

# SCIENTIFIC REPORTS



OPEN

## Defining an additivity framework for mixture research in inducible whole-cell biosensors

K. Martin-Betancor<sup>1</sup>, C. Ritz<sup>2</sup>, F. Fernández-Piñas<sup>1</sup>, F. Leganés<sup>1</sup> & I. Rodea-Palomares<sup>1</sup>

Received: 14 July 2015

Accepted: 27 October 2015

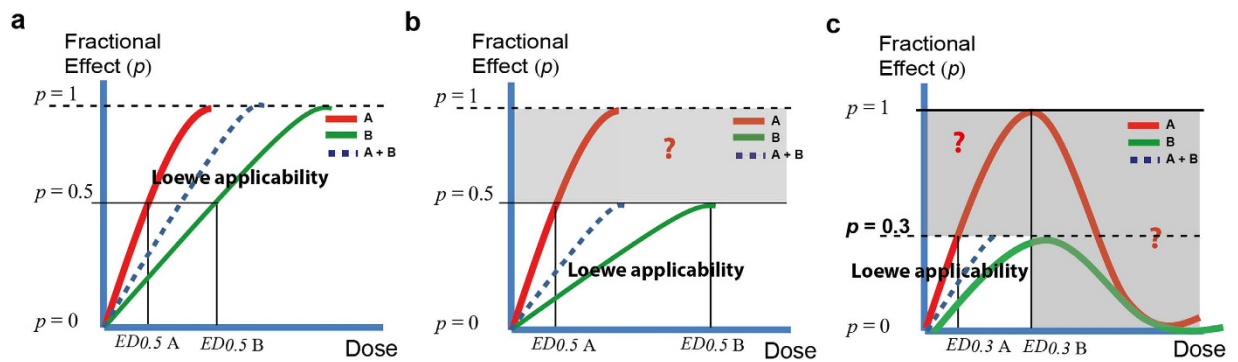
Published: 26 November 2015

A novel additivity framework for mixture effect modelling in the context of whole cell inducible biosensors has been mathematically developed and implemented in R. The proposed method is a multivariate extension of the *effective dose* ( $ED_p$ ) concept. Specifically, the extension accounts for differential maximal effects among analytes and response inhibition beyond the maximum permissive concentrations. This allows a multivariate extension of *Loewe additivity*, enabling direct application in a biphasic dose-response framework. The proposed additivity definition was validated, and its applicability illustrated by studying the response of the cyanobacterial biosensor *Synechococcus elongatus* PCC 7942 pBG2120 to binary mixtures of Zn, Cu, Cd, Ag, Co and Hg. The novel method allowed by the first time to model complete dose-response profiles of an inducible whole cell biosensor to mixtures. In addition, the approach also allowed identification and quantification of departures from additivity (interactions) among analytes. The biosensor was found to respond in a near additive way to heavy metal mixtures except when Hg, Co and Ag were present, in which case strong interactions occurred. The method is a useful contribution for the whole cell biosensors discipline and related areas allowing to perform appropriate assessment of mixture effects in non-monotonic dose-response frameworks

The prediction of the expected effect of chemical mixtures when only the effect of individual components is known is a hot topic in pharmacology, toxicology, and ecotoxicology<sup>1–3</sup>. A central element in mixture research is the definition and mathematical formulation of additivity<sup>2,4,5</sup>. At present, there are two sound pharmacological definitions of additivity: *Loewe additivity*<sup>6</sup> and *Bliss independence*<sup>7</sup>, which are the foundations of the so-called *Concentration addition* (CA) and *Independent Action* (IA) additivity models, respectively<sup>8</sup>. Departures from *Loewe additivity* can be quantitatively studied based on the *Combination Index* (CI)<sup>4,9</sup>. The crucial prerequisite for the applicability of any additivity model is to fulfill certain mathematical assumptions<sup>5</sup>. The basic mathematical feature of *Loewe additivity* is that the effects of the mixture components could be formulated in terms of a common *effective dose* ( $ED_p$ ). For instance, this requisite is trivially met when all individual mixture components present identical maximal effects, (see Fig. 1a). An important complication occurs with mixtures of chemicals that show differential maximal effects: When this happens, additivity hypothesis can only be formulated up to the effect levels achieved by the less potent compound present in the mixture due to inherent mathematical features of the additivity definition<sup>5</sup> (Fig. 1b). An additional challenge for formulating the additivity hypothesis occurs when the studied response is susceptible to result in non-monotonic biphasic dose-response patterns (see Fig. 1c). Besides, in biphasic dose-response curves, two different concentrations can result in the same *fractional effect* (Fig. 1c), resulting in misleading conclusions<sup>5</sup>. These problems have hampered the applicability of additivity models in important areas where differential maximal effects and biphasic dose-response patterns are commonly observed, such as in hormetic effects<sup>10</sup>, hormone agonists/antagonists research<sup>11</sup>,

<sup>1</sup>Departament of Biology, Facultad de Ciencias, Universidad Autónoma de Madrid, 28049 Madrid, Spain.

<sup>2</sup>Department of Nutrition, Exercise and Sports, Faculty of Science, University of Copenhagen, Rolighedsvej 30, DK-1958 Frederiksberg C, Denmark. Correspondence and requests for materials should be addressed to I.R.-P. (email: ismael.rodea@uam.es)



**Figure 1. Applicability of Loewe additivity.** Typical dose-response profiles for (a) classical monotonic dose-response curves for chemicals A and B showing identical maximal effects, (b) classical monotonic dose-response curves for chemicals A and B presenting differential maximal effects, and (c) biosensor type biphasic dose-response curves for chemicals A and B presenting differential maximal effects and toxicity threshold. The meanings of the terms presented in the figure can be found in theory section.

AhR agonists/antagonists activity research<sup>5</sup>, endocrine disruptors activity research<sup>12,13</sup>, and in general in the field of inducible (turn-on) whole cell biosensors.

To overcome these bottlenecks, some authors recently suggested the need for a formal mathematical expansion of the additivity formulations that may allow working with differential maximal effects<sup>14,15</sup>. Other authors have proposed a pragmatic numerical approximation based on a toxic unit extrapolation method to solve the problem<sup>5</sup>.

