now, we found that this effect also held true for DoD4 cells. Moreover, the TSA-induced reduction in the methylation ratio was persistent, when the drug was washed out from DoD4 to DoD6 ($p < 0.001$) (Fig. 1e, Fig. S2). Thus, altered histone methylation patterns correlated with drug effects on neurodifferentiation, and they may play a role in drug-induced developmental toxicity and persistent effects of HDACi.

Therefore, we examined the time course of histone methylations more closely. During normal development, H3K4me3 levels of PAX6 remained relatively constant, while promoter opening was indicated by a decrease in H3K27me3, especially between DoD4 and DoD6 (Fig. 2a). Treatment with TSA reduced this late decrease in the inactivating histone modification (thereby reducing PAX6 transcription) (Fig. 2b, Fig. S3). Opening of the OTX2 promoter during neurodifferentiation was indicated by early (DoD1–2) increases in H3K4me3 and simultaneous decreases in H3K27me3 (Fig. 2a). Upon treatment with TSA, the methylation ratio at the OTX2 promoter was already slightly down-regulated at DoD1 and significantly down-regulated from DoD2 on ($p < 0.05$) (Fig. 2b, Fig. S3). Thus, we found here that prolonged treatment with an HDACi can lead to an enrichment of inactivating histone modifications. This offers a mechanistic explanation for the down-regulation of important developmental genes (here PAX6 and OTX2) by TSA. These findings differ largely from those reported on short treatment (Bouddadi et al. 2013; Jergil et al. 2011; Nightingale et al. 2007), which increases the amount of chromatin-opening histone modifications. On this basis, it became highly interesting to obtain more data on the persistence of altered histone methylations upon pulsed drug treatment.

Effects of pulsed drug treatment on histone methylation and NEP differentiation

After we had found that the increase in H3K27me3 as well as the down-regulation of PAX6 and OTX2 was persistent when the cells were treated for 4 days, followed by a 2-day wash out (Fig. 1e), we investigated the minimum treatment period required to induce this stable effects.

The cells were exposed to TSA for 1, 2, or 3 days, before the drug was removed, and differentiation was continued until DoD6. These pulsed treatments (P1d, P2d, and P3d) were compared to continuous exposure to TSA (Fig. 2c). As controls of the phenotypic effect of disturbed differentiation, we quantified the protein levels of PAX6 and OTX2. Short treatments of up to 2 days had no effect, while longer drug treatment resulted in strongly decreased levels of the phenotype markers of NEP (Fig. 2d). Thus, drug treatment of about 3 days was sufficient to cause developmental disturbances.

In parallel, we examined how the pulsed treatment affected the gene expression and promoter methylation of the NEP marker genes. As seen for the protein, 3-day treatment was sufficient to cause the same extent of down-regulation as continuous drug exposure (Fig. 2e). Examination of H3K4me3 and H3K27me3 marks showed that 3-day drug treatment, followed by 3-day washout...
transcriptome changes should be similar for T6d treatment as initial experiments (Figs. 1, 2), we concluded that overall development marker expression. From this and from our drug treatment was not important anymore for neurode-
mental toxicity character of the model. The up-regulated genes strongly indicated differentiation to non-neuronal lineages (Fig. 3e, Tab. S3). Therefore, we conclude that long- but not short-term treatment indeed changes the gene expression pattern toward a wrong (non neuronal) differen-
track.

The divergence from neuronal differentiation to non-
neuronal lineages (according to oGOs) developed very strongly between T4d and T6d, i.e., during the time, when drug treatment was not important anymore for neurode-
velopmental marker expression. From this and from our initial experiments (Figs. 1, 2), we concluded that overall transcriptome changes should be similar for T6d treatment and a medium drug pulse of 4-day exposure followed by a 2-day washout (MP).
Transcriptome changes gain independence of drug presence after 4 days

We addressed the question of continued transcriptome effects in the absence of drugs by comparing T6d and MP (Tab. S2). The regulated PS showed a very high overlap of 80–87 % between MP and T6d (Fig. 5b). The oGOs among up-regulated PS pointed to the differentiation of several other cellular lineages (Fig. 4b, Tab. S4), such as the cardiovascular system, neural crest, skeletal system, and glands. Moreover, there was a high overlap of oGOs among T6d PS and the T6d/MP overlap PS (Fig. 4b). The higher GO enrichment among the latter set of genes was most likely due to the lower percentage of non-specific genes, eliminated through the overlap filter. The commonly down-regulated PS gave overwhelming evidence of disturbed neurodifferentiation (according to oGOs) (Fig. 4c, Tab. S4). We wondered whether the biological response to the drugs could have been predicted already on day 4. At this time (T4d), the majority [64 % (up)/55 % (down)] of TSA-regulated PS was the same as found after 6 days (Fig. S7A). The PS of T4d that overlapped with T6d and MP pointed already to disturbed differentiation: the oGOs among these down-regulated PS indicated a defect in forebrain development (Fig. S7B) and the up-regulated PS pointed again to an increase in unwanted differentiation tracks (Tab. S4). In summary, drug treatment for 4 days was sufficient to trigger the definite deviation from the normal differentiation path. From this point on, differentiation of wrong lineages most likely contributed to further increases in transcriptome changes (compared to control cells). This wrong differentiation track continued to deviate more and more from control cells, also in the absence of the drugs.

