Tox: a multifunctional transcription factor and novel regulator of mammalian corticogenesis

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Abstract

Major efforts are invested to characterize the factors controlling the proliferation of neural stem cells. During mammalian corticogenesis, our group has identified a small pool of genes that are transiently downregulated in the switch of neural stem cells to neurogenic division and reinduced in newborn neurons. Among these switch genes, we found Tox, a transcription factor with hitherto uncharacterized roles in the nervous system. Here, we investigated the role of Tox in corticogenesis by characterizing its expression at the tissue, cellular and temporal level. We found that Tox is regulated by calcineurin/Nfat signalling. Moreover, we combined DNA adenine methyltransferase identification (DamID) with deep sequencing to characterize the chromatin binding properties of Tox including its motif and downstream transcriptional targets including Sox2, Tbr2, Prox1 and other key factors. Finally, we manipulated Tox in the developing brain and validated its multiple roles in promoting neural stem cell proliferation and neurite outgrowth of newborn neurons. Our data provide a valuable resource to study the role of Tox in other tissues and highlight a novel key player in brain development.

Keywords brain development; DamID sequencing; HMG-box transcription factors; neural stem cells; Tox

Subject Categories Neuroscience

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Introduction

During embryonic development of the mammalian brain, neural stem and progenitor cells progressively switch from proliferative divisions that expand the stem cell pool to differentiative divisions that generate neurons and glia (Kriegstein & Alvarez-Buylla, 2009; Taverna et al, 2014). Understanding the molecular mechanisms underlying this switch is a major goal in developmental and stem cells biology both for basic research and for potential applications in regenerative therapy (Goldman, 2005; Lindvall & Kokaia, 2006).

Over the years, several transcription factors and signalling molecules have been characterized that are instrumental in the regulation of corticogenesis (Guillemot, 2007; Pinto & Gotz, 2007; Paridaen & Huttner, 2014). Yet, many other factors remain elusive or await further characterization.

Next-generation sequencing has recently provided the field with a powerful tool to identify novel factors involved in corticogenesis (Han et al, 2009; Ayoub et al, 2011; Fietz et al, 2012; Yao et al, 2012), but major limitations remain in the identification and sorting of individual cell types for comparative analyses. Specifically, in the rodent brain the two germinal layers of the ventricular and subventricular zone (VZ and SVZ, respectively) consist of a varying proportion of intermingled progenitors undergoing proliferative or differentiative divisions (Kriegstein & Alvarez-Buylla, 2009; Taverna et al, 2014). The vast majority of proliferative progenitors (PP) are confined to the VZ where they divide at its apical boundary and are for this reason also referred to as apical progenitors. In contrast, differentiating progenitors (DP) migrate to the basal boundary of the VZ to form the SVZ and are for this reason also referred to as basal progenitors (Kriegstein & Alvarez-Buylla, 2009; Taverna et al, 2014). While apical and basal progenitors represent, overall, the vast majority of PP and DP, respectively, during development an increasing proportion of apical progenitors switch their fate to become DP and generate basal progenitors or neurons (Haubensak et al, 2004; Miyata et al, 2004; Noctor et al, 2004). Conversely, a small proportion of basal progenitors can become PP to expand their pool within the SVZ (Attardo et al, 2008; Ochiai et al, 2009). Finally, newborn neurons generated in the VZ or SVZ by apical or basal DP, respectively, migrate through the intermediate zone (IZ) to form the cortical plate (CP) (Kriegstein & Alvarez-Buylla, 2009; Taverna et al, 2014). Hence, difficulties in the isolation of intermingled PP and DP, and identification of either progenitor type from neurons, limited the use of next-generation sequencing to comparisons of developmental stages and portions of tissues or species rather than individual cell types (Han et al, 2009; Ayoub et al, 2011; Fietz et al, 2012; Yao et al, 2012).

Our group has recently overcome this limitation by generating a dual-reporter, Btg2RFp/Tubb3GFP mouse line in which PP (RFp/GFP), DP (RFp/GFP ) and neurons (GFP), irrespective from RFp were identified by the combinatorial expression of two fluorescent reporters (Aprea et al, 2013). Transcriptome sequencing of the three
cell types led to the identification of ~200 transcripts being upregulated solely in DP and downregulated in both PP and neurons (on-switch genes). Conversely, a similarly abundant pool of genes was downregulated in DP and upregulated in both PP and neurons (off-switch genes). Supporting the notion that switch genes characterize the signature of neurogenic commitment, it was found that essentially all neurogenic markers were on-switches (Aprea et al, 2013). Moreover, confirming that a remarkably high proportion of switch genes are functionally relevant in corticogenesis, manipulation of novel or uncharacterized on/off-switch transcripts in vivo led to the identification of several genes with hitherto unknown functions in brain development (Aprea et al, 2013, 2015; B. Artegiani, J. Aprea, S. Bragado Alonso, F. Calegari, unpublished data).

Surprisingly, and in stark contrast to on-switch genes, our group has found that only a remarkably small number of off-switches have ever been studied in the context of cortical development (Aprea et al, 2013). This bias and under-representation of off-switch genes with functional roles in brain development was puzzling since it can be inferred that both on- and off-switches should be equally important. Hence, to challenge our assumption, we next sought to characterize novel off-switch genes with unknown roles in neural stem cell commitment.

Among these, we found Tox (thymocyte selection-associated HMG-box) that is downregulated 2.7-fold in the switch from PP to DP and upregulated 3.0-fold in newborn neurons relative to DP (Aprea et al, 2013). Tox is a highly conserved transcription factor belonging to the family of HMG-box proteins and being expressed in the thymus, liver and brain (Wilkinson et al, 2002; Allahamad et al, 2012). Tox downstream targets and its binding motif are currently unknown but, like Sox proteins, its single HMG-box domain is expected to confer sequence specificity in DNA binding (Stros et al, 2007). Yet, analysis of Tox amminoacidic sequence has led to the opposite conclusion suggesting that its binding is structure, rather than sequence, specific (O’Flaherty & Kaye, 2003).

Until now, Tox was only studied in the context of the immune system where it is required for lymphoid tissue organogenesis and regulation of T-cell and natural killer cell development during hematopoiesis (Wilkinson et al, 2002; Allahamad et al, 2004, 2010; Yun et al, 2011). Expression pattern and functional relevance of this transcription factor in any other tissue, including the nervous system, are completely uncharacterized. For these reasons, we decided to characterize: (i) the upstream molecular mechanisms controlling Tox expression during development; (ii) its downstream targets; and (iii) functional role during the fate switch of PP to DP and maturation of postmitotic neurons during mammalian cortical development.