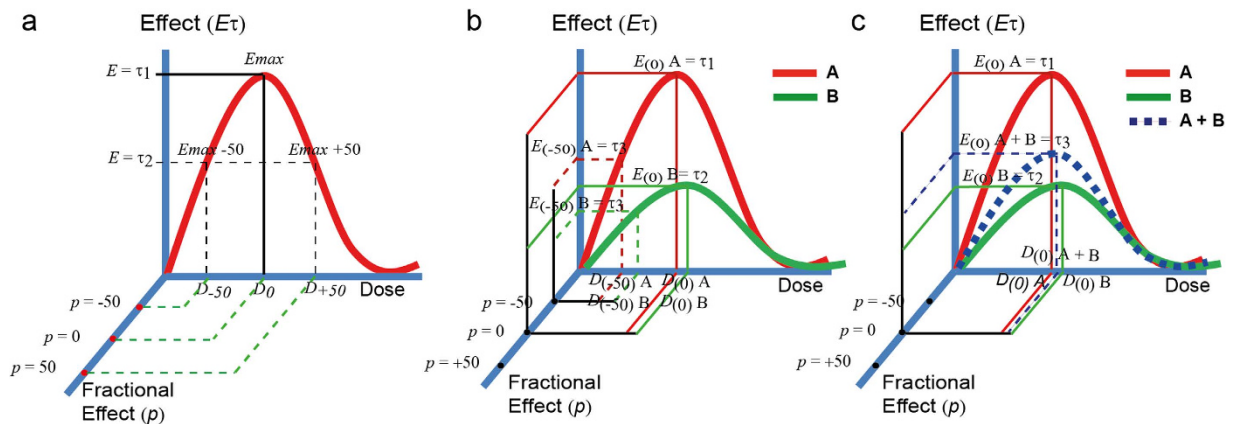
Inducible whole cell biosensors are a paradigmatic case of a biological system displaying differential maximal effects and usually biphasic dose-response profiles (e.g.,<sup>16</sup>) (Fig. 1c). Whole cell biosensors are intact, living cells genetically engineered to produce a dose-dependent measurable signal in response to a specific chemical or physical stimulus in their environment<sup>17</sup>. Inducible whole cell biosensors response is usually characterized by a dose-dependent biphasic profile presenting an induction region up to a concentration threshold (*maximum permissive concentration*), and a subsequent inhibition region where the biosensor response decays, possibly due to the inherent toxicity of the analyte above certain concentrations which affects cell viability<sup>18</sup>. Whole cell biosensors have been extensively used in the last 3 decades for the detection and quantification of different analytes and stresses of interest (for a review see<sup>17,19</sup>). Despite they have a clear vocation to be applicable in realistic conditions, mixture effect research using inducible whole-cell biosensors is presently a poorly developed research area. From our point of view, the main reason is the lack of a founded theoretical basis and experimental additivity framework.

In the present work, a novel additivity framework for mixture research in the context of whole cell inducible biosensors has been mathematically developed. The method proposes a multivariate extension of the *effective dose* ( $ED_p$ ) to take into account the occurrence of differential maximal effects and inhibition beyond the MPCs. In effect, this allows an extension of *Loewe additivity* that enables its direct application in a biphasic dose-response framework. A family of user friendly utilities has been incorporated in the (*drc*) package<sup>20</sup> for R. The method has been illustrated studying the response of the cyanobacterial biosensor *Synechococcus elongatus* PCC 7942 pBG2120 to binary mixtures of 6 heavy metals (Zn, Cu, Cd, Ag, Co and Hg). *Synechococcus* sp. PCC 7942 pBG2120 bears a fusion of the promoter region of the *smt* locus of *Synechococcus* sp. PCC 7942 to the *luxCDABE* operon of *Photobacterium luminescens*. It is an inducible self-luminescent *whole cell* biosensor able to respond to a broad range of heavy metal cations which present differential maximal effects and biphasic dose-response curves<sup>16</sup>. The method is a useful contribution for the entire whole-cell biosensors discipline and related areas which allows to perform sound mixture-effect research in the framework of biphasic dose-response curves.

## Theory

**A novel framework for mixture-effect research for whole cell biosensors.** We propose a novel framework for modelling mixture effects in *whole cell biosensors* showing biphasic dose-response curves. It is characterized by the following 5 steps: (1) Fitting biphasic dose-response profiles. (2) A dimensional extension of the *effective dose* notation. (3) *Two-dimensional formulation of Loewe additivity*. (4) Prediction of mixture effects based on individual component biphasic dose-response data. (5) Analysis of departures from additivity.

**Fitting biphasic dose-response profiles.** Inducible whole cell biosensors usually present inverted *v-shaped* biphasic dose-response profiles (Fig. 1c). This specific type of dose-response pattern may be fitted using nonlinear regression model equations Gaussian and LogGaussian. We considered 2 specific inverted *v-shaped* functions  $f$ : the *Gaussian* (Eq. 1 below) and the *log Gaussian* (Eq. 2 below) equations, which are defined as follows:



**Figure 2. The bi-dimensional fractional effect notation.** (a) Fractional effective doses  $ED_p$  in biphasic dose-response systems is proposed to be scaled as fractions ( $p$ ) of the  $E_{max}$  defining the fractional effect scale ( $E_p$ ). The empirical effect scale ( $E_{(\tau)}$ ), in the y axis is decoupled from the fractional effect scale ( $E_p$ ) which is projected in the z plane. This allows to define fractional effects covering the entire biphasic dose-response curve. (b) The proposed fractional effective notation allows to scale in a common fractional effect scale stimuli A and B showing differential maximal effects. (c) An additive biphasic dose response pattern “A + B” can be formulated for a theoretical mixture of A and B based on the univocal relationship of the fractional effect scale ( $E_p$ ) with the other two dimensions:  $D_{(p)}$  and  $E_{(p)}$ . For notation meanings in the Figures see section 2.

$$f(x) = c + (d - c) \cdot \exp(-0.5 \cdot \{(x - e)/b\}^f) \tag{1}$$

$$f(x) = c + (d - c) \cdot \exp(-0.5 \cdot |\log(x) - \log(e)|/b|^f) \tag{2}$$

where the parameters  $c$  and  $d$  correspond to the limits for  $x=0$  and  $x$  tending to infinity and the parameters  $b$  and  $e$  control the steepness of the curve and location of the peak, respectively. The parameter  $f$  describes asymmetry in the curve (i.e., asymmetry between the left and right sides of the peak).

