Developmental Neurotoxicity Testing: Recommendations for Developing Alternative Methods for the Screening and Prioritization of Chemicals

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Summary

Developmental neurotoxicity testing (DNT) is perceived by many stakeholders to be an area in critical need of alternative methods to current animal testing protocols and guidelines. An immediate goal is to develop test methods that are capable of screening large numbers of chemicals. This document provides recommendations for developing alternative DNT approaches that will generate the type of data required for evaluating and comparing predictive capacity and efficiency across test methods and laboratories. These recommendations were originally drafted to stimulate and focus discussions of alternative testing methods and models for DNT at the TestSmart DNT II meeting (http://caat.jhsphs.edu/programs/workshops/dnt2.html) and this document reflects critical feedback from all stakeholders that participated in this meeting. The intent of this document is to serve as a catalyst for engaging the research community in the development of DNT alternatives and it is expected that these recommendations will continue to evolve with the science.

Keywords: developmental neurotoxicity, screening, in vitro models

1 Introduction

The development of alternative test methods has been widely acknowledged as a critical need for toxicity testing (NRC, 2007). Two major issues are driving this need. The first is the need to provide more efficient testing methods that can provide hazard information for the thousands of untested chemicals currently used in commerce (Andersen and Krewski, 2009; Kavlock et al., 2009). The second is a need to reduce the use of animals in toxicity testing (Goldberg, 2002; Balls, 2009).

Over the past two decades, expert panels have provided criteria and guidelines for validating new testing methods. The most prominent of these are the ICCVAM guidelines on validation of alternative test methods (ICCVAM, 1997). These guidelines provide a framework for assessing the applicability domain and performance criteria of new methods, which are especially critical if the method is to be used in a regulatory context. These guidelines, however, have been criticized for implementing a tedious and lengthy process that may actually impede efficient adoption and use of alternative test methods. The need to validate the predictive nature of these methods has required the collection of extensive and expensive data sets for regulatory acceptance (Hartung, 2007). An alternative approach is to use in vitro and QSAR methods for prioritizing available data...
Developmental neurotoxicity testing is an area that is widely recognized as in need of alternative methods (Coecke et al., 2007; Lein et al., 2005; Lein et al., 2007; Bal-Price et al., 2010; Aschner et al., 2010). One issue that has restricted progress in development of new alternative methods for DNT is that funding is skewed heavily towards research on basic biological and toxicological mechanisms. This has led to the development of a wide variety of methods (Coecke et al., 2007) that are not necessarily amenable to testing large numbers of chemicals. Organotypic cultures, while providing a good model for 3-dimensional tissue organization (Sundstrom et al., 2005), also require animals for tissue harvesting and are low-throughput (Coecke et al., 2007).

The current document provides a set of principles, which, if embraced by the larger research community, will enhance the development of alternative test methods suitable for screening of large numbers of chemicals. It includes recommendations to facilitate development of alternative testing methods for screening substances for potential developmental neurotoxicity. These recommendations are not intended to be used for validation of test methods and should not be used to circumvent or substitute for any existing test method validation criteria (e.g., ICCVAM, 1997; Hartung et al., 2004; OECD, 2005). Test method validation for regulatory use involves a series of specific stages that commence with development of a method, and then proceed sequentially through test method optimization and standardization; protocol transferability assessment; studies in multiple labs to establish reliability, specificity and sensitivity; and finally peer review and regulatory review for acceptance into a regulatory framework. In contrast, this document primarily focuses on the early stages of this process: research and development, protocol optimization and protocol standardization. Taken together, these research efforts will develop sufficient data to demonstrate “proof of principle” that the test method performs adequately for the intended propose and to facilitate the comparison of data between laboratories. A favorable review of a test method’s performance at the “proof of principle” stage paves the way for moving forward towards developing the data needed for regulatory acceptance.

The goal of this document is to engage the research community in the process of developing alternative methods for developmental neurotoxicity that are amenable to high-throughput screening. The primary focus is to provide recommendations and guidance to catalyze development of a suite of assays that can be used for prioritization. Prioritization would differentiate substances that are of high concern (and may need further evaluation for a particular toxicity pathway or endpoint) from substances that are of lower concern (by virtue of having lower potential for exerting biological actions on relevant physiological or pathological processes). These same test methods could be considered for use in screening drugs or chemicals prior to commercialization and to aid in grouping commercial chemicals for read-across, or replacement of in vivo animal testing for initial hazard screening.