Comparison of continued drug exposure and short exposure of NEP

The independence of the transcriptome response from the presence of drugs suggested that the transcriptome changes did not inform on direct drug signaling. To address how the cells would react to short drug exposure, we generated NEP and exposed them to TSA only during the last 6 or 24 h of the 6-day differentiation. This treatment caused a pronounced increase in histone acetylation and tubulin acetylation (about fivefold to eightfold, p < 0.05) (Fig. 5a). These results showed that HDACi could trigger the expected biochemical response (acetylation) in NEP acutely exposed to the drugs. This effect was similar in size as the one in cells pulsed with drug during the first 6 h of differentiation (Fig. 1d). A further similarity was that also the late drug pulse (LP) triggered a pronounced transcriptome response within 6 h. However, the regulated PS differed strongly from those of the T6d data set (Fig. 5b). Only 7 % of the up-regulated T6d PS and only 1 % of the down-regulated ones were contained in LP (Fig. 5b). It may be argued that T6d contains two types of regulated genes: PS altered because of an overall altered differentiation track and PS altered because of the continued presence of the drug. The latter set of genes may be termed ‘affected by continued drug presence after long-term exposure.’ To get information on this set of genes, we subtracted the MP (washout) PS from the T6d PS. This left 393 up-regulated PS and 376 down-regulated PS. Even this subset of genes (that should in theory be enriched for PS affected by the presence of TSA) overlapped only 8 % (up) or 1 % (down) with the LP PS (Fig. 5c). These findings only corroborated our earlier conclusion that the PS of T6d or MP hardly reflected any direct responses to drug exposure.

The direct drug response (LP), related to the change in acetylation, was further characterized by analysis of GO overrepresentation. This pointed to the biological processes of ‘ion homeostasis’ and ‘phospholipase C signaling,’ and they differed strongly from the long-term response (Fig. 5d, Tab. S3).

Differences and similarities of transcriptome changes associated with early direct drug response versus continued long exposure

We explored whether the immediate drug response to TSA at the beginning of the experiment (differentiating hESC exposed during the first 6 h = T6h) was sufficiently predicting T6d. Analysis of oGO among T6h up-regulated PS pointed to altered signaling/nerve activity, and developmental GOs were not overrepresented at all among up-regulated PS (in contrast to T6d). The PS down-regulated by TSA were indicative of one major underlying biological process: Six of the 13 oGOs were related to chromatin modification and 7 to acetylation (Fig. 6a, Tab. S3). A similar response was observed for VPA (Fig. S8), and these findings are well consistent with major changes in histone acetylation. Both HDACi affected genes related to chromatin modification, such as ING5, KAT5 (histone acetylation), and several PHF genes (involved in chromatin remodeling and transcriptional regulation). Thus, one primary effect of VPA and TSA was alteration of genes involved in chromatin structure (Fig. 6a, Fig. S8). Such epigenetic mechanisms triggered early by HDACi, and fixed by longer presence of the drugs, could initialize the massive changes observed later after prolonged drug treatment.

A direct comparison of the T6h and T6d transcriptome changes showed that 70 % of the PS up-regulated early were not up-regulated late (T6d) (Fig. 6b), and 90 % of PS down-regulated early were not down-regulated late. These findings were similar for TSA and VPA (Fig. S9). Only 13 of 303 oGOs among up-regulated T6d genes (p < 0.05)
overlapped with oGOS from up-regulated T6h genes (Fig. S10). These data indicate very clearly that the early drug response predicted the overall outcome after 6 days only poorly, and vice versa the primary effects of TSA may not be identifiable from measurements at late time points.

However, there were at least some similarities of the responses: 463 PS overlapped for TSA responses after 6 h and 6 days (Fig. 6b). All oGOS among them pointed to altered signaling events, mainly related to phospholipase C (Fig. 6c; Table S4) as possibly common response feature. To narrow down the common drug responses to effects specific for HDACi, we extracted PS that were regulated by both TSA and VPA and at both time points (Fig. S9A). Eleven GOs were overrepresented among these PS (Fig. S9B), with 7 of them referring to phospholipase C regulation. They involved, e.g., EGFR, KIT, PDGFRα, and EDNRA. The other four oGOS referred either to neural crest (NC) or to their guidance and migration (comprising TWIST1, SOX10, SNAI2, and SEMA3C) (Zimmer et al. 2012). Thus, the PS up-regulated by HDACi both early and late suggested that NC is one of the cell lineages that was erroneously generated in our model when differentiation was disturbed by drugs.