**A multivariate extension of the effective dose notation.** In the present work, a notation for fractional effective doses  $ED_p$  scaled from the MPCs (hereafter,  $E_{max}$ ) has been chosen (Fig. 2a). The novelty of the approach is that we maintain the empirical effect scale ( $E_{(\tau)}$ ) in the y axis, and we project the fractional effect scale ( $E_p$ ) in the z plane (see Fig. 2a). Fractional effects ( $p$ ) in the  $E_p$  scale are defined as follows:  $-100 \leq p \leq 100$ . For  $p < 0$ , fractional effects on the left side of the  $E_{max}$  are obtained (Fig. 2a) (the induction part of the curve), for  $p > 0$ , fractional effects on the right side of the  $E_{max}$  are obtained (the inhibition part of the dose-response curve).  $p = 0 = E_{max} = MPC$ . Decoupling the fractional effect scale ( $E_p$ ) from the empirical effect ( $E_{(\tau)}$ ) allows to scale inverted  $v$ -shaped dose-response curves with differential maximum effects in a unique fractional scale independently of the maximum level of effect ( $E_{max}$ ) attained (Fig. 2b). However,  $ED_p$  needs to be dimensionally extended to account for differences in both the dose ( $D$ ) and effect ( $E$ ) scales (Fig. 2b). Therefore we define  $ED_p$  as a two-dimensional vector  $(D_{(p)}, E_{(p)})$  where  $D_{(p)}$  is the dose required to get the desired fractional effect ( $p$ ) (i.e. 50%), and  $E_{(p)}$  is the effect in the empirical effect scale ( $E(\tau)$ ) achieved at this fractional effect ( $p$ ).

**Two-dimensional formulation of Loewe additivity.** For classical monotonic dose-response curves, a uni-dimensional effective dose-notation and additivity formulation are enough to perform accurate additive predictions. In the same way that the fractional notation ( $ED_p$ ) needs to be extended to set a proper common fractional effect scale in biphasic dose-response curves, the existing additivity formulation needs to be extended as well. For this we propose a two-dimensional formulation of *Loewe additivity* computed for the two components of  $ED_p = (D_{(p)}, E_{(p)})_p$  (Fig. 2c). The additivity formulation on the dose ( $D$ ) scale is identical to the original *Loewe additivity* formulation. For notational convenience, we will formulate hereafter *Loewe additivity* as follows:

$$\frac{D_A}{(D_{(p)})_A} + \frac{D_B}{(D_{(p)})_B} = CI_{Dp} \tag{3}$$

where  $D_A$  and  $D_B$  are the doses of the components A and B which in combination produces a fractional effect  $p$  on the measured biological response.  $(D_{(p)})_A$  and  $(D_{(p)})_B$  are the doses that individually result in a fractional effect  $p$  for the components A and B, respectively. In case (Eq. 3) equals 1 *Loewe additivity* holds. Otherwise, Combination Index Theorem holds<sup>4</sup>. If  $CI_{Dp}$  denotes the Combination Index in the  $D$  dimension (see Fig. 2c) for the considered fractional effect  $p$ ,  $CI_{Dp} < 1$  indicates synergism,  $CI_{Dp} > 1$  indicates antagonism. To project *Loewe additivity* to the empirical effect dimension ( $E(\tau)$ ), we simply assume that the effect ( $E$ ) dimension is equivalent to the Dose ( $D$ ) dimension in its relationship with the fractional effect ( $Ep$ ) dimension (Fig. 2c). Therefore, it holds that *Loewe additivity* is computed in the  $E$  dimension as follows:

$$\frac{E_A}{(E_{(p)})_A} + \frac{E_B}{(E_{(p)})_B} = CI_{Ep} \quad (4)$$

where  $E_A$  and  $E_B$  are the effects (in the empirical scale ( $\tau$ )) of the components A and B which in combination results in a fractional effect  $p$  on the measured biological response.  $(E_{(p)})_A$  and  $(E_{(p)})_B$  are the effects in the empirical effect scale of the components A and B resulting individually in the desired fractional effect  $p$ .  $CI_{Dp}$  is the Combination Index in the  $E$  dimension for the considered fractional effect  $p$ .

The proposed two-dimensional formulations are susceptible of extension to  $n$  components as follows:

$$\sum_{i=1}^n \frac{D_i}{(D_{(p)})_i} = CI_{Dp} \quad (5)$$

$$\sum_{i=1}^n \frac{E_i}{(E_{(p)})_i} = CI_{Ep} \quad (6)$$

where  $D_i$  is the dose of the  $i^{\text{th}}$  component which in combination produces a fractional effect  $p$ .  $(D_{(p)})_i$  is the dose of the  $i^{\text{th}}$  components that individually produces the same fractional effect  $p$ .  $E_i$  is the effect ( $\tau$ ) of the  $i^{\text{th}}$  component which in combination produces a fractional effect  $p$ .  $(E_{(p)})_i$  is the effect ( $\tau$ ) of the  $i^{\text{th}}$  components that individually produce the same fractional effect  $p$ . Thus,  $CI_{Dp}$  and  $CI_{Ep}$  are the combination index scores for the  $D$  and  $E$  dimensions at any fractional effect  $p$ .

