Editorial
Whither the impending european regulation of presumed endocrine disruptors?

The legislative impulse to regulate presumed endocrine disrupting chemicals (EDCs) was born as an appendage to the US Food Quality Protection Act of 1996, focusing on public health rather than environmental issues. Pressed by advocacy claims, US legislators were persuaded by a study in animals - shortly after officially labeled as scientific misconduct1 - and by epidemiologic claims that could not be validated. Twenty years later, the momentum to regulate EDCs has spread worldwide, even though many studies over the last decades have yet to yield credible epidemiologic evidence of public health adversities linked to xenoendocrine contaminants. Absent a confirmed public health target, what could justify a program to regulate presumed EDCs?

The European Commission in June 2016 issued draft criteria for EDCs identification and regulation, still set in the conjectural frameworks of regulatory science and precautionary considerations.2 Animal and reductionist laboratory tests are to be used because tests in humans are not possible, and arbitrary definitions of adverse effects are to be adopted as valid clinical proxies for humans. As a novel challenge, the drafted criteria disregard potency in identifying EDC hazards, contrary to plain empirical evidence, common sense and elementary thermodynamics. Clearly, without sufficient causal potencies nothing stirs in the natural universe, including endocrine-dependent events.

A group of experts hosted by the German Federal Institute for Risk Assessment (BfR), included potency in the identification of EDC hazards and proposed that any substance could be considered an EDC, if acting by an endocrine mode of action (MoA) and causing adverse effects in a daily dose range up to 1000 mg/Kg bw.3 The equivalent upper dose would be 70g daily for a 70 Kg person. On these grounds, and invoking precaution, regulators likely would prescribe the highest doses as the standard for EDC testing, parallel to the prescribed maximum tolerated dose (MTD) for carcinogen bioassays. Such a testing regimen would likely indict numerous substances, even though typical xenoendocrines show receptor binding affinities thousands of times lower than human physiologic hormones, and are commonly experienced at very low concentrations. A regulatory scheme on these terms would considerably reduce vegetal food supplies by banning a large segment of staple foods that carry low levels of phytoestrogens. It would also ban many cosmetics, medicines and other compounds containing low levels of natural and synthetic hormones.

With these prospects, and conceding that regulation of putative EDCs may not be resisted, the adoption of pertinent World Health Organization (WHO/IPCS) testing guidelines has been considered.4 Excluding reductionist laboratory assays, the guidelines endorse tests in appropriate whole animal models to reach estimates of potencies and NOAELs against human hormone standards. On this evidence, it would be questionable to estimate human risks in the absence of clinical epidemiologic benchmarks, but it would be sensible to exonerate and remove from public anxieties those substances testing positive below appropriate thresholds of toxicologic or regulatory concern (TTC/TRC), at realistic exposures. Substances exceeding those thresholds would be regulated, although few of such instances could likely be found, due to the absence of clinical epidemiologic signals and the low concentrations and receptor binding affinities of putative xenoendocrines.

Opposing such options, a European EDC regulatory program disregardng potency would reinforce a dangerous precedent by further encouraging the creative regulation of putative hazards for public health adversities. The Commission did ask for public comments on the drafted criteria, but the first question is why the Commission chose to embark on this course. Is the Commission intrigued by the prospect of a new open season of authoritarian regulations justified by the flimsiest conjectures? Does the Commission hope for a flood of protests to counteract pressure, and thus to reinstate potency as a core justification of EDCs regulation?

Potency or no potency, however, EDC regulation in Europe — and similarly worldwide — is poised to continue in an autocratic rather than factual mode: it will be set by the same administrators writing rules, policing, prosecuting, judging, and penalizing. They also will appoint juries of advisors, selected by conflict of interest criteria

designed to preclude dissent. The arbitrary extent of such proceedings is generally unknown to the public and to most elected legislators, happy to believe that regulations are grounded on testable evidence. If sufficiently informed of this whimsical, authoritarian and legally enforced reality, free societies would be hard pressed to tolerate regulations that cause massive economic misallocations and pervasive public anxieties. More so if the public realized that even putative health and longevity benefits of EDC regulation could not be assessed, in the absence of testable clinical and epidemiologic benchmarks of public health adversities linkable to EDCs.

The social, civic and moral implications of such regulatory scenarios should advise restraint while raising some key questions. Remembering how the raison d’etre of regulation is to provide public health benefits, is it reasonable to issue costly regulations for the relief of putative public health problems bereft of clinical epidemiologic footprints? If a case is made for preventive and precautionary regulations, should those be justified by direct or indirect evidence that is factually measurable, or should they be triggered by whimsical conjectures? More pointedly, are certain regulations becoming a pretext for lucrative debates about angels on pin heads? What special interests stand to benefit from regulation, other than public health at large? Have biases and conflicts of interest of all participants – regulators included - been made transparent and openly perceived by the public?

Ethical and rational answers to these questions could put regulation on the right course, but it will not be. What inflames controversial regulations at the center of public, administrative and legislative attention is not a yearning for public health or a respect for science. Rather, it is the achieving of compromises to balance the open and ulterior interests of advocacies, academicians, regulators, advisors, industry, politicians and the media. Public health hardly emerges in those discussions.

Under irresistible pressure from advocates, the market, the media and an ill-informed European parliament, the European Commission is proposing to regulate EDCs based on criteria that cannot be justified scientifically or by common sense. They would enable an arbitrary approach to precaution, unconcerned that precaution itself becomes a very costly if not a paralyzing hazard, when its potency and dimensions were not accounted for. Indeed we seem to have reached a decision point: either we return to embrace the evidentiary ethic of science and a sensible approach to socially affordable precaution, or we proceed to slouch toward an “everything goes” regulatory morass dominated by opportunist special interests. For the moment, and unless improbable legislative miracles may happen, the balance rests in the hands of the European Commission.

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