



To: Ambassador Miriam Sapiro, Deputy US Trade Representative
 Daniel Mullaney, Assistant US Trade Representative for Europe and the Middle East
 Boris Bershteyn, Acting Administrator, Office of Information and Regulatory Affairs
 Daniel Calleja Crespo, Director General, DG Enterprise and Industry
 Jean-Luc Demarty, Director General, DG Trade

Re: Response to Sept. 2012 Joint Solicitation by US-EU High-Level Regulatory Cooperation Forum

Date: 31 October 2012

These comments are submitted on behalf of Humane Society International and The Humane Society of the United States and our more than 11 million members across North America, Europe and the globe in response to the USTR comment request of 28 September 2012 (77 Fed. Reg. 59702). We appreciate the initiative of the US-EU High-Level Regulatory Cooperation Forum to promote greater transatlantic regulatory compatibility across economic sectors, and would like to highlight one sector in particular where we believe there is room for concrete, near-term progress—the pesticide sector.

Regulatory data requirements for pesticide registration differ somewhat between European, U.S. and other global markets, such that companies are generally unable to prepare a single registration dossier that will satisfy authorities in different regions. *In practical terms, this can lead to duplicative testing costs and delays in market access.* Accordingly, we are submitting these comments to highlight key areas that can be improved and we are providing recommendations as to how this can be done.

A 2010 study commissioned by the European Crop Protection Association and CropLife America, which compares the costs of new crop protection product discovery and development between 2000 and 2008, reflects an cost escalation of 39.1% (US\$184 million to \$256 million) over this period. The most dramatic cost increases (77.7% and 116%) are attributable to regulatory testing for product registration, i.e., toxicological safety assessment and field trials, respectively (*www.croplife.org/view_document.aspx?docId=2478*).

It should also be noted that from an animal welfare perspective, upwards of 10,000 dogs, rabbits and other animals can be used in dozens of separate toxicological and ecotoxicological studies to satisfy regulatory data requirements for a single active ingredient. For many endpoints, there is substantial redundancy among *in vivo* data requirements, e.g., acute toxicity testing by up to 3 different exposure routes (oral, dermal, inhalation) for both active ingredients and finished products, subchronic toxicity testing using both dogs and rats, developmental toxicity testing using both rabbits and rats, etc.

Recent and ongoing revisions to EU regulations for biocides and plant protection products are leading to the uptake of numerous state-of-the-art toxicological testing methods and strategies, which maintain a high level of regulatory rigor to protect human health and the environment while making substantial strides toward elimination of unwarranted redundancies, cost savings for industry, and the replacement, reduction and refinement (3Rs) of vertebrate animal testing (*http://eusaat.org/images/*

Celebrating Animals | Confronting Cruelty Worldwide 2012/presentations/seidle_troy_2012_09_07_pesticide_regulation_eusaat_2012.pdf).

	European Union	United States	
Agency / Directorate General	DG Environment	Environmental Protection	
C .	European Chemicals Agency	Agency	
Relevant regulatory / statutory	Annexes II-III of Regulation	40 CFR § 158W	
provisions	(EU) No. 528/2012	-	
Regulatory differences & their	In June 2012, the EU replaced its former Biocidal Products Directive		
negative effects on stakeholders	98/8/EC with a new Biocidal Products Regulation cited above.		
	Among other changes, the information requirements specified in		
	Annexes II and III for registration of new active ingredients and		
	formulated products have been substantially amended to reflect		
	scientific best practices in the toxicology and ecotoxicology based on		
	the latest test guidelines and guidance	ce promulgated by the	
	Organization for Economic Coopera	ation and Development (OECD)	
	and others. These regulatory amendments allow for a more flexible		
	and efficient approach to safety testing that reduces both economic		
	costs to industry and welfare costs to animals while maintaining a		
	high level of regulatory scrutiny for the protection of human health		
	and the environment. A cursory overview of changes to specific		
	regulatory endpoints is provided below and at <i>http://eusaat.org/</i>		
	images/2012/presentations/seidle_troy_2012_09_0/_		
	pesuciae_regulation_eusaat_2012.paj:		
	1. Acute systemic toxicity, through conditional waiving of the		
	 Carcinogenicity, through the conditional waiving of the mouse bioassay 		
	 Subchronic toxicity, through in assessments in lieu of standalor genotoxicity endpoint 	clusion of micronucleus ne <i>in vivo</i> testing for this	
	4. Calculation approaches for haza products based on the propertie	ard classification of finished s of their constituent ingredients	
	 Skin sensitization, through acce lymph node assay where an ass required 	eptance of the reduced local essment of potency is not	
	 Fish acute toxicity, through use strategy 	of the threshold approach/tiered	
	7. Reproductive toxicity, through extended 1-generation study	the adoption of the new OECD	
	8. Developmental toxicity, throug rat teratogenicity study where the rodent reproductive toxicity stu- effects on fertility or development	h the conditional waiving of the he rabbit study and a generational dy reveal no signs of adverse ent	
	9. Avian reproduction, through wa	aiving of the study requirement	

1. NON-FOOD ANTIMICROBIAL PESTICIDES / BIOCIDES

when the dietary LC_{50} is in excess of 2000 mg/kg.