### Predictions of the joint effect of a mixture under two-dimensional Loewe additivity.

Equation (5) for  $CI_{Dp} = 1$  can be rewritten in a predictive formulation (allowing for *in-silico* predictions of the joint effect of  $n$  mixture components based on individual chemical information only) according to Faust, *et al.*<sup>21</sup>, if  $D_i$  are expressed as relative proportions  $j_i$  of the total dose ( $D_{mix}$ ), where  $j_i = D_i/D_{mix}$ . Under the condition that the mixture elicits a  $p$  total fractional effect, the total dose of the mixture  $D_{mix}$  is defined as the effective dose of the mixture ( $D_{pmix}$ ) required to produce a fractional effects  $p$ . If  $CI = 1$ ,  $\Rightarrow D_i$  can be substituted in Eq. (5) by  $j_i D_{pmix}$ , and by rearrangement, it holds:

$$D_{pmix} = \left( \sum_{i=1}^n \frac{j_i}{D_{pi}} \right)^{-1} \quad (7)$$

According to Gonzalez-Pleiter, *et al.*<sup>22</sup>, equation (5) can be similarly rewritten as follows when  $CI \neq 1$ :

$$D_{pmix} = \left( \sum_{i=1}^n \frac{j_i}{D_{pi} \cdot CI_{Dp}} \right)^{-1} \quad (8)$$

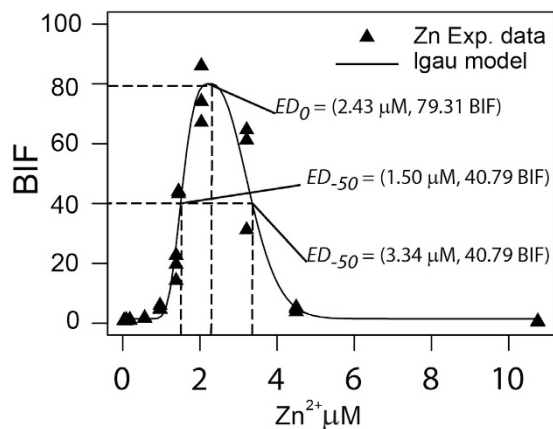
Considering the proposed two-dimensional notation  $ED_p = (D_{(p)}, D_{(p)})$ , the formulation of Equation (7) in the  $E$  dimension gives:

$$E_{(p)mix} = \left( \sum_{i=1}^n \frac{j_i}{E_{(p)i}} \right)^{-1} \quad (9)$$

In case  $CI \neq 1$ :

$$E_{(p)mix} = \left( \sum_{i=1}^n \frac{j_i}{E_{(p)i} \cdot CI_{Ep}} \right)^{-1} \quad (10)$$

**Analysis of departures from additivity.** The definition in equations (5) and (6) also allows investigating departures from additivity in biphasic dose-response systems. The result from evaluating



**Figure 3. Fitting biphasic profiles with non-linear regression models.** Experimental response of *Synechococcus elongatus* PCC 7942 pBG2120 to increasing concentrations of  $\text{Zn}^{2+}$ . Best fit model for  $\text{Zn}^{2+}$  logGaussian “lgau2” (see Supplementary Material SM 1) is presented in the figure. Position of selected  $ED_p$  vectors ( $-50, 0, +50$ ) are marked in the figure.

equations (5) and (6) are two numerical values of the Combination Index ( $CI$ ) for the  $D$  and  $E$  dimensions, which can be expressed as a two-dimensional vector  $CI_{p(D,E)} = (CI_{Dp}, CI_{Ep})$ . This approach yields 9 theoretical combinations of  $CI_{p(D,E)}$  as follows:  $CI_{p(D,E)} = (1, 1)$ ;  $(1, <1)$ ;  $(1, >1)$ ;  $(<1, 1)$ ;  $(<1, <1)$ ;  $(<1, >1)$ ;  $(>1, >1)$ ;  $(>1, 1)$ ;  $(>1, <1)$ .

For global assessment of departures from additivity, a simplification of the two-dimensional information can be obtained by calculating a weighted index  $CI_{wp}$  for any fractional effect  $p$  as follows:

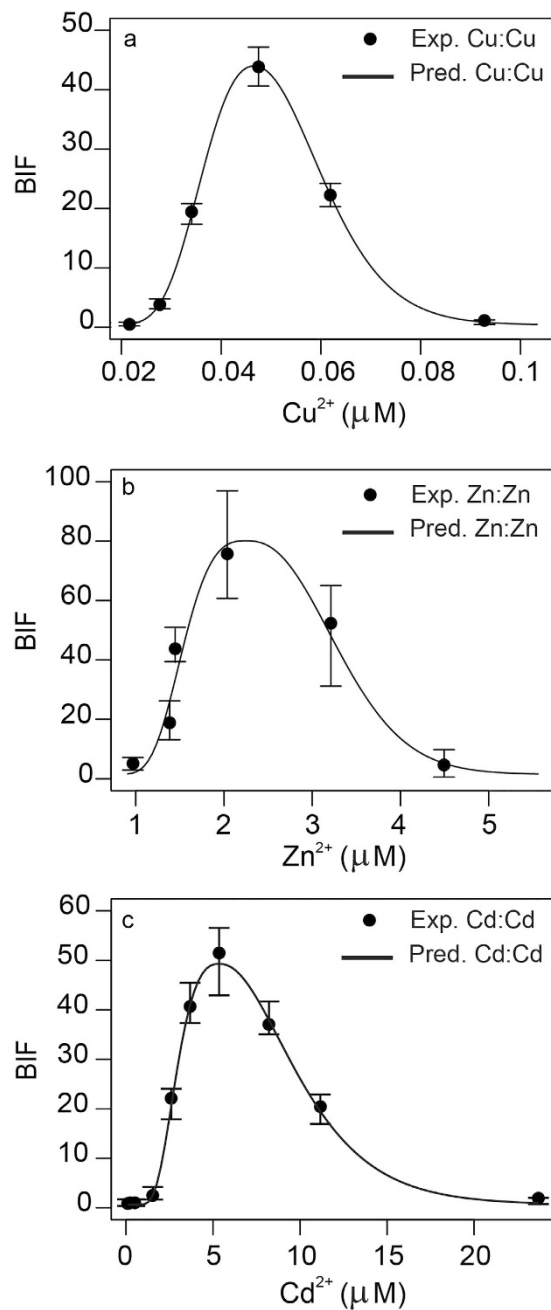
$$CI_{wp} = CI_{Dp} \cdot CI_{Ep} \quad (11)$$

where  $CI_{wp} > 1$  indicates overall antagonism,  $CI_{wp} = 1$  an overall additive effect, and  $CI_{wp} < 1$ , an overall synergistic effect.

