November 8, 2018

Ian C. Read Chairman of the Board and CEO Pfizer, Inc. Via email: <u>Ian.read@pfizer.com</u>

Mikael Dolsten, M.D., Ph.D. President, Worldwide Research and Development Pfizer, Inc. Via email: <u>mikael.dolsten@pfizer.com</u>

Dear Chairman Read and Dr. Dolsten,

On behalf of People for the Ethical Treatment of Animals (PETA) and our more than 6.5 million members and supporters, I am writing to ask that Pfizer Inc. (Pfizer) discontinue use of the Forced Swim Test (FST) in its behavioral experiments involving animals.

Since 1989, Pfizer has published at least nine manuscripts that describe the use of the FST in experiments involving mice and rats. I have listed these references below. In publications, Pfizer-affiliated authors have described the FST as a model or test of depression,ⁱ "depression-like behavior,"ⁱⁱ and a test capable of demonstrating "antidepressant-like"ⁱⁱⁱ effects of compounds. However, the applicability of an animal's behavior during the FST to their mood, or to human depression, or to the utility of a compound for treating human depression has been substantially refuted. A thorough discussion of this matter is presented in the document, "The Invalidity of the Forced Swim Test" (attached).

In brief, animals, typically mice or rats, are made to swim in a cylinder of water. They swim frantically, trying to find an escape, until they stop struggling and subsequently float. The claim is that when mice spend more time floating, they are deemed to be more "depressed." This claim is made in spite of the evidence that floating is actually a learned and adaptive behavior, one that saves energy and is beneficial for survival.^{iv} Individual animals who are quicker to float also save energy and are less likely to sink, meaning that animals who more rapidly pick up on this reality, and spend less time struggling, are simply learning this adaptive behavior more readily.

Some claim that the forced swim test is a screening tool for antidepressant activity, since, sometimes, mice who are given drugs like fluoxetine will swim more and float less. However, the immobility response also occurs after treatment with drugs that do not have antidepressant effects at all, such as antihistamines and other miscellaneous drugs.^v Time spent swimming vs. floating is also influenced by the genetic strain of an animal and experimental variances, such as water depth or temperature.^{vi}



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Five^{vii} compounds identified in Pfizer's published animal experiments have been tested in humans. For only two^{viii} of these compounds did the authors' *interpretation* of an animal's behavior during the FST predict a *potential* efficacy or inefficacy of the compound's antidepressant-like action in humans; however, *none* of the compounds identified is currently approved as a treatment for human depression. For over half of the compounds identified, the authors' interpretation of what an animals' behavior during the FST means for humans, or the efficacy of the compound in human depression, was *not* corroborated in human trials. **This data suggest that, in your studies, a** *certain interpretation* of an animal's behavior during FST will predict the *potential* efficacy of a compound for use in human depression only 40 percent of the time, which is less than chance (50 percent), and has a zero percent chance of predicting the successful use of a compound for human depression.

There is a clear need to develop new therapeutics to treat human depression. Only small numbers of patients respond to available treatments, which themselves have severe shortcomings.^{ix} However, the use of animal experiments in an effort to generate these treatments has been criticized as a major contributor to failure rates in this area.^x Animal models of human depression lack many important aspects of model validity. Hendrie and Pickles argue that multiple failures on the part of animal experimenters are to blame for lack of progress in this field, namely falling trap to "logical flaws" and "false assumptions."^{xi}

The FST is so traumatic to animals that it is often used as a stressor in itself,^{xii} in an effort to create a sense of helplessness. To quote Dutch animal behaviorists Franz Josef van der Staay, Saskia S. Arndt, and Rebecca E. Nordquist, "If evidence accumulates that the intended goal/purpose cannot be reached, then one should consider abandoning further development of the model."^{xiii} This group also pointed out that in all cases, "benefits must outweigh the ethical costs of the animals. These costs include pain and suffering, distress and death."^{xiv}

In summary, the FST does not reliably predict successful treatments for human depression—nullifying any scientific justification for carrying out the test; and it causes acute suffering and distress to the animals who are used—presenting a compelling ethical argument against using the test. We therefore ask that Pfizer immediately discontinue its use of the FST in behavioral experiments involving animals.

May we meet to discuss this important matter?

Sincerely,

Egnel

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Use of the Forced Swim Test by Pfizer Inc.