The rest of this document provides definitions for important terms that describe test systems and a series of recommendations that can be used when developing new alternative test methods. Consideration of the recommendations made herein, while important for development of all test methods, should assist in the transition of methods from early development stages to use in screening and validation efforts. For each recommendation there are examples and references.

2 Definitions

For the purpose of this draft we defined the following terms:

1. Endpoint (E): the biological or chemical process, response, or effect assessed by a test method (OECD Guidance Document 34, 2005)
2. Test system (TS): any animal, cellular, subcellular, chemical, or physical system, or a combination thereof, used in a study (modified from OECD, GLP principle, directive 87)
3. Test method (TM): A process or procedure used to obtain information on the biological effects of a substance or agent. Toxicological test methods generate information regarding the ability of a substance or agent to produce a specified biological effect under specified conditions. Used interchangeably with “test” and “assay.” (OECD Guidance Document 34, 2005). The test method should assess one or more key aspects of human neurodevelopment.

3 Guidance

This guidance was drafted to stimulate and focus discussions of alternative testing methods and models for DNT at the TestSmart DNT II meeting (http://caat.jhsph.edu/programs/workshops/dnt2.html), and the current version reflects critical feedback from all stakeholders that participated in this meeting. In the present document, PC12 cells as a test system and neurite outgrowth as an endpoint will be used as examples. This is not intended to be exclusionary of other test methods or approaches. Table 1 provides an overview of the 15 guidance items for developing alternative screening test methods for developmental neurotoxicity.

1. Key event of neurodevelopment (TS, TM, E). Test methods should incorporate one or more endpoints that model key aspects of human neurodevelopment.
4. Parametric controls (E). Assay parameters that result in predictable changes in the endpoint should be characterized. These experimental parameters can be used to optimize the test method. Examples: i) Increasing nerve growth factor (for NS-1 cells) or retinoic acid (for SH-SY5Y cells) concentration will increase neurite outgrowth; ii) Increasing days in culture yields greater neurite growth; iii) Cell density influences neurite outgrowth. Reference: Radio and Mundy, 2008

5. Response characterization (E). The level of change in the response associated with an effect should be characterized. This is the degree of change that, if exceeded, results in a positive response (a “hit”). Importantly, one needs to have a fairly robust understanding of the variability in the control response levels in order to interpret results. Generally, there are two ways to determine the positive response level. The first approach, commonly used in pharmaceutical screening, defines a hit as any response greater than 3 SD from the control. This conservative statistical approach is used to ensure a very small number of false positives: false positives would be costly to pursue. In toxicological screening and prioritization for further testing, it may be acceptable to have a higher rate of false positives. Thus, a second approach defines a positive response level based on biological significance. Professional

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**Tab. 1: Overview of the 15 guidance items for developing alternative screening test methods for developmental neurotoxicity.**

(See text for details and definitions)

| Methods Area                      | Method Elements | Description                                                                                                                                 |
|-----------------------------------|-----------------|--------------------------------------------------------------------------------------------------------------------------------------------|
| 1 Key Event of Neurodevelopment   | TM, TS, E       | Endpoints should model key aspects of neurodevelopment                                                                                     |
| 2 Endpoint Measurement            | TM, E           | Correct and accurate measurement of the endpoint                                                                                           |
| 3 Dynamic Range                   | E               | Determination of the extent of measurable change                                                                                           |
| 4 Parametric Controls             | E               | Assay parameters that predictably change the endpoint                                                                                       |
| 5 Response Characterization       | E               | Level of change determined to be an effect                                                                                                  |
| 6 Concentration                   | TS, E           | Methods must be designed to allow determination of concentration-response                                                                  |
| 7 Endpoint Selectivity            | E               | Discrimination of the endpoint of concern from non-specific outcomes                                                                      |
| 8 Endpoint-Selective Controls      | E               | Chemicals known to reliably and consistently alter the endpoint at a mechanistic level                                                     |
| 9 Training Set Chemicals          | TM, E           | Goal is proof-of-concept that the test method can rapidly and efficiently screen moderate numbers of chemicals. Should include chemicals known to reliably elicit a response, or no response, based on *in vitro* findings. |
| 10 Testing Set Chemicals          | TS, TM, E       | Goal is to demonstrate ability to test large number of chemicals. Should include chemicals known to affect, and lack effects, *on in vivo* developmental neurotoxicity endpoints. |
| 11 Specificity and Sensitivity    | TM, E           | Analysis to determine ability to correctly differentiate active and non-active chemicals                                                   |
| 12 High Throughput                | TS, E           | Test system and endpoint should be amenable to automation                                                                                   |
| 13 Documentation                  | TM              | Full and published documentation of the test method                                                                                           |
| 14 Transferability                | TS, E           | Resources for use should be available for any laboratory                                                                                    |
| 15 Data Sharing                   | Data            | Open access databases are highly desirable.                                                                                                 |