## Results

Dose-response profiles of *Synechococcus elongatus* PCC 7942 pBG2120 exposed to Zn, Cu, Cd, Ag, Co and Hg and their binary combinations were fitted using the proposed set of  $v$ -shaped non-linear functions described in Theory section. Best fit models for each metal and metal mixture were selected accordingly to the minimum of the residual sum of squares<sup>20</sup>. Supplementary Material SM1 shows a summary of the selected model function and parameters for each individual metal and metal mixture. Overall, the majority of single metal experimental data were better fitted by log Gaussian function, and only Ag was fitted better by a Gaussian model. On the contrary, the responses of the biosensor to half of the binary metal mixtures were fitted better by Gaussian, and the other half by logGaussian model (see Supplementary Material SM1). Figure 3 shows as an example, the goodness of fit of the  $v$ -shaped non-linear function logGaussian “lgau2” to Zn response experimental data (see Supplementary Material SM1 for model parameters). As can be observed, the function fits adequately Zn experimental data at both the induction and the inhibition part of the dose-effect curve. Using the *drc* package we can get  $ED_p$  vectors ( $D_{(p)}, E_{(p)}$ ) at any desired fractional effect  $p$ . For example, we can calculate the Zn  $ED_{-50} = (1.50 \mu\text{M}, 40.79 \text{ BIF})$ , Zn  $ED_0 = (2.43 \mu\text{M}, 79.31 \text{ BIF})$  and the Zn  $ED_{+50}$  to be  $(3.34 \mu\text{M}, 40.79 \text{ BIF})$  (Fig. 3) (definitions in Methods section). Since  $ED_p$  is two-dimensional, it includes not only the concentration required to produce a specific fractional effect  $p$ , but also the actual BIF that the biosensor signal achieves at this specific fractional effect for a specific analyte. A summary of  $ED_p$  vectors ( $-50, 0, +50$ ) for the 6 metals can be found in Supplementary Material SM2. As can be seen, the bi-dimensional notation allows to explicitly predict concentrations of the analyte resulting in fractional effects in both the induction and the inhibition regions of the dose-response curves. An illustrative example on how to get  $ED_p$  vectors can be found in Supplementary Material SM3.

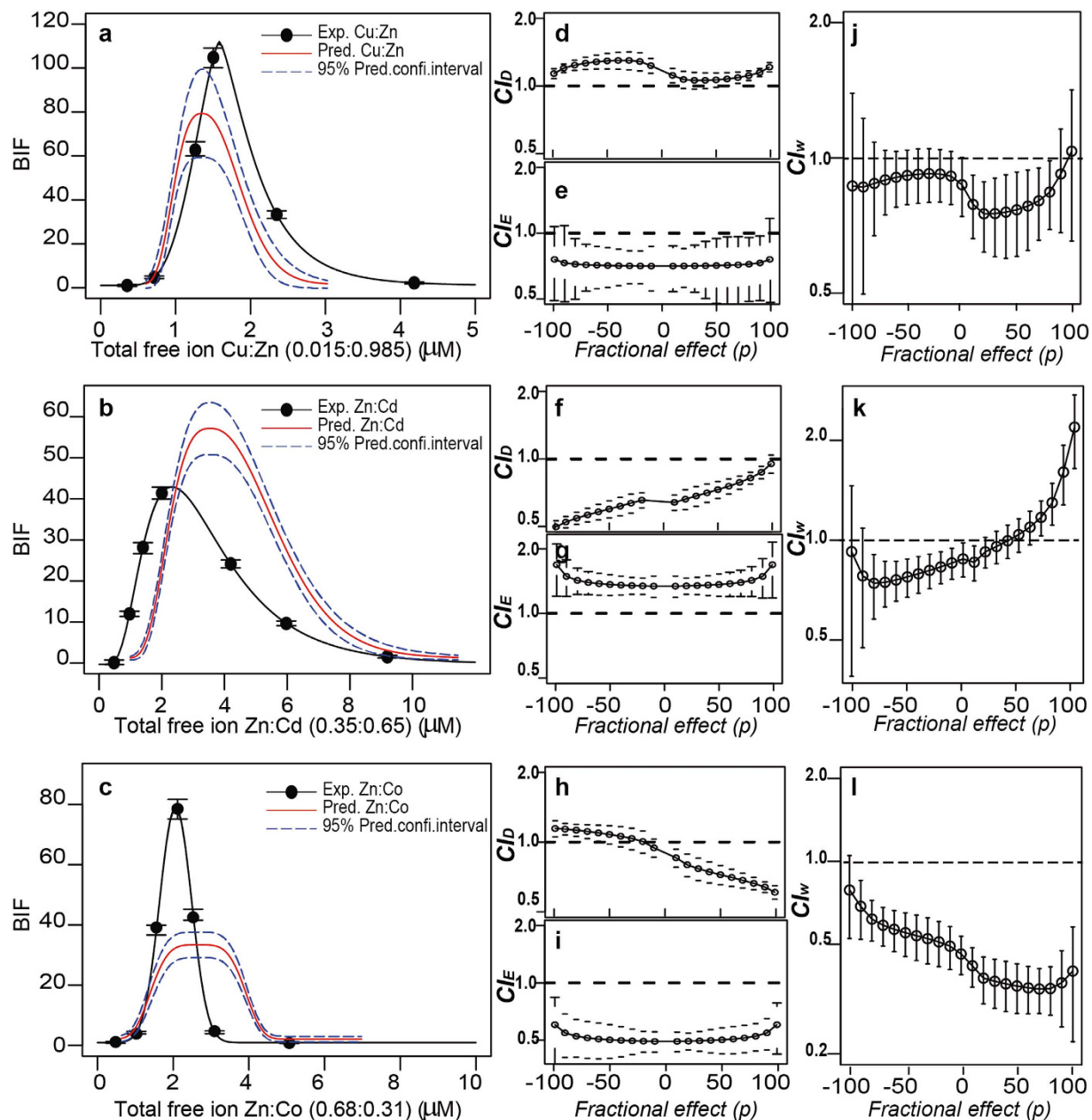
Once individual metal biphasic response profiles are adequately fitted, and  $ED_p$  vectors can be obtained to any fractional effect  $p$ , it is possible to predict the biosensor response to any combination of the individual metals evaluating Equations (7) and (9). However, prior to any further analysis, it is required to validate the proposed two-dimensional definition of *Loewe additivity*. Figure 4 shows predicted vs experimental dose-response patterns of sham binary mixtures for Zn:Zn, Cd:Cd and Cu:Cu. A near perfect overlap occurred between the experimental and predicted dose-response curves of sham mixtures for the 3 metals. This validates the proposed multivariate formulation of *Loewe additivity* model for biphasic dose-response curves, even at mixture concentrations where inhibition of the signal occurs (See Fig. 1b).