Dependence of immediate transcriptome responses to HDACi on the developmental stage

The above findings suggest that longer exposure is not suitable for defining the mode of action of drugs. Some literature data suggest that short drug pulses of few hours may be more suited (Jergil et al. 2011; Theunissen et al. 2012). However, this raises the question, when such short-term exposure should be performed. A developing biological system does not only consist of one defined cell type, but it rather reflects all biological stages between the starting point and the final population. On this basis, we hypothesized that the ‘cellular’ response, i.e., short-term transcriptome alterations would strongly depend on the time point of measurement, i.e., on the differentiation stage. To test this, we compared the responses to 6-h TSA treatment of cells at the early stage of differentiation (T6h) and at the final stage (LP). The down-regulated PS of the two conditions showed hardly any overlap at all. For the up-regulated PS, 13% of the T6h PS were also found in LP and 46% of the PS of LP were found in T6h (Fig. 6d). GOs overrepresented in commonly up-regulated PS referred only to regulation of phospholipase activity. Besides this hint toward signal transduction being affected in both cases, there was little further indication of common functional effects. On the contrary, the representation of superordinate biological processes by the oGOS was largely divergent between the two 6-h pulse experiments (Fig. 6c, Tab. S3). We conclude from this that typical HDACi target genes that inform on the mode of action do not exist as such. They depended to a very high degree on the experimental setup, the time point of measurement, as shown here, and they also depended to a high extent on the length of exposure, as shown in the above paragraphs.

Discussion

Transcriptomics studies have been instrumental for characterizing the development of model organisms, the human brain, or various stem cells (Kang et al. 2011; Mariani et al. 2012; Xie et al. 2013; Yang et al. 2013). Most available data refer to fixed time points, but some studies already showed the highly dynamic behavior of such systems (Theunissen et al. 2012; Zimmer et al. 2011). An additional layer of complexity is added, when experimental disturbances are studied in systems that change over time. A stressor that switches a static system from one state to another one would be expected to shift such a dynamic system from one developmental track (dynamic series of changes) to another. Second and third levels of additional complexity are added, if the type and extent of such shifts depend on the length of exposure and on the time point of exposure relative to the normal differentiation track. Although the understanding of the temporal evolution of a developing biological model system under stress is essential in toxicology and pathophysiology, we are not aware of any study that has addressed this issue in a quantitative and systems-wide way. Three major conclusions from our study can form a basis for further exploration of this field:

First, it became clear that the usual (long-term exposure) toxicogenomics data recorded from developing stem cells describe to a large extent the cellular phenotypic changes and only to a small extent the direct drug action as such. Second, drug actions in a dynamic stem cell system can be fully reversible, even when the expression of hundreds of genes has been changed. However, at a certain point, permanent and persistent changes are triggered that can become independent of the continued presence of drug. We found that chromatin alterations in particular histone methylation could act as persistence detector or irreversibility switch. Third, even the short-term immediate effects on the transcriptome depended on the time point of drugs application. This apparently trivial finding, considering the dynamic behavior of a differentiation system, has important implications for future systems toxicology studies: Exposures at multiple times need to be performed in order to obtain data that sufficiently describe drug effects and responses of the cells.

Do transcriptomics data reveal the mechanisms by which a compound interferes with neural development? (Theunissen et al. 2012). Our data suggest unambiguously
that final changes in the transcriptome at the end of the differentiation period mainly define the cellular phenotype. This is supported by the following six lines of evidence. (a) Primary drug responses to HDACi are dominated by up-regulated genes (Berger 2007). The chromatin opening due to acetylation may be enhanced and then stabilized by increased H3K4me3 (Nightingale et al. 2007). We find here equal numbers of up- and down-regulated PS for the T6d condition. This corroborates findings in the literature (Krug et al. 2013; Theunissen et al. 2012) and suggests that indirect responses play a large role after longer (>6 h) incubation periods. (b) The primary biochemical drug response (acetylation) was not detectable after prolonged exposure. Thus, it is unlikely that the transcriptome response at late time points was triggered by a direct response to HDACi. (c) The long-term (T6d) response differed considerably from the acute response at the same time (LP). About 95 % of the T6d PS were not predicted by the LP direct drug response data. (d) The transcriptome response after a 2-day washout of the drugs (MP) was very similar to the response without washout (T6d). This makes it very unlikely that a direct drug response was measured at T6d. (e) Moreover, it should be assumed that the transcriptome response data should be to some degree independent of the experimental system, if they would characterize the MoA of drugs. However, we observed that closely related hESC-based systems showed very different responses to VPA (Krug et al. 2013) and apparent HDACi consensus genes (Jergil et al. 2009, 2011) defined in murine cells by overlapping responses of different drugs were not regulated in our cells. In summary, this corroborates our hypothesis that transcriptome data from disturbed developmental systems reflect mainly an altered phenotype and that information on direct drug effects requires short drug treatments of up to 6 h. (f) When the overlap of drug-affected ‘toxicant-response’ genes (T-genes) and developmentally regulated gene (D-gene) clusters was examined, we found that 90 % of the genes regulated late during differentiation were affected by HDACi. This high percentage of overlap is hard to explain by primary drug action; in particular as TSA acted early, and the changes occurring between day 4 and day 6 did not even require the presence of the drug. Instead, the observation is plausible, if it is assumed that the drugs changed the overall development, and therefore nearly all later phase D-genes were affected. Thus, the apparently specific effects of HDACi on neurodevelopment may be mainly due to the fact that the system studied was based on neurodevelopment. One may then hypothesize that HDACi may preferentially target cardiac genes in a cardiac development model. Such findings have indeed been obtained from differentiating mESC. VPA affected their neural development upon neurodifferentiation and cardiac development upon cardiac differentiation (Theunissen et al. 2013).