Results

Tox expression during corticogenesis recapitulates the gradients of neurogenesis

First, we analysed Tox expression by immunohistochemistry at the onset, mid and end of cortical neurogenesis as well as in the adult brain. At embryonic day (E) 9.5, Tox displayed a rostral-high/caudal-low and lateral-high/medial-low gradient of expression (Fig 1A, top-left) that closely resembled the known gradients of differentiation at the onset of cortical neurogenesis (Caviness et al, 2009). Accordingly, the proportion of Tox+ neuroepithelial cells varied from nearly 100% to essentially 0%. Moreover, the very few Tubb3+ neurons that could be found at this stage were Tox+ (Supplementary Fig S1A), suggesting that newborn neurons do not express Tox. Similar results were obtained 3 days later at E12.5, again showing a rostral-high/caudal-low and lateral-high/medial-low gradient of expression within the neuroepithelium with, additionally, Tox+ neurons particularly at the level of the prospective olfactory bulb and mid/hindbrain regions (Fig 1A, top-right; and data not shown).

At E15.5, the rostro-caudal gradient of Tox expression essentially disappeared with virtually all cells in the VZ being Tox+ (Fig 1A, bottom-left). In contrast, a lateral-medial gradient was still detectable (Fig 1A, bottom-right) that was reminiscent of the gradients of expression reported for the neurogenic markers Tbr2 and Ngn2 (Englund et al, 2005; Britz et al, 2006). Although generally abundant in progenitors of the VZ, Tox was undetectable in basal progenitors of the SVZ and was reinduced in most neurons of all areas analysed (Fig 1A, bottom and Supplementary Fig S1B). With regard to the latter cell pool, we observed that immature, migrating neurons in the IZ displayed a much lower, if any, expression of Tox as compared to more differentiated neurons in the CP that were essentially all Tox+ (Fig 1B). This observation is consistent with analyses at E9.5 and E12.5 showing that Tox expression is restricted within the subset of more mature neurons. Overall, Tox immunoreactivity recapitulated its mRNA levels previously assessed by transcriptome analysis of PP, DP and neurons (Aprea et al, 2013) although the overlap with the neurogenic gradients and difference among IZ and CP neurons could not be detected by pooling together cortical areas and neuron subtypes.

Co-labelling with cellular markers identifying the different classes of neural progenitors was performed to characterize the identity of Tox+ cells at mid-corticogenesis at E15.5. In the VZ, essentially all (95.8 ± 2.6%) Tox+ cells were positive for the proliferative marker Sox2 (Fig 1C and D, left). Conversely, many Sox2+ cells did not express Tox due to its gradients of expression already mentioned (Supplementary Fig S1B). Interestingly, quantification of Sox2 immunoreactivity within Tox+ nuclei of individual cells showed a clear positive correlation with the levels of Tox (Fig 1D, middle). This held true in different regions of the VZ despite the fact that each region varied with regard to the proportion of Tox+ cells (Supplementary Fig S1B and data not shown). To identify VZ progenitors committed to differentiative divisions, we next used the Btg2BFP reporter mouse that revealed not only that in the VZ only a small proportion of Tox+ cells were Btg2+ (14.5 ± 2.4%) but also that in Btg2+ cells the level of Tox was significantly lower than in Btg2- cells (Fig 1D). In contrast, the second germinal layer, the SVZ, was virtually deficient of Tox+ cells in all regions analysed (Fig 1C), thus indicating a remarkably tight control in Tox protein levels in different progenitor types.

We next analysed Tox expression at the end of the neurogenic interval at E18.5. Tox was maintained in the germinal layers, and at this stage, the lateral-medial gradient spread much further than at E15.5 and was almost undetectable with essentially all cells in the VZ of the lateral cortex being Tox+ (Fig 1A, bottom-right). The only germinal zone in which Tox+ cells were undetectable was the prospective hippocampus that is known to have a prolonged neurogenic interval extended to postnatal life. Again, no significant
expression of Tox was observed within the residual SVZ being detectable at this stage (data not shown). With regard to neurons, Tox was enriched in deeper layer neurons of the CP with its expression broadly overlapping with the deep layer marker Ctip2 and detected also in a proportion of double-positive Ctip2+/Satb2+ neurons (Fig 1A, bottom-right, Supplementary Fig S1C and further discussed below).

Finally, we found that Tox in the adult brain was maintained in the sub-ventricular zone but, surprisingly, not in the second adult neurogenic niche of the hippocampus that was never found to express this transcription factor throughout development (Supplementary Fig S1D and D0, and data not shown). In addition, Tox in the adult cortex was still expressed in the majority (71.7 ± 10.2%) of Ctip2+ neurons (Supplementary Fig S1D and D0), suggesting that Tox in the neuronal compartment identifies a specific sub-class of cortical neurons during both embryonic and postnatal life.

All together, these data show that Tox expression in progenitors cells of the developing VZ strongly correlates with the temporal and spatial gradients of neurogenesis. Perhaps counterintuitively, Tox expression within neurogenic areas was found to be higher in cells undergoing proliferative, rather than neurogenic, division such as Sox2+ and/or Btg2+ cells and apical, rather than basal, progenitors. The substantial decrease in Tox expression in DP of the VZ and SVZ and reinduction in CP neurons further indicated that this transcription factor is tightly regulated at the transcriptional and, possibly, post-translational level. In fact, analysis of Tox protein sequence identified four sites of ubiquitination suggesting that this transcription factor is actively degraded in the switch from PP to DP (data not shown). Its reinduction in a specific sub-class of neurons, primarily Ctip2+, may in turn suggest that Tox plays different roles in progenitors and neurons and, possibly, in different neurogenic niches of the adult brain.

**Tox is regulated by the calcineurin/Nfat signalling pathway**

To gain insight into the molecular mechanisms regulating Tox expression during corticogenesis, we took advantage of previous evidence linking calcineurin activity to Tox levels in thymocytes (Aliahmad et al., 2004) as well as our own analysis by NetPhos predicting several phosphorylation sites in the Tox sequence (data not shown). Calcineurin is a Ca2+-dependent phosphatase involved in several biological functions including stem cell commitment and brain function (Crabtree & Olson, 2002; Horsley et al., 2008; Mukherjee & Soto, 2011; Kujawski et al., 2014). Calcineurin acts by controlling nuclear trafficking of target proteins including several transcription factors, among which the Nfat family members are some of the most characterized (Crabtree & Olson, 2002; Horsley et al., 2008; Mukherjee & Soto, 2011; Kujawski et al., 2014). Calcineurin acts by controlling nuclear trafficking of target proteins including several transcription factors, among which the Nfat family members are some of the most characterized (Crabtree & Olson, 2002; Horsley et al., 2008; Mukherjee & Soto, 2011; Kujawski et al., 2014).
Figure 2. Tox is regulated by calcineurin/Nfat4 signalling.