**Figure 4. Validation of the additivity definition using sham mixtures.** Predicted dose-response patterns of sham binary mixtures for (a) Cu:Cu, (b) Zn:Zn, (c) Cd:Cd. Predicted dose-response patterns were calculated solving the multivariate extension of *Loewe additivity* equation in predictive formulation (eqs (7) and (9)). Error bars are standard errors ( $n = 3-4$ ).

Once the multivariate extension of *Loewe additivity* was validated with the sham mixtures, we applied the method to perform additivity predictions and to study the nature of the interaction (if any) of the response of *Synechococcus elongatus* PCC 7942 pBG2120 to the 15 possible binary mixtures of Zn, Cd, Cu, Ag, Hg and Co. Complete results of the analysis can be found in Supplementary Material SM5. A representative selection of the results is presented in Fig. 5. In the case of the binary mixture Cu:Zn (Fig. 5a), the additivity prediction fitted reasonably well the experimental *dose-response* profile. However, the additivity predictions of the binary mixtures Zn:Cd (Fig. 5b) and Zn:Co (Fig. 5c), deviated from their respective experimental *dose-response* pattern, suggesting a departure from additivity. However, which kind of departure are we observing?

To answer this question, a quantitative analysis of the nature of the interactions is required<sup>3,4</sup>. Departures from additivity can be quantified solving Equations 5 and 6 for any fractional effect  $p$ . As illustrative example, we used *extended p-CI* plots to graphically present the results of the quantification



**Figure 5. Analysis of departures from additivity.** Experimental vs predicted (under additivity) dose-response patterns for Cu:Zn (a), Zn:Cd (b) and Zn:Co (c) binary mixture's, respectively. Extended  $p$ -CI plots presenting departures from additivity (as CI values) as a function of the effect level ( $p$ ) for the  $D$  dimension of Cu:Zn (d), Zn:Cd (f) and Zn:Co (h). Extended  $p$ -CI plots presenting departures from additivity (as CI values) as a function of the effect level ( $p$ ) for the  $E$  dimension of Cu:Zn (e), Zn:Cd (g) and Zn:Co (i). Extended  $p$ - $CI_w$  plots presenting weighted departures from additivity (as  $CI_w$  values) as a function of the effect level ( $p$ ) for Cu:Zn (j), Zn:Cd (k) and Zn:Co (l). Error bars are standard errors ( $n = 3-4$ ). Total free ion concentrations presented in the Figures are those of Supplementary Material SM2 presented as  $ED_0$  in Supplementary Material SM2 corrected by MINTEQA2 calculations due to the presence of the two metals used in each mixture.

of the interactions of the three selected mixtures (Cu:Zn, Zn:Cd and Zn:Co). Extended  $p$ -CI plots allow to observe the nature of the interactions along the  $D$  and  $E$  dimensions covering the complete biphasic dose-response curves, representing the deviation from the additivity line ( $CI = 1$ ) for any fractional effect  $p$ . Figures 5d,f,h and Figure 5e,g,i show extended  $p$ -CI plots for the  $D$  and  $E$  dimensions (respectively), for the three selected metal binary mixtures (Cu:Zn, Zn:Cd and Zn:Co). Extended  $p$ -CI plots for  $D$  and  $E$  axis (Fig. 5d,e, respectively) for the Cu:Zn combination showed CI values near 1 (the additivity line)

along the entire range of fractional effects ( $p$ ) in both dimensions, confirming the near additive behaviour of this metal combination. The Zn:Cu binary combination presented statistically significant synergism ( $CI < 1, p < 0.05$ ) along the entire range of fractional effects ( $p$ ) in the  $D$  dimension (Fig. 5f), but statistically significant antagonism ( $CI > 1, p < 0.05$ ) in the  $E$  dimension (Fig. 5g). On the other hand, the Zn:Co combination showed a  $p$ -dependent interaction patterns in the dose ( $D$ ) axis (going from antagonistic to synergistic), and consistent synergism ( $CI < 1, p < 0.05$ ) in the effect ( $E$ ) axis along the entire range of effect levels ( $p$ ) (Fig. 5h,i). See Supplementary Material SM4 for specific  $p$ -values for three selected fractional effect levels ( $-50, 0, +50$ ). Figure 5j–l showed *Extended  $p$ - $CI_w$  plots* for the mixtures Cu:Zn, Zn:Cu and Zn:Co, respectively. *Extended  $p$ - $CI_w$  plots* (see equation 11) can be used as a measure of the overall fitness to additivity of the response of a biosensor to mixtures of analytes. As can be seen in the Fig. 5j, the overall effect of the combination of Cu:Zn combination was additive, that of Zn:Cu combination was  $p$ -dependent (Fig. 5k), going from synergistic (below the MPC) to antagonistic (above the MPC). That of Zn:Co was consistently synergistic  $CI < 0.5$  along the entire range of effect levels ( $p$ ) (Fig. 5l).