The effects of a toxicant may depend not only on the time point of measurement, but also on the duration of exposure. Some developmental toxicity responses may require the activation of a ‘persistence detector’ to distinguish between short reversible interactions on the one hand and toxicity on the other hand (Lim et al. 2013). For instance, epigenetic changes have been suggested as persistence switches for ethanol sensitization (Botia et al. 2012; Qiang et al. 2011). An altered state of cells is also fixed by so-called ‘gateway drugs.’ These are compounds allowing the later action of other drugs, even when they have been washed out for a long time. This is particularly important in the field of addiction, and the mode of action has also been explained by histone modifications (Levine et al. 2011). Molecular persistence mechanisms are of high importance in the field of developmental toxicity, in which compounds might show effects years after the exposure has taken place (Kadereit et al. 2012). Evidence is emerging that altered behavior and late-onset disorders that are triggered early in life are associated with epigenetic alterations (Rudenko and Tsai 2013; Weaver et al. 2004). Early developmental exposure to toxicants could result in an accumulation of epigenetic changes that, when reaching a certain threshold, result in transcriptome alterations associated with adverse health effects. There are several examples for toxicants or stressors that can trigger diseases in later life when exposure takes place in utero or childhood or that can even trigger trans-generational effects (McGowan et al. 2009). Early exposure to lead has, for example, been associated with Alzheimer’s disease (Wu et al. 2008). The model system chosen here was based on the known neurodevelopmental disturbances of HDACi and the associated transcriptome responses (Fig. 7a, b). We used this as a test case to study time-dependent transcriptome responses. A short pulse of TSA (EP) triggered a strong transcriptional response that was fully reversible, while a longer exposure (MP) triggered a persistent effect, even after discontinuation of drug exposure. The transient response correlated well with acetylation. The persistent response did not correlate with histone acetylation, but rather with a shift in the histone methylation ratio (Fig. 7c, d). This shift required more time and was specific for the gene studied. For instance, OTX2 was affected earlier than PAX6, and the direction of methylation changes in the OCT4 promoter (Balmer et al. 2012) was opposite to the one described here for PAX6. Histone methylations are well suited for deciding on cell fate and long-term regulations, as they can favor transcription (H3K4me) or attenuate transcription (H3K27me), depending on the site of modification. Histone methylations do not usually change globally (i.e., their overall cellular level remains constant), but the pattern of the different methylases and demethylases can change dramatically during early neural differentiation (Weng et al.
and individual promoters are affected in a highly specific manner. In particular, the ratio of promoter H3K4/H3K27 trimethylation has been shown to reflect the phenotypic plasticity of stem cells during neural fate decision. Changes in this ratio at promoters of genes associated with neural differentiation frequently precede the changes in gene expression (Burney et al. 2013).

If it is assumed that cells can react reversibly to short stimuli, as shown here, and that the duration of exposure plays a role for the overall outcome, then a mechanism functioning as drug persistence detector is required. The easiest way of imagining such a mechanism could be a superordinate regulator of a wrong pathway (e.g., a neural crest organizer) whose promoter chromatin structure functions as Boolean AND element being switched by acetylation plus a second alteration that takes more time in response to drug exposure. Short drug exposure would only trigger acetylation and thus not be sufficient for activation/switching. Longer exposure would still allow for acetylation and now also for the second change. Together, they would lead to activation of the superordinate regulator, and the activation state would be fixed by the histone methylation pattern. At present, this is a speculative hypothesis just intended to give an idea how underlying regulations may be imagined; the model system chosen may be too complex to provide causal evidence for such a mechanism with the technology presently available. Further studies will require considerable technical optimizations. ChIP can easily be accommodated to a whole genome level by applying sequencing instead of PCR as endpoint. Yet, the real issue lies in the quantification across several independent experiments with human stem cells, and the handling of the

**Fig. 7** Summary. a Neuroepithelial precursors (NEP) were generated from hESC within 6 days of differentiation (DoD). Main cell type markers are indicated. b Continuous drug treatment led to an altered NEP differentiation (NEP*) as indicated by transcriptome changes (blue), reduced marker expression (PAX6, OTX2), and permanent changes in histone methylation (after 2–3 days) at the promoters of the markers. Acetylation changes were only transient. c Drug exposure for 4 days resulted in the same disturbed NEP differentiation as continued drug exposure. The number of altered PS and the alterations of histone methylation were not affected by drug washout. d Short-term drug treatment induced a strong, but fully reversible change in gene expression. Methylation of marker promoters was not altered, and marker expression was normal. The transition of transient drug-induced gene expression and histone acetylation changes to a permanently altered transcriptome and NEP differentiation correlated with the permanence of promoter histone methylations (color figure online)
type of information resulting from this. Quantitative detection of compound-induced histone changes requires a high level of robustness and sensitivity of the method, and this is much harder to achieve for ChiP than for methods such as PCR, DNA microarray, or Western blot. Already in the present study, considerable method optimization was required to obtain reproducible and statistically significant data on the chromatin changes taking place at few selected marker genes.