A–B′ Layout of CsA, or vehicle as control, administration during embryonic development (A) and Western blot analysis (B) of Tox levels in cytoplasmic (cyt) or nuclear (nuc) fractions of E15.5 brains. Quantifications (B′) were performed upon internal normalization for Gadph or Tbp and referred to treatment with vehicle. n = 3; error bars = SD; *P < 0.05.

C Map of Tox locus and Nfat-binding recognition sites (red) identified by MatInspector.

D–E′ Layout of CsA or vehicle administration and Western blot analysis as in (A–B′) upon in utero electroporation with CA-Nfat3 or CA-Nfat4 vectors at E13.5. Tox in extracts of E15.5 brains was quantified normalized to Tbp and referred to CsA treatment. n = 3; error bars = SD; ***P < 0.001.

F–G′ Fluorescent pictures of the mouse cortex upon electroporation with GFP, CA-Nfat3 or CA-Nfat4 (as indicated) followed by GFP (green) or Tox (G, red) immunolabelling and DAPI counterstaining (blue). Tox upregulation after CA-Nfat4 overexpression was assessed within nuclei of GFP+ cells (n = 30) in the VZ and represented as a box plot relative to GFP− cells (n = 30) used as internal control (G′). Tox ectopic expression within the SVZ is indicated (G, arrowheads). Scale bar, 50 μm (F) and 5 μm (G); **P < 0.01.
et al, 2008). Hence, we first sought to investigate whether Tox was a direct target of calcineurin.

To this aim, we administered the highly specific, placentapermeable calcineurin inhibitor cyclosporin A (CsA) (Liu & Janeway, 1991; Graef et al, 2003) by repetitive injections in pregnant females at E13.5 and collected embryos 48 h later (Fig 2A). Western blot analyses of nuclear versus cytosolic extracts derived from either vehicle or CsA-treated mice did not show either a change in the cellular localization of Tox nor a shift in its apparent molecular weight resulting from hyper-phosphorylation (Fig 2B). In contrast, we observed a 30% decrease in Tox expression levels upon CsA treatment as compared to controls (Fig 2B and B’), suggesting that Tox is indirectly regulated by calcineurin rather than being a direct target of dephosphorylation and nuclear transport.

To calculate calcineurin-dependent transcription factors responsible for the regulation of Tox expression by calcineurin, we then analysed its 5’-region by MatInspector and found with high-confidence three binding sites for Nfat (Fig 2C). Furthermore, transcriptome analysis of PP, DP and neurons (Aprea et al, 2013) indicated that only two of the four calcineurin-dependent Nfat members are expressed at any significant level during corticogenesis allowing us to focus our attention on Nfat3(c4) and Nfat4(c3) as two candidate regulators of Tox expression through calcineurin. To address each, we next used in utero electroporation and overexpressed calcineurin-independent, constitutively active versions of either of the two Nfat members and investigated whether this manipulation rescued the downregulation of Tox induced by CsA.

Electroporation was performed at E13.5 with plasmids encoding either a constitutively active (CA) Nfat3 or Nfat4 and using a co-injected GFP vector as a reporter of targeted cells. Subsequently, pregnant mice were administered CsA for 48 h (Fig 2D), embryos collected at E15.5 and the GFP+ portion of the cortex microdissected together with its contralateral side serving as an internal negative control. Western blot analyses of nuclear extracts showed unchanged levels, hence no rescue, of Tox upon overexpression of CA-Nfat3 while, in contrast, overexpression of CA-Nfat4 triggered a massive upregulation of Tox sufficient not only to rescue the effects of CsA but also to raise its levels by 30-fold (Fig 2E and E’).

Consistent with the lack of effect of CA-Nfat3 on Tox, and as previously reported (Kurabayashi & Sanada, 2013), we found that the distribution of GFP+ cells across the cortical layers of the E15.5 brain upon CA-Nfat3 electroporation was indistinguishable from that of brains electroporated with control plasmids (Fig 2F), indicating that overexpression of CA-Nfat3 had no obvious consequence on brain development. In contrast, interestingly, electroporation of CA-Nfat4 showed an almost complete lack of GFP+ cells in the IZ/CP (Fig 2F), showing that this Nfat family member plays crucial roles in neurogenic commitment, neuronal maturation, migration and/or survival, some of which may, at least in part, depend on its effect on Tox.

Finally, we sought to corroborate our biochemical assessment of Tox upregulation induced by CA-Nfat4 (Fig 2E and E’) at the cellular level and quantified Tox immunoreactivity in nuclei of randomly chosen GFP+ cells after electroporation while using GFP- cells as an internal negative control. This showed that CA-Nfat4 induced an increase by ~35% in Tox immunoreactivity within the VZ (Fig 2G and G’) and that within the SVZ several GFP+ cells started to be characterized by levels of Tox immunoreactivity comparable to VZ cells (Fig 2G). The latter increase of GFP+ relative to GFP- cells in the SVZ, however, could not be assessed mathematically given the fact that the latter do not express any detectable level of Tox (Fig 1C). Notwithstanding the many differences in specificity, sensitivity and saturation of signal inherent in the two methods, we conclude that the increase in Tox levels by Nfat4 as assessed by Western blot (Fig 2E’) results from the combined effects observed by immunohistochemistry in the VZ and SVZ (Fig 2G). Hence, these data suggest that Tox expression during corticogenesis is controlled by the calcineurin/Nfat signalling pathway and that among Nfat family members Nfat4 is the main upstream regulator of Tox.