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Example: Neurite outgrowth measurement in differentiating PC-12 cells. The test method models neurite outgrowth, the test system employs differentiating PC-12 cells, and the endpoints include measurement of neurite length, number of neurites, number of branch points, etc. References: Pool et al., 2008; Kim et al., 2007; Lein et al., 2005; Cоеcke et al., 2007

2. Endpoint measurement (TM, E). All test methods must demonstrate the ability to correctly and accurately measure the intended endpoint (see Sections 4 and 8). Example: Automated image analysis of GFP-tagged neurofilament as a biomarker for neurite outgrowth in PC12 cells should yield similar results as *camera lucida* drawings of neurite outgrowth in PC12 cells (i.e., does automated read-out really reflect increased neurite outgrowth or increased cell proliferation?) References: Sette, 1987; Cоеcke et al., 2007

3. Characterization of dynamic range (E). One should determine the extent of change that can be detected for a DNT endpoint and whether both increases and decreases from untreated control can be measured. Example: Both increased and decreased neurite outgrowth can be assessed in PC12 cells. Reference: Endo et al., 2008

4. Parametric controls (E). Assay parameters that result in predictable changes in the endpoint should be characterized. These experimental parameters can be used to optimize the test method. Examples: i) Increasing nerve growth factor (for NS-1 cells) or retinoic acid (for SH-SY5Y cells) concentration will increase neurite outgrowth; ii) Increasing days in culture yields greater neurite growth; iii) Cell density influences neurite outgrowth. Reference: Radio and Mundy, 2008

5. Response characterization (E). The level of change in the response associated with an effect should be characterized. This is the degree of change that, if exceeded, results in a positive response (a “hit”). Importantly, one needs to have a fairly robust understanding of the variability in the control response levels in order to interpret results. Generally, there are two ways to determine the positive response level. The first approach, commonly used in pharmaceutical screening, defines a hit as any response greater than 3 SD from the control. This conservative statistical approach is used to ensure a very small number of false positives: false positives would be costly to pursue. In toxicological screening and prioritization for further testing, it may be acceptable to have a higher rate of false positives. Thus, a second approach defines a positive response level based on biological significance. Professional
judgment should be used to balance the biological and statistical relevance of the response level.
References: Tierno et al., 2007; Radio and Mundy, 2008; Breier et al., 2008

6. Concentration range (TM, E). Each test method should be designed to characterize the concentration-response relationship. One recommendation is to minimally test five concentrations ranging from the solubility limit to five logs below the solubility limit. Concentration-response is critical to comparison of sensitivity between test methods, or endpoints within a test method. Concentrations above the level known to induce cytotoxicity should not be used.

7. Endpoint selectivity (E). The ability of the test method to discriminate the endpoint of concern from other outcomes. Example: The ability to determine the concentration-relationships for both cytotoxicity and the endpoint in the same assay provides assurance that chemical-induced endpoint changes do not primarily result from cell death (Fig. 1).
References: Schmuck et al., 2000; Cristòfol et al., 2004

8. Endpoint-selective controls (TM, E). Endpoint-selective control chemicals reliably and consistently alter the endpoint by known mechanisms. Both positive and negative control chemicals should be tested. A positive control is a chemical or stressor which is known from previous experience to reliably affect the endpoint. A negative control is a chemical that reliably causes no effect on the endpoint of interest. A negative control demonstrates the base-line result obtained when a test chemical does not produce a measurable positive result.