	In the best-case scenario, these amendments could reduce animal testing by approximately 40%, with substantial cost savings as well. However, such savings can only be realized if regulations in other major markets are aligned with the EU's revised data requirements. In the worst case, if there is no move toward alignment with new EU requirements, industry could be forced to double-test in some instances, e.g., non-animal test for the EU market and the conventional animal test for the US, leading to a relative escalation in testing costs
Possible solutions	In the interests of minimizing redundant testing and preventing undue costs, market access delays and animal use, we urge the US to act swiftly to bring its registration data requirements into alignment with the new state-of-the-art embodied in the EU biocides regulation. We understand from bilateral discussions with EPA's Office of Pesticide
	Programs that the 158W rulemaking for antimicrobial pesticides has not yet been finalized, thus providing the US with an ideal route for enhancing regulatory alignment with the EU in a manner that will benefit all concerned stakeholders.

	European Union	United States	
Agency / Directorate General	DG Health & Consumers	Environmental Protection	
	(SANCO)	Agency	
	European Food Safety Authority		
Relevant regulatory / statutory	Regulation (EC) No. 544/2011 &	40 CFR § 158	
provisions	Regulation (EC) No. 545/2011		
	to be amended in 2013 by		
	SANCO/11802/2010 Rev. 7		
	(POOL/E3/2010/11802/11802R7-		
	EN.doc)		
Regulatory differences & their	The EU is currently in the advanced stages of revising its data requirements for plant protection products, i.e., food-use pesticides. In July of this year, we were informed that the revised requirements detailed in the above mentioned SANCO document were received favorably by the EU Member State Standing Committee on the Food Chain and Animal Health, and have now moved forward in the political process for scrutiny by the European Parliament. Once this process is complete, the new data requirements for pesticide active ingredients and formulated products will be adopted as separate regulations, repealing those referenced above. The anticipated amendments are substantially similar to those outlined above for		
negative effects on stakeholders			
	biocides, as are the benefits of regulatory alignment and drawbacks of		
	divergences.		
Possible solutions	40 CFR Part 158 was revised in 2007	and EPA has indicated that it	
	does not intend to embark on additional rulemaking in the foreseeable		
	future. However, another option could be the issuance of science-		
	policies that would authorize departur	res from Part 158 requirements	
	under well-defined circumstances.		

2. FOOD-USE PESTICIDES / PLANT PROTECTION PRODUCTS

	European Union	United States	
Agency / Directorate General	DG Enterprise and Industry	Environmental Protection	
		Agency	
Relevant regulatory / statutory provisions	Regulation (EC) No. 1272/2008	40 CFR § 156	
Regulatory differences & their negative effects on stakeholders	In 2008 the EU repealed Directives 67/548/EEC and 1999/45/EC and replaced them with the above-mentioned regulation with a view to implementing the United Nations Globally Harmonized System of Classification and Labeling of Chemicals (GHS) and facilitating worldwide trade through harmonized criteria for classification and labeling. More recently the US Occupational Safety and Health Agency has aligned its Hazard Communication Standard with the GHS; however, EPA's Office of Pesticide Programs, despite many years of discussion with stakeholders, has yet to take similar action in relation to its Label Review Manual and regulations.		
	Although classification and labeling (C&L) criteria are often said to be "test method neutral," meaning that the results of any internationally accepted test could in theory be used as a basis for C&L, some authorities and companies cite divergent C&L criteria as a basis for, e.g., not utilizing available 3R alternative methods. Practical examples of divergences between EU and US C&L criteria include:		
	 Acute oral/dermal toxicity limit and 5,000 mg/kg in the US. The limit dose on both practical and higher dose level is still retained divergence could lead EU comp 2000 mg/kg limit dose to repeat studies to satisfy US C&L requi A single "irritant" category for s vs. two irritant categories (sever accepted <i>in vitro</i> skin irritation to validated to distinguish between do not support sub-classification requirement for sub-classification been cited as the major barrier to non-animal test methods in the barrier 	dose of 2000 mg/kg in the EU e GHS discourages the higher animal welfare grounds, yet this d by EPA (but not OSHA). This panies with products tested at the cone or more acute toxicity irements. skin and eye irritation in the EU re / moderate) in the US. OECD- test methods are currently n irritants and non-irritants, but n. The US EPA (but not OSHA) on of skin and eye irritants has o the widespread use of these US.	
Possible solutions	EPA has indicated that it does not in rulemaking in the foreseeable future be the issuance of science-policies th from classification criteria specified	tend to embark on additional . However, another option could nat would authorize departures in § 156.	

3. CLASSIFICATION AND LABELLING OF CHEMICALS AND MIXTURES

We believe these points are responsive to the High-Level Regulatory Cooperation Forum's objectives of reducing excessive regulatory costs, unjustified regulatory differences, and unnecessary red tape while respecting each other's right to protect public health, safety, welfare and the environment.

Thank you for your consideration of this submission. Please direct any questions to the undersigned.

Sincerely, 6

Troy Seidle Director of Research & Toxicology +1 647 236 3889 (North America) +32 491 317 072 (Brussels) tseidle@hsi.org