The acute responses to HDACi differed significantly from the response to continued treatment. More importantly, they differed also from one another. No matter whether the experiment was performed at the beginning or toward the end of differentiation, a vastly larger number of genes were up-regulated than down-regulated (as expected of HDACi). But different genes were affected. The 6-h time point has been found to be optimal in previous studies, e.g., on stem cells (Jergil et al. 2011) or a large number of human tumors (Cohen et al. 2011) to record direct drug effects. We conclude that the direct drug effect was different at distinct times of differentiation. This corroborates earlier findings of VPA acute cytotoxicity being highly dependent on the developmental stage (Fujiki et al. 2013). Taken together, these findings imply that a potential mode of action of a developmental toxicant is not an intrinsic drug property, but it is a combined feature of the experimental system and the chemical used. In this respect, developmental toxicity may differ from other fields. Its testing may therefore require a battery of parallel tests and particularly complex systems toxicology approaches (Hermse et al. 2013; Tonk et al. 2013) relative to more acute forms of toxicity. It needs to be established whether approaches, as in the ToxCast program (Sipes et al. 2011, 2013), that rely mainly on hundreds of simple assays for biochemical/cellular targets can help to substitute or to complement the complex differentiation models used in a test battery (Piersma et al. 2013).

Our study has major implications for the design and interpretation of toxicogenomics data in development. We found that classical transcriptome data from a disturbed/stressed differentiation model strongly reflect the altered phenotype. In the case of HDACi, the phenotype contribution is >90%. This number may be smaller or larger for other stressors, but it will most likely always be sizeable. The question arises what this transcriptome information can be used for, if it does not inform on the mode of action of a chemical. We assume that different types of stressors result in different developmental disturbances and therefore also different transcriptome patterns (Balmer et al. 2012; Krug et al. 2013). Thus, the information will be useful for compound classification, differentiation, and possibly potency ranking (Schulpen et al. 2014). Beyond this, there is a further dimension of information contained in the transcriptome data. If they are largely independent of direct compound effects, then they constitute a comprehensive phenotypic description of the culture state. In the classical toxicological literature, it has always been assumed that transcriptome endpoints require phenotypic anchoring to other types of endpoints (Paules 2003; Waters and Fostel 2004). Our data suggest now that the comprehensive transcriptome data can be a phenotypic anchor as such. Transcriptome data may be more comprehensive and robust than classical endpoints (such as immunostains). It appears worthwhile to develop quantification tools that indicate, on the basis of the transcriptome as phenotypic descriptor, how big a developmental insult is, and that allow the ranking of unknown drugs, or of different concentrations or exposure times of one given drug (Waldmann et al. 2014).

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References

Abranches E, Silva M, Pradier L et al (2009) Neural differentiation of embryonic stem cells in vitro: a road map to neurogenesis in the embryo. PLoS ONE 4(7):e6286. doi:10.1371/journal.pone.0006286

Andersen ME, Krewski D (2010) The vision of toxicity testing in the 21st century: moving from discussion to action. Toxicol Sci 117(1):17–24. doi:10.1093/toxsci/kfq188

Balmer NV, Weng MK, Zimmer B et al (2012) Epigenetic changes and disturbed neural development in a human embryonic stem cell-based model relating to the fetal valproate syndrome. Hum Mol Genet 21(18):4104–4114. doi:10.1093/hmg/ddk239

Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful approach to multiple testing. J Roy Stat Soc 57(1):289–300

Berger SL (2007) The complex language of chromatin regulation during transcription. Nature 447(7143):407–412. doi:10.1038/nature05915

Berthold MR, Cebron N, Dill F et al (2007) KNIME: the konstanz information miner. In: Preisach C, Burkhardt H, Schmidt-Thieme L, Decker R (eds) Data analysis, machine learning and applications. Springer, Berlin, pp 319–326

Botia B, Legastelois R, Alaux-Cantin S, Naassila M (2012) Expression of ethanol-induced behavioral sensitization is associated with alteration of chromatin remodeling in mice. PLoS ONE 7(10):e47527. doi:10.1371/journal.pone.0047527