Examples: i) NGF-induced neurite outgrowth in PC12 cells requires MAP kinase signaling, therefore the MAP kinase inhibitor U0126 could be used as a positive control; ii) NGF-induced neurite outgrowth in PC12 cells does not involve signaling via the JAK/STAT signaling pathway, thus a JAK/STAT inhibitor could be used as a negative control.
Reference: Radio et al., 2008

9. Training set of chemicals (TS, TM, E). Once the method has been demonstrated to exhibit the correct characteristics described above (see sections 1-8), a “training set” of chemicals should be developed and tested. This training set should be composed of two types of chemicals: chemicals that are known to reliably elicit a response of concern (needed to assess sensitivity) and chemicals that are known to reliably elicit no response of concern (needed to assess specificity). Evidence for an effect, or lack thereof, should come from in vitro data. However, additional evidence from in vivo studies, if available, is highly recommended. Selection of the training set of chemicals should also consider the purpose of the assay, as discussed above in the introduction. The goal of the training set is to evaluate the test method, including: 1) testing the practical ability of the method to efficiently process moderate numbers of chemicals; 2) confirming positive and negative controls; and 3) generating historical control data to characterize the inherent response range for the endpoint.
References: Radio et al., 2008; Breier et al., 2008

10. Testing set of chemicals (TS, TM, E). The testing set of chemicals should include a large number of substances
known to affect endpoints of developmental neurotoxicity in vivo, as well as chemicals that reliably do not affect developmental neurotoxic endpoints. For in vitro screening assays (as opposed to replacement assays), it is important at this stage to demonstrate the ability of the method to rapidly and efficiently screen large numbers of chemicals with an adequate degree of sensitivity and specificity, and to provide data that can be used in determining future steps in the process of method development, validation, and regulatory acceptance of the endpoint (E) and test method.

An example of a list of chemicals with demonstrated effects on neurotoxicity endpoints in vivo is provided in Table 2. Reference: Padilla et al. (unpublished)

11. Specificity and Sensitivity (TM, E). Sensitivity is defined as the proportion of active substances that are correctly identified by the new test, and specificity is defined as the proportion of inactive substances that are correctly identified. Positive and negative predictivity are the frequencies of correct predictions obtained from the new test.

References: Cooper et al., 1979; ICCVAM, 1997

12. High throughput (TS, E). In the case of in vitro screening assays, it is highly desirable for the test method to have the potential for automation. For any new, revised, or replacement assay, it is critical that the method be more efficient than the current testing scheme (OECD Test Guideline 426, 2007). Testing one chemical with this guideline can take up to 6 months, require hundreds of animals, and cost hundreds of thousands of dollars. Automation of the method could lead to testing of hundreds or even thousands of chemicals for one endpoint in one day.

13. Documentation (TM). For any test method published with the intent to demonstrate feasibility for screening large numbers of chemicals, the test method needs to be fully documented and readily available to allow for implementation across laboratories. Experimental details critical to replication of methods must be included. This kind of information

**Tab. 2: Draft list of a table of chemicals to consider when developing new test methods for developmental neurotoxicity.**

Chemicals on this list were derived from published reports or regulatory data from humans, non-human primates, or laboratory mammals. Findings were deemed to be suggestive of adverse neurological outcomes following developmental exposure. To be included on the list there must be positive results from more than one laboratory (Mundy et al., 2009).

| Acrylamide | Diamorphine hydrochloride | Naloxone |
| Aldicarb | Diazepam | Naltrexone |
| Allethrin | Diazinon | Nicotine |
| Aluminum | Diethylstilbestrol | Parathion |
| Aminonicotinamide, 6- | Epidermal Growth Factor | PCBs |
| Amphetamine, d- | Ethylenetriurea | Permethrin |
| Aspartame | Fluourouracil, 5- | Phenylacetate |
| Azocytidine | Haloperidol | Phenylalanine |
| Benomyl | Halothane | Phthalates |
| Benzene | Heptachlor | Propylthiouracil |
| Bioallethrin | Hexachlorobenzene | Salicylate |
| Bis(tri-n-butyltin)oxide | Hydroxyurea | Tellurium |
| Butylated hydroxyanisol | Iminodiproprionitrile | Thalidomide |
| Butylated hydroxytoluene | Lindane | Toluene |
| Carbamazepine | LSD | Triamcinolone |
| Carbon monoxide | Maneb | Tributyltin chloride |
| Chlordecone | Methadone | Trichlorfon |
| Chlordiazepoxide | Methanol | Trichloroethylene |
| Chlorine dioxide | Methimazole | Triethyllead |
| Chlorpromazine | Methoxysteranol, 2- | Triethyltin |
| Colcemid | Methylazoxymethanol Monosodium glutamate | Trimethyltin |
| Colchicine | | Trypan blue |
| Cytosine arabinoside | | Urethane |
| DEET | | Urethane |
| | | Vincristine |
can be within published reports or as supplementary files available from the publisher’s website.