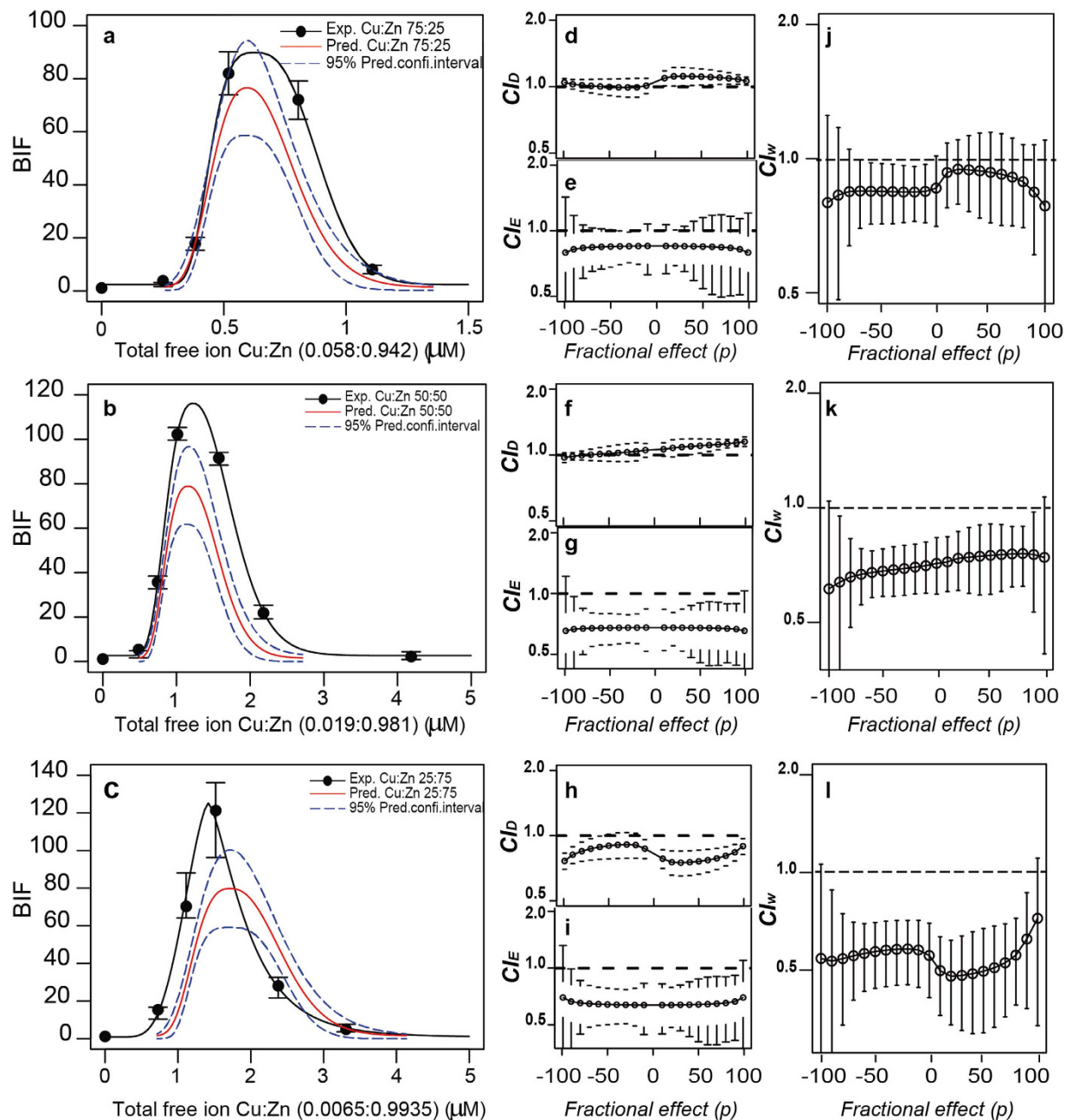
The effect of the metal ratio on the predictive power of the multivariate extension of Loewe additivity was addressed for selected binary mixtures. Figure 6a–c shows the experimental dose-response patterns and the respective additivity predictions for the three different metal ratios (75:25, 50:50, 25:75, respectively) of the binary metal mixture Cu:Zn. *Extended  $p$ - $CI$  plots* for the  $D$  and  $E$  dimensions for the different metal ratios are presented in Fig. 6d–i. In addition, *extended  $p$ - $CI_w$  plots* are presented in Fig. 6j–l. The selection of metal ratios allows to specifically address the possibility of predicting the dose response patterns including metal combinations in which eventually one of the metal may be present below the MPC and the other above the MPC and vice versa. As can be seen in Fig. 6, the main features of the dose-response pattern of the 50:50 mixture of Cu:Zn (Fig. 6b), that is additivity in  $D$ , synergism in  $E$ , and overall additive effect ( $CI_w > 0.5$ ) is essentially conserved in the 25:75 and the 75:25 ratio. The only differences were the occurrence of a slight tendency to synergism in  $D$  in the 25:75 ratio (Fig. 6h), and a slight tendency to synergism in both the 75:25 and 25:75 ratios based on  $CI_w$  (but still additive based on the management criterion:  $0.5 < CI_w < 2$ ). Similar results were obtained for ratio variations for the mixture Zn:Cu (Supplementary Material SM4).

The analysis of the departures from additivity for the 15 possible binary combinations of the studied metals revealed a complex scenario where the 9 possible theoretical combinations of the  $CI_{D,E}$  vectors anticipated in Theory section were actually found (Supplementary Material SM5). In order to get a global idea on the fitness to additivity of the response of the biosensor to the binary mixtures of metals, we computed  $CI_w$  values according to Equation (11) which were summarized as well as *polygonograms*<sup>4</sup> (for  $p$  levels  $-50, 0, +50$ ) in Fig. 7. Interestingly, additive or near additive effects (according to the management criterion:  $0.5 < CI_w < 2$ ) hold for binary mixtures of Zn, Cu, Ag and Cu at the three representative  $p$  levels (Fig. 7). However, some mixtures containing Hg, Co and Ag resulted in significant departures from additivity: The mixtures Hg:Co and Hg:Ag resulted in synergism and antagonism, respectively at the three representative  $p$  levels ( $-50, 0, +50$ ). In addition, some mixtures resulted in effect-level dependent departures from additivity: the mixture Co:Zn resulted in synergism at  $p = 0$  and  $p = +50$ , and the mixture Ag:Cu resulted in synergism at  $p = +50$ .