Boudadi E, Stower H, Halsall JA et al (2013) The histone deacetylase inhibitor sodium valproate causes limited transcriptional change in mouse embryonic stem cells but selectively overrides Polycomb-mediated Hoxb silencing. Epigenetics & Chromatin 6(1):11. doi:10.1186/1756-8935-6-11
Burney MJ, Johnston C, Wong KY et al (2013) An epigenetic signature of developmental potential in neural stem cells and early neurons. Stem Cells. doi:10.1002/stem.1431

Chambers SM, Fasano CA, Papapetrou EP, Tomishima M, Sadelain M, Studer L (2009) Highly efficient neural conversion of human ES and iPS cells by dual inhibition of SMAD signaling. Nat Biotechnol 27(3):275–280. doi:10.1038/nbt.1529

Cohen AL, Soldi R, Zhang H et al (2011) A pharmacogenomic method for individualized prediction of drug sensitivity. Mol Syst Biol 7:513. doi:10.1038/msb.2011.47

Colleoni S, Galli C, Gaspar JA et al (2011) Development of a neural teratogenity test based on human embryonic stem cells: response to retinoic acid exposure. Toxicol Sci 124(2):370–377. doi:10.1093/toxsci/kfr245

Collins FS, Gray GM, Bucher JR (2008) Toxicology. Transforming environmental health protection. Science 319(5865):906–907. doi:10.1126/science.1154619

Coskun V, Tsoa R, Sun YE (2012) Epigenetic regulation of stem cells differentiating along the neural lineage. Curr Opin Neurobiol 22(5):762–767. doi:10.1016/j.conb.2012.07.001

Crofton KM, Mundy WR, Shafer TJ (2012) Developmental neurotoxicity testing: a path forward. Congenit Anom 52(3):140–146. doi:10.1111/j.1741-4425.2012.00377.x

Dufour-Rainfray D, Vourc'h P, Le Guisquet AM et al (2010) Behavioral and serotonergic disorders in rats exposed prenatally to valproate: a model for autism. Neurosci Lett 470(1):55–59. doi:10.1016/j.neulet.2009.12.054

Fass DM, Shah R, Ghosh B et al (2010) Effect of inhibiting histone deacetylase with short-chain carboxylic acids and their hydroxamic acid analogs on vertebrate development and neuronal chromatin. ACS Med Chem Lett 2(1):39–42. doi:10.1021/ml1001954

Fujiki R, Sato A, Fujitani M, Yamashita T (2013) A proapoptotic effect of valproic acid on progenitors of embryonic stem cell-derived glutamatergic neurons. Cell Death Dis 4:e677. doi:10.1038/cddis.2013.205

Gautier L, Cope L, Bolstad BM, Irizarry RA (2004) Affy---analysis of Affymetrix GeneChip data at the probe level. Bioinformatics 20(3):377–380. doi:10.1093/bioinformatics/btg405

Gentleman RC, Carey VJ, Bates DM et al (2004) Bioconductor: open software development for computational biology and bioinformatics. Genome Biol 5(10):R80. doi:10.1186/gb-2004-5-10-r80

Gottlicher M, Minucci S, Zhu P et al (2001) Valproic acid defines a novel class of HDAC inhibitors inducing differentiation of transformed cells. EMBO J 20(J24):6969–6978. doi:10.1093/emboj/20.24.6969

Graff J, Rei D, Guan JS et al (2012) An epigenetic blockade of cognitive functions in the neurodegenerating brain. Nature 483(7388):222–226. doi:10.1038/nature10849

Hartung T, van Vliet E, Jaworska J, Bonilla L, Skinner N, Thomas R (2013) Systems toxicology. AlteX 286(41):35977–35988. doi:10.1074/jbc.M111.266254

Hezroni H, Sailaja BS, Meshorer E (2011) Pluripotency-related, valproic acid (VPA)-induced genome-wide histone H3 lysine 9 (H3K9) acetylation patterns in embryonic stem cells. J Biol Chem 286(41):35977–35988. doi:10.1074/jbc.M111.266254

Jergil M, Kultima K, Gustafson AL, Dencker L, Stigson M (2009) Valproic acid-induced deregulation in vitro of genes associated in vivo with neural tube defects. Toxicol Sci 108(1):12–148. doi:10.1093/toxsci/kfp002

Kadereit S, Zimmer B, van Thriel C, Hengstler JG, Leist M (2012) Compound selection for in vitro modeling of developmental neurotoxicity. Front Biosci 17:2442–2460

Kamieniarz K, Izzo A, Dundur M et al (2012) A dual role of histone H1.4 lys 34 acetylation in transcriptional activation. Genes Dev 26(8):797–802. doi:10.1101/gad.182014.111

Kang HJ, Kawasawa YI, Cheng F et al (2011) Spatio-temporal transcriptome of the human brain. Nature 478(7370):483–489. doi:10.1038/nature10523

Katouka S, Takuma K, Hara Y, Maeda Y, Ago Y, Matsuda T (2013) Autism-like behaviors with transient histone hyperacetylation in mice treated prenatally with valproic acid. Int J Neuropsychopharmacol 16(1):91–103. doi:10.1017/S1461145711001714