**DamID-Seq identifies the features of Tox binding to chromatin and its downstream targets**

We next sought to identify the downstream transcriptional targets of Tox. However, commercially available Tox antibodies were not validated for ChIP and, in our conditions, we did not achieve immunoprecipitation with any of the antibodies tested (data not shown). Hence, as an alternative method, we decided to use DNA adenine methyltransferase identification (DamID) to characterize DNA binding of proteins by their fusion to the prokaryotic Dam-methylase resulting in adenine methylation of GATC sequences (van Steensel & Henikoff, 2000; Vogel et al, 2006, 2007; Southall & Brand, 2007) (Supplementary Fig 2A). In this system, biases in the identification of targets are minimized by the fact that Dam methylation can occur at a considerable distance from the transcription factor binding site, up to about ±2 kb, while GATC motifs have a median occurrence every 460 nucleotides in the human genome (own bioinformatic analysis), that is about eight motifs within methylation range. Furthermore, differences in chromatin accessibility, untargeted binding and methylation are controlled for by the use of unused Dam. Of note, even recently DamID is typically combined with microarray analysis (van Bemmel et al, 2013; Southall et al, 2014), which is known to provide a lower spatial resolution than ChIP-Seq in defining binding domains (Shimbo et al, 2013). Therefore, inspired by reports on nuclear envelop proteins (Wu & Yao, 2013), we explored the use of high-throughput sequencing combined with DamID (DamID-Seq) to increase both the spatial resolution and quantitative read-out of transcription factor binding to chromatin regions.

DamID was performed in HEK-293T and Neuro-2a cells in parallel since both were expected to provide specific advantages. In particular, we considered HEK-293T as a system that should in principle allow the identification of Tox targets independently of any bias towards a specific lineage. Conversely, Neuro-2a may have provided an in vitro model for the identification of neural-specific Tox targets. As a recognized key factor for the successful use of DamID (Southall & Brand, 2007; Vogel et al, 2007), we first confirmed that fusion to Dam did not perturb the nuclear localization of Tox as assessed upon its overexpression by transfection with a constitutive Tox-Dam expression vector in vitro (Supplementary Fig S2B). Moreover, DamID relies on the minimal expression of the transgenes, which is necessary for a highly specific methylation pattern without saturation. Therefore, we used highly diluted lentiviruses expressing Dam or Tox-Dam under two inducible promoters
neither of which was ever induced and resulting in undetectable levels of Tox-Dam expression derived from double-leakiness. Genomic DNA was collected 2 days later, methylated-GATC fragments isolated, and high-throughput DNA sequencing performed (Supplementary Fig S2A).

Pearson’s correlation analysis revealed the high reproducibility of the two biological replicates and the diversity between samples obtained after infection of HEK-293T cells with Dam or Tox-Dam viral suspensions (Fig 3A). In contrast, surprisingly, Neuro-2a cells did not show any significant difference between conditions, neither using the same viral titre applied to HEK-293T cells nor a 10-fold diluted suspension (data not shown). We do not know the reason for this discrepancy among cell lines, but it is reasonable to assume that the expression of endogenous Tox will compete with Tox-Dam.

Figure 3. DamID-Seq identifies Tox targets.
A Pearson’s correlation analysis between Tox-Dam versus Dam reads of HEK-293T cells sampling the genome in windows of 10 kb.
B Meta-analysis of normalized Tox-Dam at the transcription start site (TSS) of Ensembl-annotated genes.
C Profile of normalized Tox-Dam reads on cell-specific enhancers (colours; categories ordered according to enrichment). Background (grey) was assessed based on the Tox signal at random genomic locations.
D Relationship between the fold-change of Tox peaks and their distance to the TSS. Note the enrichment in peaks at the boundary of ±15 kb from TSS and the correlation between their fold-change and proximity to the TSS.
D’ Distribution of Tox peaks within regions of loci (as indicated). The relative length of each genomic region was chosen to represent its average length across the human genome and circles distributed to represent the proportion of Tox peaks mapping on that given region (1 circle = 1%).
E PhastCons scores of 100 vertebrate genomes averaged for the Tox peak locations (blue) and for a matched genomic background (grey) showing higher conservation within Tox peaks (start-end).
F Putative 10-mer binding motif of Tox identified with gimmeMotifs.
for DNA binding. This is unlikely to occur in HEK-293T cells since Tox is not expressed at any detectable level in this cell line (Supplementary Fig S2B). Hence, we continued our study using data obtained from HEK-293T cells.

We next pooled the HEK-293T replicates together and mapped reads in each condition to assess their local enrichment. Selection of Tox-Dam peaks ($P < 0.01$) that were 2.5-fold enriched relative to Dam resulted in the identification of ~13,000 chromatin regions (Supplementary File S1). Among these, as expected for a transcription factor, we found that Tox-Dam peaks were characterized by a substantial increase in the coverage of regions in proximity of transcriptional start sites (TSS) relative to Dam controls (Fig 3B).

Interestingly, enrichment in Tox-Dam peaks mapping on active enhancers of different tissues (Andersson et al., 2014) was revealed to be even more prominent in enhancers of the telencephalon, neural stem cells and neurons than those of the immune system, such as macrophages and T cells (Fig 3C) in which Tox function was first characterized. Furthermore, we found a strong, positive correlation between the fold-change of Tox-Dam peaks and their vicinity to a TSS (Fig 3D). This feature, together with a drastic increase in the number of Tox-Dam peaks within a boundary of ±15 kb from a TSS (Fig 3D), suggested that Tox has a higher affinity and/or specificity of binding within this region. Hence, we identified annotated genes within ±15 kb from a Tox-Dam peak resulting in 9,368 loci encoding potential Tox targets (Supplementary File S1). Both the number of loci and the distribution of peaks within intronic, exonic and untranslated regions of annotated genes (Fig 3D') compared well with previous characterizations of transcription factor binding by either ChIP or DamID followed by either microarray or sequencing analysis (Chen et al., 2008; Southall & Brand, 2009; Shimbo et al., 2013).

Consistent with the high evolutionary conservation of Tox family members (O’Flaherty & Kaye, 2003; and our own analysis, data not shown), we found that Tox-Dam peaks were characterized by a substantial increase in non-exonic nucleotide conservation across 100 vertebrates species (Fig 3E). Finally, discovery of over-represented sequences within the 1,000 Tox peaks with the highest fold-change and up to 1 kb in length allowed us to identify a 10-mer binding motif of Tox that is rich in GC sequences (Fig 3F).

Altogether, the use of DamID-Seq gave us a pipeline for studying the features of chromatin binding of this transcription factor together with a list of its putative targets, both of which were essential for further dissecting its biological function.

**Tox can act as a repressor or activator of genes critical for corticogenesis**

We next investigated whether the 9,368 putative targets of Tox were proportionally over-represented in functional, gene ontology (GO) terms for: (i) cellular compartment, (ii) molecular function and (iii) biological process. Using DAVID and clustering genes with redundant terms together, we found a substantial enrichment in (i) nucleoplasm and cellular protrusions (Fig 4A, left), (ii) DNA binding, transcription factors, kinases and cytoskeletal proteins (Fig 4A, middle) and (iii) axonogenesis, CNS development, neurogenesis and regulation of transcription (Fig 4A, right). This enrichment in biological processes closely related to brain development and neuronal protrusions is particularly striking if one considers that the use of HEK-293T cells, contrary to Neuro-2a, did not imply a priori any bias towards the expression of neuronal genes.