14. Transferability (TS, E). The required resources need to be accessible and widely available to allow for implementation across laboratories. For example, proprietary cell lines not available to other researchers would not be acceptable. Equipment should be commercially available or well documented (see #13).

Example: Commercially available ArrayScan™ technology and a specialized PC12 cell line (ThermoScientific) for use in neurite outgrowth assays.

Reference: Radio et al., 2008

15. Data sharing through open access databases is highly desirable. It is extremely important for data from testing methods to be openly reported in publicly accessible databases (Gruber and Hartung, 2004). This will allow inter-laboratory and intra-laboratory comparisons of test methods.

Examples include:

   i) PubChem website: http://pubchem.ncbi.nlm.nih.gov
   ii) AltTox website: http://www.alttox.org/ttc/way-forward
       iii) ToxCast™ website: http://www.epa.gov/nct/toxcast

4 Conclusion

Alternative test methods with potential for developmental neurotoxicity screening will require demonstration of their predictive capacity and this will entail development and acceptance of a strategy for data interpretation. These two issues, prediction and interpretation, cannot be determined without the generation of data. This document provides guidance for methods development and data collection. Consideration of the recommendations herein are important for all new testing methods and should assist in the transition of methods from the early development stages to use in screening and validation efforts. Hopefully, this document will also stimulate the generation of data needed to begin devising strategies to prioritize chemicals for further testing.

References

Andersen, M. E. and Krewski, D. (2009). Toxicity testing in the 21st century: Bringing the vision to life. Toxicol. Sci. 107, 324-330.

Aschner, M., Crofton, K. M. and Levin, E. D. (2010). Introduction to the special issue on emerging high throughput and complementary model screens for neurotoxicology. Neurotoxicol. Teratol. 32, 1-3.

Bal-Price, A. K., Hogberg, H. T., Buzanska, L. et al. (2010). In vitro developmental neurotoxicity (DNT) testing: Relevant models and endpoints. Neurotoxicology 31, 545-554.

Balls, M. (2009). The origins and early days of the Three Rs concept. ATLA 37, 255-265.

Breier, J. M., Radio, N. M., Mundy, W. R. and Shafer, T. J. (2008). Development of a high-throughput screening assay for chemical effects on proliferation and viability of immortalized human neural progenitor cells. Toxicol. Sci. 105, 119-133.

Coecke, S., Goldberg, A. M., Allen, S. et al. (2007). Workgroup report: incorporating in vitro alternative methods for developmental neurotoxicity into international hazard and risk assessment strategies. Environ. Health Perspect. 115, 924-931.

Cooper, J. A., 2nd, Saracci, R. and Cole, P. (1979). Describing the validity of carcinogen screening tests. Br. J. Cancer 39, 87-89.

Cristófol, R. M., Gasso, S., Vilchez, D. et al. (2004). Neurotoxic effects of trimethyltin and triethyltin on human fetal neuron and astrocyte cultures: a comparative study with rat neuronal cultures and human cell lines. Toxicol. Lett. 152, 35-46.

Endo, Y., Beauchamp, E., Woods, D. et al. (2008). Wnt-3a and Dickkopf-1 stimulate neurite outgrowth in Ewing tumor cells via a Frizzled3- and c-Jun N-terminal kinase-dependent mechanism. Mol. Cell Biol. 28, 2368-2379.

Gruber, F. P. and Hartung, T. (2004). Alternatives to animal experimentation in basic research. ALTEX 21, Suppl. 1, 3-31.