Khan N, Jeffers M, Kumar S et al (2008) Determination of the classification of small-molecule histone deacetylase inhibitors. Biochem J 409(2):581–589. doi:10.1042/BJ20070779

Krug AK, Kolde R, Gaspar JA et al (2013) Human embryonic stem cell-derived test systems for developmental neurotoxicity: a transcriptomics approach. Arch Toxicol 87(1):123–143. doi:10.1007/s00204-012-0967-3

Levine A, Huang Y, Drisaldi B et al (2011) Molecular mechanism for a gateway drug: epigenetic changes initiated by nicotine prime gene expression by cocaine. Sci Transl Med 3(107):107ra109. doi:10.1126/scitranslmed.3003062

Lim WA, Lee CM, Tang C (2013) Design principles of regulatory networks: searching for the molecular algorithms of the cell. Mol Cell 49(2):202–212. doi:10.1016/j.molcel.2012.12.020

Mariani J, Simonini MV, Palejev D et al (2012) Modeling human cortical development in vitro using induced pluripotent stem cells. Proc Natl Acad Sci USA 109(31):12770–12775. doi:10.1073/pnas.1202944109

Marinova Z, Leng Y, Leeds P, Chuang DM (2011) Histone deacetylase inhibition alters histone methylation associated with heat shock protein 70 promoter modifications in astrocytes and neurons. Neuropharmacology 60(7–8):1109–1115. doi:10.1016/j.neuropharm.2010.09.022

McGowan PO, Sasaki A, D’Alessio AC et al (2009) Epigenetic regulation of the glucocorticoid receptor in human brain associates with childhood abuse. Nat Neurosci 12(3):342–348. doi:10.1038/nn.2270

Meador KJ, Baker GA, Browning N et al (2009) Cognitive function at 3 years of age after fetal exposure to antiepileptic drugs. N Engl J Med 360(16):1597–1605. doi:10.1056/NeJMoa0803531

Menegola E, Cappelletti G, Di Renzo F (2012) Epigenetic approaches and methods in developmental toxicology: role of HDAC inhibition in teratogenic events. Methods Mol Biol 889:373–383. doi:10.1007/978-1-61779-867-2_23

Nightingale KP, Gendreizig S, White DA, Bradbury C, Hollfelder F, Turner BM (2007) Cross-talk between histone modifications in response to histone deacetylase inhibitors: MLL4 links histone H3 acetylation and histone H3K4 methylation. J Biol Chem 282(7):4408–4416. doi:10.1074/jbc.M60773200

NRC (2007) Toxicity testing in the 21st century: a vision and a strategy. National Academies Press, Washington

Paules R (2003) Phenotypic anchoring: linking cause and effect. Environ Health Perspect 111(6):A338–A339

Piersma AH, Bosgra S, van Duursen MB et al (2013) Evaluation of an alternative in vitro test battery for detecting reproductive toxicants. Reprod Toxicol 38:53–64. doi:10.1016/j.reprotox.2013.03.002

Qiang M, Denny A, Lieu M, Carreon S, Li J (2011) Histone H3K9 modifications are a local chromatin event involved in...
ethanol-induced neuroadaptation of the NR2B gene. Epigenetics 6(9):1095–1104. doi:10.4161/epi.6.9.16924

Reimand J, Anark T, Vilo J (2011) g: Profiler—a web server for functional interpretation of gene lists (2011 update). Nucleic Acids Res 39(Web Server Issue):W307–W315. doi:10.1093/nar/gkr378

Robinson JF, Pennings JL, Piersma AH (2013) Toxicogenomic approaches in developmental toxicology testing. Methods Mol Biol 947:451–473. doi:10.1007/978-1-62703-131-8_31

Robinson JF, Pennings JL, Piersma AH (2012a) A review of toxicogenomic approaches in developmental toxicology. Methods Mol Biol 889:347–371. doi:10.1007/978-1-61779-867-2_22

Robinson JF, Pennings JL, Pront TE, Piersma AH (2012b) A comparison of gene expression responses in rat whole embryo culture and in vivo: time-dependent retinoic acid-induced teratogenic response. Toxicol Sci 126(1):242–254. doi:10.1093/toxsci/kfr342

Rudenko A, Tsai LH (2013) Epigenetic regulation in memory and cognitive disorders. Neuroscience. doi:10.1016/j.neuroscience.2012.12.034

Sanchez-Serrano SL, Cruz SL, Lamas M (2011) Repeated toluene exposure modifies the acetylation pattern of histones H3 and H4 in the rat brain. Neurosci Lett 489(3):142–147. doi:10.1016/j.neulet.2010.12.004

Schulpen SH, Pennings JL, Tonk EC, Piersma AH (2014) A statistical approach towards the derivation of predictive gene sets for potency ranking of chemicals in the mouse embryonic stem cell test. Toxicol Lett 225(3):342–349. doi:10.1016/j.toxlet.2014.01.017

Sipes NS, Martin MT, Reif DM et al (2011) Predictive models of prenatal developmental toxicity from ToxCast high-throughput screening data. Toxicol Sci 124(1):109–127. doi:10.1093/toxsci/kfr220