Beyond genomewide analyses and clustering by GO terms, we next sought to manually inspect individual Tox peaks for at least some of the genes that are known to play key roles in CNS development including proliferation versus differentiation of neural progenitors and maturation and specification of postmitotic neurons. These included: (i) key members of the Shh, Wnt, Notch and Fgf signalling pathways (Gli1 is shown as an example in Fig 4B; top); (ii) well-characterized regulators of neural stem cell fate such as Tbr2/Eomes, Sox2, Prox1, Foxp1/2 and Yap1 (e.g. Tbr2/Eomes: Fig 4B; middle); and (iii) genes involved in neuronal maturation and specification such as Robo2, Tbr1, Satb2 and ErbB4 (e.g. Robo2: Fig 4B; bottom). In almost all cases analysed, we found that Tox-Dam peaks were characterized by very sharp and defined increases in coverage relative to Dam controls and that the vast majority of these peaks well exceeded the 2.5-fold threshold of confidence originally considered to identify the 9,368 putative target of Tox (Supplementary File S1).

These data revealed the high resolution and confidence achieved by DamID-Seq for transcription factor binding to chromatin regions. Yet, this did not provide us with any information about the effect of this binding on the expression of those targets in tissue. In order to achieve this, and validate our approach in vivo, we performed expression profiling of the putative Tox targets mentioned above by qRT–PCR upon Tox overexpression in developing mouse embryos. For this purpose, only multi-exon targets were chosen to discriminate transcripts from genomic DNA contamination after PCR amplification.

**In utero electroporation** was performed at E13.5 with control or Tox-expressing vectors, each encoding GFP to identify targeted cells. Embryos were collected 48 h later and the targeted area dissected to isolate GFP+ cells by FAC sorting. RNA extraction was followed by qRT–PCR using primers (Supplementary Table S1) spanning across at least two exons of putative Tox targets whose up/downregulation could potentially explain the high enrichment in GO terms related to neurogenesis, axonogenesis, CNS developmental and regulation of transcription (Fig 4A).

Validating DamID-Seq in vivo, we found that the abundance of all transcripts being investigated was significantly up- or downregulated upon Tox overexpression compared to GFP controls (Fig 4C). This further indicated that Tox can act both as a transcriptional activator or repressor of its targets.

**Tox inhibits the differentiation of cortical progenitors**

To investigate the function of Tox during cortical development, we next decided to manipulate its expression by in utero electroporation. Vectors encoding Tox together with GFP, or GFP alone as control, were delivered at E13.5 and embryos collected 48 h later. Distribution of GFP+ electroporated cells, contrary to Neuro-2a, did not imply a priori any bias towards the expression of neuronal genes.

Beyond genomewide analyses and clustering by GO terms, we next sought to manually inspect individual Tox peaks for at least some of the genes that are known to play key roles in CNS development including proliferation versus differentiation of neural progenitors and maturation and specification of postmitotic neurons. These included: (i) key members of the Shh, Wnt, Notch and Fgf signalling pathways (Gli1 is shown as an example in Fig 4B; top); (ii) well-characterized regulators of neural stem cell fate such as Tbr2/Eomes, Sox2, Prox1, Foxp1/2 and Yap1 (e.g. Tbr2/Eomes: Fig 4B; middle); and (iii) genes involved in neuronal maturation and specification such as Robo2, Tbr1, Satb2 and ErbB4 (e.g. Robo2: Fig 4B; bottom). In almost all cases analysed, we found that Tox-Dam peaks were characterized by very sharp and defined increases in coverage relative to Dam controls and that the vast majority of these peaks well exceeded the 2.5-fold threshold of confidence originally considered to identify the 9,368 putative target of Tox (Supplementary File S1).

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Validating DamID-Seq in vivo, we found that the abundance of all transcripts being investigated was significantly up- or downregulated upon Tox overexpression compared to GFP controls (Fig 4C). This further indicated that Tox can act both as a transcriptional activator or repressor of its targets.
Figure 4. Tox controls neurogenic determinants.

A Functional enrichment scores for cellular compartment, molecular function and biological process (left to right).

B Examples of Tox-Dam (green) or Dam (grey) peaks (superimposed in each panel) for Gli1, Eomes/Tbr2 and Robo2 (top to bottom) to represent factors involved in signalling, neurogenic commitment and neurite outgrowth, respectively.