Goldberg, A. M. (2002). Use of animals in research: a science–society controversy? The American perspective: animal welfare issues. ALTEX 19, 137-139.

Hartung, T. (2007). Food for thought ... on validation. ALTEX 24, 67-80.

Hartung, T., Bremer, S., Casati, S. et al. (2004). A modular approach to the ECVAM principles on test validity. ATLA 32, 467-472.

ICCVAM (1997). Validation and regulatory acceptance of toxicological test methods: A report of the ad hoc interagency coordinating committee on the validation of alternative methods. http://iccvam.niehs.nih.gov/docs/about_docs/validate.pdf

Judson, R. S., Houck, K. A., Kavlock, R. J. et al. (2010). In vitro screening of environmental chemicals for targeted testing prioritization: the ToxCast project. Environ. Health Perspect. 118, 485-492.

Kavlock, R. J., Austin, C. P. and Tice, R. R. (2009). Toxicity testing in the 21st century: Implications for human health risk assessment. Risk Anal. 29, 485-487.

Kim, J.-H., Ha, H.-C., Lee, M.-S. et al. (2007). Effect of Tremella fuciformis on the neurite outgrowth of PC12h cells and the improvement of memory in rats. Biol. Pharm. Bull. 30, 708-714.

Lein, P., Silbergeld, E., Locke, P. and Goldberg, A. M. (2005). In vitro and other alternative approaches to developmental neurotoxicity testing (DNT). Environ. Toxicol. Pharmacol. 19, 735-744.

Lein, P., Locke, P. and Goldberg, A. (2007). Meeting report: Alternatives for developmental neurotoxicity testing – Test-Smart developmental neurotoxicology. Environ. Health Perspect. 115, 764-768.

Mundy, W., Padilla, S., Shafer, T. et al. (2009). Building a database of developmental neurotoxicants: Evidence from human and animal studies. Toxicologist 108, 284.
National Research Council – NRC (2007). Toxicity testing in the 21st century: A vision and a strategy. Washington, DC: National Academy Press.
OECD (2005). OECD Series on Testing and Assessment, Number 34: Guidance document on the validation and international acceptance of new or updated test methods for hazard assessment. Organization for Economic Cooperation and Development. ENV/JM/MONO(2005)14, 18 Aug 2005, Paris.
OECD (2007). Test Guideline 426. OECD Guideline for Testing of Chemicals. Developmental Neurotoxicity Study. http://www.oecd.org/document/55/0,3343,en_2649_34377_2349687_1_1_1_1,00.html
Parran, D. K., Mundy, W. R. and Barone, S. Jr. (2001). Effects of methylmercury and mercuric chloride on differentiation and cell viability in PC12 cells. Toxicol. Sci. 59, 278-290.
Pool, M., Thiemann, J., Bar-Or, A. and Fournier, A. E. (2008). NeuriteTracer: a novel ImageJ plugin for automated quantification of neurite outgrowth. J. Neurosci. Methods 168, 134-139.
Radio, N. M., Breier, J. M., Shafer, T. J. and Mundy, W. R. (2008). Assessment of chemical effects on neurite outgrowth in PC12 cells using high content screening. Toxicol. Sci. 105, 106-118.
Radio, N. M. and Mundy, W. R. (2008). Developmental neurotoxicity testing in vitro: models for assessing chemical effects on neurite outgrowth. Neurotoxicology 29, 361-376.
Schmuck, G., Ahr, H. J. and Schluter, G. (2000). Rat cortical neuron cultures: an in vitro model for differentiating mechanisms of chemically induced neurotoxicity. In Vitro. Mol. Toxicol. 13, 37-50.
Schulte, P. A. (1989). A conceptual framework for the validation and use of biologic markers. Environ. Res. 48, 129-144.
Sette, W. F. (1987). Complexity of neurotoxicological assessment. Neurotoxicol. Teratol. 9, 411-416.
Sundstrom, L., Pringle, A., Morrison, B. and Bradley, M. (2005). Organotypic cultures as tools for functional screening in the CNS. Drug Discovery Today 10, 993-1000.
Tierno, M. B., Johnston, P. A., Foster, C. et al. (2007). Development and optimization of high-throughput in vitro protein phosphatase screening assays. Nat. Protoc. 2, 1134-1144.

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