Sipes NS, Martin MT, Kothiya P et al (2013) Profiling 976 ToxCast chemicals across 331 enzymatic and receptor signaling assays. Chem Res Toxicol 26(6):878–895. doi:10.1021/tx400021f

Smyth GK (2005) Limma: linear models for microarray data. In: Gentleman R, Carey V, Dudoit S, Irizarry W, Huber W (eds) Bioinformatics and computational biology solutions using R and Bioconductor, 2006/05/02 edn. Springer, New York, pp 397–420

Theunissen PT, Robinson JF, Pennings JL et al (2012) Transcriptomic concentration-response evaluation of valproic acid, cyproconazole, and hexaconazole in the neural embryonic stem cell test (ESTn). Toxicol Sci 125(2):430–438. doi:10.1093/toxsci/kfr293

Theunissen PT, Pennings JL, van Dartel DA, Robinson JF, Kleinjans JC, Piersma AH (2013) Complementary detection of embryotoxic properties of substances in the neural and cardiac embryonic stem cell tests. Toxicol Sci 132(1):113–120. doi:10.1093/toxsci/kfs333

Tonk EC, Robinson JF, Verhoef A, Theunissen PT, Pennings JL, Piersma AH (2013) Valproic acid-induced gene expression responses in rat whole embryo culture and comparison across in vitro developmental and non-developmental models. Reprod Toxicol. doi:10.1016/j.reprotox.2013.06.069

Tung EW, Winn LM (2010) Epigenetic modifications in valproic acid-induced teratogenesis. Toxicol Appl Pharmacol 248(3):201–209. doi:10.1016/j.taap.2010.08.001

Umlauf D, Goto Y, Feil R (2004) Site-specific analysis of histone methylation and acetylation. Methods Mol Biol 287:99–120. doi:10.1385/1-59259-828-5:099

van Darrel DA, Zeijen NJ, de la Fonteyne LJ, van Schooten FJ, Piersma AH (2009) Disentangling cellular proliferation and differentiation in the embryonic stem cell test, and its impact on the experimental protocol. Reprod Toxicol 28(2):254–261. doi:10.1016/j.reprotox.2009.03.017

Vojnits K, Ensenat-Waser R, Gaspar JA et al (2012) A transcriptomic study to elucidate the toxicological mechanism of methylmercury chloride in a human stem cell based in vitro test. Curr Med Chem 19(36):6224–6232

Waldmann T, Rempel E, Balmer NV et al (2014) Design principles of concentration-dependent transcriptome deviations in drug-exposed differentiating stem cells. Chem Res Toxicol. doi:10.1021/tr400402j

Waters MD, Fostel J (2004) Toxicogenomics and systems toxicology: aims and prospects. Nat Rev Genet 5(12):936–948. doi:10.1038/nrg1493

Weaver IC, Cervoni N, Champagne FA et al (2004) Epigenetic programming by maternal behavior. Nat Neurosci 7(8):847–854. doi:10.1038/nn1276

Weng MK, Zimmer B, Polt D et al (2012) Extensive transcriptional regulation of chromatin modifiers during human neurodevelopment. PLoS ONE 7(5):e36708. doi:10.1371/journal.pone.0036708

Wobus AM, Loser P (2011) Present state and future perspectives of using pluripotent stem cells in toxicology research. Arch Toxicol 85(2):79–117. doi:10.1007/s00204-010-0641-6

Wu J, Bashra MR, Brock B et al (2008) Alzheimer’s disease (AD)-like pathology in aged monkeys after infantile exposure to environmental metal lead (Pb): evidence for a developmental origin and environmental link for AD. J Neurosci 28(1):3–9. doi:10.1523/JNEUROSCI.4405-07.2008

Xie W, Schultz MD, Lister R et al (2013) Epigenomic analysis of multilineage differentiation of human embryonic stem cells. Cell 153(5):1134–1148. doi:10.1016/j.cell.2013.04.022

Xu X, Duan S, Yi F, Ocampo A, Liu GH, Izpisua Belmonte JC (2013) Mitochondrial regulation in pluripotent stem cells. Cell Metab. doi:10.1016/j.cmet.2013.06.005

Yang H, Zhou Y, Gu J et al (2013) Deep mRNA sequencing analysis to capture the transcriptome landscape of zebrafish embryos and larvae. PLoS ONE 8(5):e64058. doi:10.1371/journal.pone.0064058

Zimmer B, Kugler PB, Baudis B et al (2011) Coordinated waves of gene expression during neuronal differentiation of embryonic stem cells as basis for novel approaches to developmental neurotoxicity testing. Cell Death Differ 18(3):383–395. doi:10.1038/cdd.2010.109

Zimmer B, Lee G, Stiegler NV et al (2012) Evaluation of developmental toxicants and signaling pathways in a functional test based on the migration of human neural crest cells. Environ Health Perspect. doi:10.1289/ehp.1104489