C qRT–PCRs of GFP+ FAC-sorted cells from the E15.5 mouse brain upon electroporation at E13.5 with control (white) or Tox (black) constructs. Primers (Supplementary Table S1) were selected to amplify multi-exon transcripts (indicated) to represent the classes of genes that would explain functional annotations in (A). n ≥ 3; error bars = SD; *P < 0.05; **P < 0.01; ***P < 0.001.
To this aim, we performed electroporation in investigated whether its overexpression altered the proportion of was perturbed by the fusion of Tox to Dam. Supplementary Fig S2C) making it unlikely that chromatin binding vectors induced essentially identical phenotypes (Fig 5A and our DamID-Seq analysis, electroporation with Tox or Tox-Dam of CA-Nfat4 (Figs 2F and 5A). Moreover, as an additional control of while, conversely, increasing it in the SVZ by 10% (32.6 26.4 respectively correlated in physiological conditions (Fig 1C and C not significant; *P < 0.05, **P < 0.01, ***P < 0.001. Scale bars, 50 μm. of CA-Nfat4 (Figs 2F and 5A). Moreover, as an additional control of our DamID-Seq analysis, electroporation with Tox or Tox-Dam vectors induced essentially identical phenotypes (Fig 5A and Supplementary Fig S2C) making it unlikely that chromatin binding was perturbed by the fusion of Tox to Dam. Being Tox an off-switch gene downregulated in DP, we first investigated whether its overexpression altered the proportion of DP. To this aim, we performed electroporation in Btg2RFP reporter mice and found a reduction of Btg2RFP+ cells in the VZ by 30% in Tox-overexpressing brains relative to control (from 37.7 ± 3.1% to 26.4 ± 4.9%, respectively) (Fig 5B). Moreover, and consistent with a decrease in DP, we observed that Tox overexpression reduced the proportion of Tbr2+ newborn basal progenitors in the VZ by 40% while, conversely, increasing it in the SVZ by 10% (32.6 ± 7.1% versus 20.8 ± 0.7% in VZ and 75.9 ± 2.9% versus 85.2 ± 1.4% in SVZ), such that the two effects ultimately compensated each other (Fig 5C). Finally, we found that Tox-overexpressing cells in the VZ expressed significantly higher levels of the proliferative marker of apical progenitors Sox2 (Fig 5D, left). This increase in Sox2 immunoreactivity is intriguing since (i) Tox and Sox2 expression positively correlated in physiological conditions (Fig 1C and C') and (ii) DamID-Seq identified Sox2 as a direct target of Tox, although this single-exon transcript could not be assessed by qRT–PCR. These data show that Tox overexpression inhibits the differentiation of apical progenitors in the VZ. Yet, its overexpression increased primarily progenitors in the SVZ, and not in the VZ (Fig 5A). To address this counterintuitive finding, we inspected the SVZ to assess whether overexpression of Tox promoted the mislocalization of PP from the VZ into the SVZ. As a marker of apical progenitors, we again used Sox2 and found an increase by almost 50% (18.2 ± 1.1% versus 26.4 ± 0.4%) in Sox2+ progenitors in the SVZ (Fig 5D, right). Moreover, considering that cortical progenitors include not only apical and basal progenitors but also a small fraction of basal radial glia (Shitamukai et al, 2011; Wang et al, 2011), we next investigated whether Tox overexpression altered the proportion of one specific progenitor type as identified by their bi-polar, a-polar and uni-polar morphology in mitosis, respectively. To address this, we acquired high-resolution, Z-stack images of p-Vim+ mitotic cells in the SVZ/IZ upon electroporation with control or Tox-expressing vectors. This showed that Tox-overexpressing cells in these layers were neither apical progenitors with extended interkinetic nuclear migration nor newly generated basal radial glia because the almost totality (34/36; i.e. 94%) maintained an a-polar morphology as it was the case in both GFP+ unmanipulated cells or GFP+ electroporated with control plasmids (together 132/138; i.e. 96%) (Fig 5D and data not shown). This was further confirmed by the analysis of interphase, GFP+ cells in the SVZ, with the vast majority displaying a-polar morphology, thus, showing that the increase in the proportion of Sox2+ cells in the SVZ is primarily due to its ectopic expression in basal progenitors (Fig 5D and data not shown). Altogether, these data show that Tox controls neural stem cell fate by inhibiting their switch from PP to DP and decreasing neurogenesis.

**Tox affects neurite outgrowth and neural specification**

Given the enrichment in functional terms related to axonogenesis and neurite outgrowth resulting from DamID-Seq data (Fig 4A; right), we next investigated whether Tox overexpression had any effect on neuronal migration or neurite growth. To address the former, we performed electroporation at E13.5 followed 24 h later by administration of a single dose of BrdU to label progenitors in S phase (Fig 6A). Embryos were collected at E15.5 and proportion and distribution of GFP/BrdU+ cells assessed. First, we found that...
Figure 6. Tox promotes neurite outgrowth and neuronal fate.

A–B’ Experimental layout (A), fluorescence pictures (B) and quantifications (B’; as indicated) plasmids at E13.5 followed by BrdU exposure and sacrifice as shown in (A). Neuronal migration was assessed by calculating the distribution of GFP+/BrdU+ cells within five equidistant bins of the IZ (B and B’, left), while output was quantified as the proportion of GFP+/BrdU+ in the IZ and CP relative to all GFP+ cells (B’; right). n = 2; error bars = SEM. Scale bar, 50 μm.

C, D 3D reconstruction (left) and quantification (right) of newborn (C) or polarized (D) neurons in the IZ or CP, respectively. Individual Z-stacks (0.1 μm optical thickness; green) and computer reconstructions (grey) were used to count cell protrusions (C; arrowheads) or neurite length (D; purple line) that were statistically evaluated (box plots; right, n ≥ 49; ***P < 0.001). Scale bars, 10 μm.

E Fluorescence pictures (left) of the E18.5 mouse cortex upon electroporation of GFP or Tox plasmids at E13.5 followed by immunohistochemistry for GFP (green), Ctip2 (white), Satb2 (red) and DAPI counterstaining (blue). Quantification of Ctip2+/GFP+ cells is shown in either condition (right). n = 3; **P < 0.01; error bars = SD. Scale bar, 50 μm.
the distribution of GFP+/BrdU+ cells in five equidistant bins within the IZ, hence, the pattern of migration of Tox-overexpressing neurons generated 24 h prior to dissection, was essentially identical after GFP or Tox overexpression (Fig 6B and B’, left). Moreover, this experiment allowed us to assess the proportion of neurons generated from a pool of progenitors labelled with BrdU 24 h earlier. Reinforcing our observations on cell fate change (Fig 5), we found a significant decrease in the proportion of GFP+/BrdU+ neurons in Tox-overexpressing brains relative to control by 30% (Fig 5B’, right) (from 50.9 ± 2.1% to 36.6 ± 0.7%).

Having excluded an effect on neuronal migration by Tox, we next investigated its possible effects on process outgrowth. To this aim, we acquired high-resolution Z-stacks images of randomly chosen GFP+ cells within the IZ or CP of GFP or Tox electroporated brains and 3D-reconstructed individual cells to assess their protrusions (Fig 6C and D). Newly born neurons in the IZ are primarily characterized by relatively short processes being dynamically extended and retracted (Kriegstein & Noctor, 2004; Barnes & Polleux, 2009). In contrast, more mature and polarized neurons are characterized by a trailing and a leading process. After reaching the CP, these neurons grow an unipolar dendrite directed towards the extended and retracted (Kriegstein & Noctor, 2004; Barnes & Polleux, 2009). Looking for these two features of newborn and polarized neurons, we found that Tox overexpression induced a doubling (from 3.6 ± 1.6 to 8.0 ± 2.8) in the number of processes of non-polarized neurons in the IZ (Fig 6C). Moreover, independently from the fact that remarkably few neurons could be found in the CP upon Tox overexpression (Fig 5A), the few that were found displayed a substantially longer dendrite (from 21.2 ± 7.2 to 32.10 ± 11.6 μm) (Fig 6D).

Finally, we investigated whether Tox-overexpressing neurons were able to undergo full maturation and specification irrespective of their change in numbers and altered formation of processes. To this aim, we electroporated E13.5 embryos with control or Tox-expressing vectors as previously described but collecting brains at a longer survival time at E18.5. This showed that the majority of Tox electroporated neurons reached the CP (Fig 6E), supporting our conclusion about a lack of phenotype on migration. However, the proportion of GFP+ cells in the CP that expressed Ctip2 decreased after Tox overexpression by ~60% compared to control (from 24.1 ± 3.6% to 8.7 ± 2.6%). In contrast, the overwhelming majority of GFP+ cells in upper and deeper layers remained Satb2+ in either condition. Importantly, this reduction in the proportion of Ctip2+ neurons induced by Tox can be explained either as a consequence of a delayed differentiation of PP or as a cell-intrinsic effect on neuronal specification.