- 1. Saccomano NA, Vinick FJ, Koe BK, Neilsen JA, et al. Calcium-independent phosphodiesterase inhibitors as putative antidepressants: [3-(bicycloalkyloxy)-4-methoxyphenyl]-2-imidazolidinones. *Journal of Medicinal Chemistry*. 1991;34:291-298.
- 2. Siuciak JA, Fujiwara RA. The activity of pramipexole in the mouse forced swim test is mediated by D2 rather than D3 receptors. *Psychopharmacology*. 2004;175:163-169.
- 3. Suiciak JA, McCarthy SA, Chapin DS, Fujiwara RA, et al. Genetic deletion of the striatum-enriched phosphodiesterase PDE10A: evidence for altered striatal function. *Neuropharmacology*. 2006;51:374-385.
- 4. Siuciak JA, Chapin DS, McCarthy SA, Guanowsky V, et al. CP-809,101, a selective 5-HT2C agonist, shows activity in animal models of antipsychotic activity. *Neuropharmacology*. 2007;52:279-290.
- 5. Siuciak JA, McCarthy SA, Chapin DS, Reed TM, Vorhees CV, Repaske DR. Behavioral and neurochemical characterization of mice deficient in the phosphodiesterase-1B (PDE1B) enzyme. *Neuropharmacology*. 2007;53:113-124.
- 6. Siuciak JA, McCarthy SA, Chapin DS, Martin AN. Behavioral and neurochemical characterization of mice deficient in the phosphodiesterase-4B (PDE4B) enzyme. *Psychopharmacology*. 2008;197:115-126.
- 7. Rollema H, Guanowsky V, Mineur YS, Shrikhande A, Coe JW, Seymour PA, Picciotto MR. Varenicline has antidepressant-like activity in the forced swim test and augments sertraline's effect. *European Journal of Pharmacology*. 2009;605(1-3):114-116.
- 8. Beyer CE, Dwyer JM, Piesla MJ, Platt BJ, et al. Depression-like phenotype following chronic CB1 receptor antagonism. *Neurobiology of Disease*. 2010;39;148-155.
- Grimwood S, Lu, Y, Schmidt AW, Vanase-Frawley MA, et al. Pharmacological characterization of 2methyl-N-((2'-(pyrrolidin-1-ylsulfonyl)biphenyl-4-yl)methyl)propan-1-amine (PF-04455242), a highaffinity antagonist selective for κ-opioid receptors. *The Journal of Pharmacology and Experimental Therapeutics*. 2011;339(2):555-566.

¹ Suiciak JA, McCarthy SA, Chapin DS, Fujiwara RA, et al. Genetic deletion of the striatum-enriched phosphodiesterase PDE10A: evidence for altered striatal function. *Neuropharmacology*. 2006;51:374-385.; Beyer CE, Dwyer JM, Piesla MJ, Platt BJ, et al. Depression-like phenotype following chronic CB1 receptor antagonism. *Neurobiology of Disease*. 2010;39;148-155.

ⁱⁱ Siuciak JA, McCarthy SA, Chapin DS, Reed TM, Vorhees CV, Repaske DR. Behavioral and neurochemical characterization of mice deficient in the phosphodiesterase-1B (PDE1B) enzyme. *Neuropharmacology*. 2007;53:113-124.; Siuciak JA, McCarthy SA, Chapin DS, Martin AN. Behavioral and neurochemical characterization of mice deficient in the phosphodiesterase-4B (PDE4B) enzyme. *Psychopharmacology*. 2008;197:115-126.

ⁱⁱⁱ Siuciak JA, Fujiwara RA. The activity of pramipexole in the mouse forced swim test is mediated by D2 rather than D3 receptors. *Psychopharmacology*. 2004;175:163-169.; Siuciak JA, Chapin DS, McCarthy SA, Guanowsky V, et al. CP-809,101, a selective 5-HT2C agonist, shows activity in animal models of antipsychotic activity. *Neuropharmacology*. 2007;52:279-290.; Rollema H, Guanowsky V, Mineur YS, Shrikhande A, Coe JW, Seymour PA, Picciotto MR. Varenicline has antidepressant-like activity in the forced swim test and augments sertraline's effect. *European Journal of Pharmacology*. 2009;605(1-3):114-116.; Grimwood S, Lu, Y, Schmidt AW, Vanase-Frawley MA, et al. Pharmacological characterization of 2-methyl-N-((2'-(pyrrolidin-1-ylsulfonyl)biphenyl-4-yl)methyl)propan-1-amine (PF-04455242), a high-affinity antagonist selective for κ-opioid receptors. *The Journal of Pharmacology and Experimental Therapeutics*. 2011;339(2):555-566.

^v Arai I, Tsuyuki Y, Shiomoto H, Satoh M, Otomo S. Decreased body temperature dependent appearance of behavioral despair in the forced swimming test in mice. *Pharmacological Research*. 2000;42:171-176.

^{vi} De Pablo JM, Parra A, Segovia S, Guillamon A. Learned immobility explains the behavior of rats in the forced swimming test. *Physiology and Behavior*. 1989;46:229-237.; Jeffrys D, Funder J. The effect of water temperature on immobility in the forced swimming test in rats. *European Journal of Pharmacology*. 1994;253:91-94.; Lucki I, Dalvi A, Mayorga AJ.

Sensitivity to the effects of pharmacologically selective antidepressants in different strains of mice. *Psychopharmacology* 2001;155:315-322.

vii Pramipexole; CP-809,101; varenicline; rimonabant; PF-04455242

^{ix} Hendrie C, Pickles A. The failure of the antidepressant drug discovery process is systemic. *Journal of Psychopharmacology*. 2013;27(5):407-416.

^x Garner JP. The significance of meaning: Why do over 90% of behavioral neuroscience results fail to translate to humans, and what can we do to fix it? *ILAR Journal*. 2014;55(3):438-456.; Hendrie 2013

xi Hendrie 2013

^{xii} de Kloet ER, Molendijk ML. Coping with the forced swim stressor: Towards understanding an adaptive mechanism. *Neural Plast.* 2016;2016:6503162.

^{xiii} van der Staay FJ, Arndt SS, Nordquist RE. Evaluation of animal models of neurobehavioral disorders. *Behavioral and Brain Functions*. 2009;5:11.

xiv van der Staay 2009

viii CP-809,101; rimonabant