Hence, we propose that Tox is a multifunctional factor involved in corticogenesis by promoting not only proliferative divisions of neural progenitors but also neurite outgrowth and fate of newborn neurons. These functions were predicted by DamID-Seq and likely due to the regulation of different transcriptional targets of Tox in different cell types.

Discussion

Here, we found that the expression of Tox at the tissue level recapitulates the temporal and spatial gradients of cortical neurogenesis. At the cellular level, Tox positively correlated with markers of proliferation while, conversely, markers of differentiation were less abundant in Tox− cells. Possibly underlying these highly specific patterns of expression in tissues and individual cells, we found that Tox is regulated by calcineurin/Nfat4 signalling. Interestingly, both calcineurin and Nfat4 are known as key players implicated in stem cell commitment and regeneration (Crabtree & Olson, 2002; Horsley et al., 2008; Mukherjee & Soto, 2011; Kujawski et al., 2014). Moreover, and possibly underlying the coordination of calcineurin/Nfat signalling, VZ progenitors were described to undergo spontaneous Ca2+ oscillations during development (Owens & Kriegstein, 1998). In this frame, our study provides one additional link between Ca2+ and neurogenesis (Aprea & Calegari, 2012) in which at least some of these effects may act through Nfat4-dependent upregulation of Tox.

Until now, Tox has been studied solely in the immune system and during hematopoiesis (Wilkinson et al., 2002; Aliahmad et al., 2004, 2010; Yun et al., 2011), but even in this context, its downstream targets were unknown and it was unclear whether this transcription factor binds DNA in a sequence- or structure-dependent manner (O’Flaherty & Kaye, 2003; Stros et al., 2007). To address this, we combined powerful experimental and bioinformatical tools developed by other groups (van Steensel & Henikoff, 2000; Vogel et al., 2006, 2007; Southall & Brand, 2007; Wu & Yao, 2013) and performed the first DamID-Seq profiling of a transcription factor. This revealed a level of resolution comparable to ChIP-Seq with the great advantage of avoiding the use of antibodies and giving us not only important information about the properties of DNA binding of Tox, including its motif, but also a comprehensive list of its downstream targets that should be devoid of biases towards any specific lineage. Here we extended the use of DamID as pioneered by other groups (van Steensel & Henikoff, 2000; Orian et al., 2003; Bianchi-Frias et al., 2004; Southall et al., 2013, 2014) and developed DamID-Seq for a transcription factor providing a powerful tool to identify the features of, in principle, any DNA-binding protein. In addition, our list of Tox targets represents an important resource for the study of this transcription factor in, virtually, any tissue.

Since Tox is an off-switch gene, we expected that its targets would also be either off- or on-switches. Yet, we found among Tox targets a mixture of on- and off-switches as well as genes up- or downregulated in PP, DP and/or neurons. This in turn suggests that different genes are differentially regulated by Tox in different cell types, which is supported by the fact that Tox overexpression in the developing cortex could either inhibit or induce its different targets. Corroborating this, DamID-Seq analyses revealed that in addition to the generic terms “transcription factors” and “CNS development” the primary and most significant functional terms associated with Tox targets were “neurogenesis” and “dendrite outgrowth”, which are related to stem cell fate and neuronal maturation, respectively. From this, we propose that Tox is involved in the regulation of two classes of downstream targets in the two cell types, which might depend on the expression of different cofactors in each population (Fig 7).

Finally, we overexpressed Tox by in utero electroporation and investigated its effects during corticogenesis. We found that Tox inhibited the differentiation of neural progenitors resulting in a decrease in neurogenesis. Concerning this, DamID-Seq and qRT–PCR revealed that in certain cases Tox overexpression had a seemingly counterintuitive effect on the expression of specific transcription
factors, specifically Tbr2; in that Tox was found to upregulate Tbr2 despite the fact that this neurogenic factor is expected to be downregulated in a context of decreased neurogenesis. This, however, is easily explained by the fact that the effect of Tox overexpression is the result of the concerted action of all of its targets simultaneously. Hence, upregulation of potent proliferative factors such as Gli1, Yap1 and Sox2 may substantially overcome the concomitant effect on neurogenic factors such as Tbr2. Since Tox expression recapitulates the gradients of neurogenesis in physiological conditions, we propose that Tox upregulation in neurogenic cortical areas is important to maintain a pool of proliferating progenitors among these areas. In contrast, cortical areas in which the neurogenic output is physiologically low, such as caudal/medial cortical regions in early embryos, may not require Tox upregulation, at least until the gradients of differentiation have reached those areas.

Beyond progenitor cells, we observed that newborn neurons migrating through the IZ to reach the CP were characterized by an increase in the number and length of protrusions, respectively. Tox was additionally involved in neuronal fate with a decrease in the proportion of deeper layer neurons. This effect can be due to a delay in neurogenesis resulting from the increased expansion of progenitors and/or by a cell-intrinsic affect in neuronal maturation, which is reminiscent of other transcription factors, such as Brn2 and Cux2 expressed, similar to Tox, in progenitors and subsequently a subset of neurons (Franco et al., 2012; Dominguez et al., 2013).

All together, here we identified Tox as a multifunctional, off-switch transcription factor controlling brain development, neural stem cell differentiation and dendritogenesis. Interestingly, Tox was found among the group of transcription factors regulated by active enhancers of the developing forebrain (Visel et al., 2013; Pattabiraman et al., 2014) and another Tox family member, Tox3, was found to be differentially regulated in neural cell types of the adult sub-ventricular zone (Beckervordersandforth et al., 2010). Hence, these data point to the identification of the Tox family as novel regulator of neural stem cells commitment.

Material and Methods

Constructs

Tox cDNA was generated by RT–PCR of E13.5 cortices (Supplementary Table S1) and cloned into pLgwV5EcoDam (Vogel et al., 2006) or pBI-CMV1 (Clontech) vectors, the latter also used to express GFP from its bidirectional promoter, and used for DamID or in utero electroporation, respectively. The vectors pcdNA3.T7-Δ-Nfat3 (CA-Nfat3) (Addgene plasmid 28224) (Fougere et al., 2010) and pEGFP-RHR4 (CA-Nfat4) (kind gift of Dr. S. Matsuda) (Matsuda et al., 2000) were co-electroporated together with pBI-CMV1-GFP.

Manipulation in mice

In utero electroporation was performed as previously described (Artegiani et al., 2012) by injecting ~1–5 µg of DNA into C57BL/6J or B6.GaTAP mice followed by the delivery of 6, 50 ms long, electric pulses of 30 V with intervals of 1 s. Embryos were electroporated at E13.5 and sacrificed 48 h later eventually upon intraperitoneal administration of 1 mg BrdU (1 pulse at E14.5) or 0.5 mg of CsA (two injections per day) (dissolved in 100 µl PBS or 400 µl of NaCl 0.9%, respectively).

Biochemistry

Electroporated areas of the brain were dissociated (MACS dissociation kit; Miltenyi), cells washed with PBS and incubated 15 min at 4°C in hypotonic solution (Hepes 10 mM, MgCl2 1.5 mM, KCl 10 mM, DTT 0.5 mM and Complete Protease Inhibitors Cocktail...
(CPIC; Roche), and extracts centrifuged at 16,000 g for 15 min at 4°C to separate cytoplasmic and nuclear fractions. The latter was further resuspended in nuclear extraction buffer (Hepes 20 mM, MgCl2 1.5 mM, NaCl 420 mM, EDTA 0.2 mM, DTT 0.5 mM, glyc erol 25% and CPIC) for 30 min at 4°C and centrifuged as above. Extracts were stored at ~80°C and 20 µg analysed after boiling in LDS buffer (Invitrogen) using NuPage 4–12% Bis-Tris gels (Invitrogen) and transferred to nitrocellulose. Membranes were blocked 1 h at room temperature in 3% BSA in TBS 0.3% Tween and incubated overnight with 1:1,000 Tox (Abcam), Gadph (NovusBiological) or Tbp (Abcam) antibodies followed by washing, incubation with HRP-conjugated secondary antibody for 2 h and chemiluminescence reaction (ECL Dura or Pico, Pierce). Non-saturated radiograms were analysed using “Gels” on Fiji 9 software.

Immunohistochemistry

Immunohistochemistry was performed as described (Lange et al., 2009; Ar tegiani et al., 2011). Briefly, after fixation (4% PFA), cryo- protection (30% sucrose) and cryosectioning, slices (10 µm thick) were permeabilized and blocked (0.3% Triton X-100 and donkey serum 10% in PBS) (IHC buffer) for 2 h at room temperature. All primary antibodies are described in Nonaka-Kinoshita et al (2013) except for anti-Tox (Atlas Antibodies 1:500) and incubated overnight at 4°C in IHC followed by 2-h washing in PBS and 3-h incubation with DyLight-secondary antibodies (1:1,000; Jackson Laboratory) at room temperature. DAPI was used to counterstain nuclei. ApoTome (Zeiss) composite images were acquired and processed by AxioVision or Zen (Zeiss). Assessment of immunoreactivity within individual cells was performed by outlining randomly chosen, DAPI-labelled nuclei and quantifying pixels’ density in the relevant channel. 3D reconstruction of randomly selected neurons was performed using 40 or 63× objectives and 0.1 µm Z-stacks. Analyses were performed on digital images using Photoshop (Adobe) and/or Fiji 9 (http://fiji.sc/Fiji).

DamID-Seq

DamID was performed according to Vogel et al (2007) using lentivirus es obtained as previously described (Artegiani et al., 2012, 2013) but without concentration. HEK-293T or Neuro-2a cells were infected with viral supernatants diluted 1:2 (or even 1:10 in the case of Neuro-2a), and 48 h later, their genomic DNA was extracted, digested with DpnI, ligated to adaptors, further digested with DpnII and PCR-amplified. Sequencing libraries were prepared as described (Aprea et al, 2013) and subjected to 75-bp single-read sequencing on a HiSeq 2000 platform (Illumina), resulting in ~20 million reads per sample. Sequencing raw data were deposited in GEO.

Bioinformatics

Reads alignment to the human genome (Ensembl67) was performed with bowtie (v0.12.7) using “-b best” and “-m 1” to report uniquely mapped reads. Replicate reproducibility was tested by bamCorrelate from deepTools (Ramirez et al, 2014) and alignments of replicates pooled before peak calling. Genomic regions of Tox-Dam enrichment were identified by SICER (Zang et al, 2009; Wu & Yao, 2013) considering genes within 15 kb upstream of the TSS, or inside the transcript body. Peak annotation with genomic features was done by ChiPseeker (Bioconductor). For Tox read profile around the TSS, aligned reads were normalized to Dam using bamCompare (Ramirez et al, 2014) setting the parameters: “–ratio; –normalizeTo1x 245196000; and –scaleFactorsMethod read-Count”. Cell-specific enhancers (Andersson et al, 2014) were converted from hg18 to hg19 using liftOver (UCSC) and Tox read density calculated by computeMatrix scale regions (GitHub), scaling to the average size of enhancers (3 kb). Conservation scores for placental species (phastCons46way) were downloaded from the UCSC genome browser and exonic regions masked. Matched random genomic locations to the Tox peaks were created with shuffleBed (bedTools) using the parameters: “–seed 927442958; -chrom; -noOverlapping”. Tox putative binding motif was predicted by gimmeMotifs clustering results from several other softwares (van Heeringen & Veenstra, 2011) and using 1,000 peaks with ≥ 2.5-fold enrichment and ≤ 1 Kb.

qRT–PCR

Electroporated cells were dissociated as described above and GFP+ cells sorted using a FACSaria III (Becton Dickinson). RNA was extracted using the RNeasy micro kit (QIAGEN) and cDNA generated with Superscript kit (Invitrogen) starting with 100 ng of RNA and using 1 µl of cDNA diluted 1:10 for qPCRs. Primers (Supplementary Table S1) were designed to span across exons to distinguish cDNA from genomic DNA.

Statistical analysis

All the analyses presented in this work were performed with at least three biological replicates (n ≥ 3) except for DamID-Seq (n = 2). Values obtained were used to calculate mean, standard deviation and significance by two-tailed Student’s t-test, assuming 0.05 as a threshold for significance. Quantifications of individual cells were performed using a minimum of 24–80 cells as indicated in each figure panel.

Supplementary information for this article is available online: http://emboj.embopress.org

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Author contributions

BA and FC planned the experiments and wrote the manuscript; BA performed all experiments supported by SBA, EB and SM; and AMJD performed all bioinformatic analyses supported by AD. All authors approved the manuscript.

Conflict of interest

The authors declare that they have no conflict of interest.
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