## Discussion

Here we present a theoretical framework which allows to perform sound mixture-effect research in inducible whole cell biosensors and related fields. To have a mathematical formulation of additivity is crucial in order to obtain accurate and comprehensive results in mixture research as demonstrated in the last 20 years in pharmacology and eco/toxicology<sup>2–4</sup>. In these disciplines, *Loewe additivity* is the gold standard for additivity formulations<sup>2–4</sup>. However, the practical applicability of *Loewe additivity* is historically hampered in biological systems presenting differential maximal effects and non-monotonic responses such as biphasic dose-response curves<sup>5,14</sup>. In the present work we propose a multivariate extension of *Loewe additivity* which allows its application in the context of differential maximal effects and biphasic dose-response curves. The proposed methodology was validated and tested using the inducible whole-cell self-luminescent metal biosensor *Synechococcus elongatus* PCC 7942 pBG2120 as case study. Our solution is in agreement with the conceptual formulations which Belz, *et al.*<sup>14</sup> proposed in order to extend the applicability of *Loewe additivity* in the context of hormesis (an stimulatory effect found at low doses of toxicants). They postulated that differential maximal hormetic effects among individual mixture components force the need to perform additivity formulations in both dose ( $D$ ) and effect ( $E$ ) dimensions independently. As solution, they proposed to use the original *Loewe* equation (Equ. 3) for predictions in the ( $D$ ) dimension, but a simple summation of fractional effects for predictions in the  $E$  dimension. Their predictions were reasonable for the dose ( $D$ ) dimension, but were, in their own words “more dubious” for the ( $E$ ) dimension<sup>14</sup>. This is because their proposal of *Loewe additivity* formulation for the  $E$  dimension was not accurate. We have demonstrated with the sham mixtures that our definition (Eqs 4 and 6) is a true projection of *Loewe additivity* in the ( $E$ ) dimension. Scholze, *et al.*<sup>5</sup> recently proposed an elegant approach to the differential maximal effect problem. Since, without multivariate extension, it is impossible to solve *Loewe additivity* equation for systems presenting differential maximal effects, they proposed a numerical approximation as a solution. Basically, they constructed an extrapolated interval of mixture predicted effects based on reasonable maximal and minimal hypothetical contributions of the less potent mixture components. The method worked well with a biosensor based

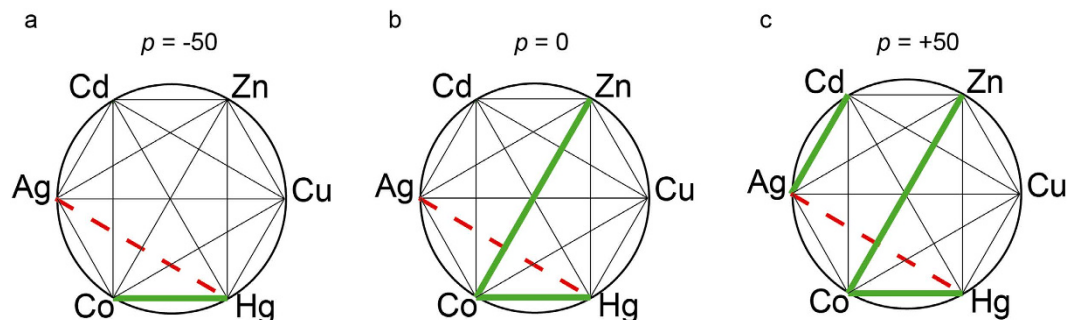




**Figure 6. Effect of mixture ratio.** Experimental vs predicted (under additivity) dose-response patterns for the mixture Cu:Zn at 75:25 (a), 50:50 (b) and 25:75 (c) ratios based on the  $D_0$  concentration of the metals. Extended  $p$ -CI plots presenting departures from additivity (as CI values) as a function of the effect level ( $p$ ) for the  $D$  dimension (d,f,h), and the  $E$  dimension (e,g,i). Extended  $p$ - $CI_w$  plots presenting weighted departures from additivity (as  $CI_w$  values) as a function of the effect level ( $p$ ) for the binary mixture Cu:Zn at 75:25 (j), 50:50 (k), 25:75 (l) mixture ratios. Error bars are standard errors ( $n=3-4$ ). Total free ion concentrations presented in the Figures are those presented as  $ED_0$  in Supplementary Material SM2 corrected by MINTEQ calculations due to the presence of the two metals used in each mixture.

on partial agonists of aryl hydrocarbon receptor (AhR) used for mixtures including components with varying maximal effects. However, they recognized that it cannot work in systems presenting inhibitory thresholds beyond the MPCs<sup>5</sup>.

The availability of a *Loewe additivity* formulation conceived for inducible whole-cell biosensors is an important milestone in this field of research which may allow for a wider generalization of mixture-effect research. This is especially true if a user friendly utility is available. We have made available all the presented mathematical equations, statistical and graphical utilities in the “dose-response curve” (*drc*) package for R<sup>20</sup>. Currently the methodology is presently set up for binary mixtures, and only two biphasic



**Figure 7.  $CI_w$  Polygonograms.** Polygonograms summarizing weighted combination index ( $CI_w$ ) for all the binary combinations of the 6 studied metal cations (Zn, Cu, Cd, Ag, Co and Hg) at three representative fractional effects ( $p$ ):  $-50$  (a),  $0$  (b),  $+50$  (c). Departures from additivity are based on the risk management criterion (see Methods). Thin solid black lines represent additive effect ( $0.5 < CI_w < 2$ ). Solid green lines indicate synergism ( $CI_w < 0.5$ ). Dashed red lines indicate antagonism ( $CI_w > 2$ ).

models (Gaussian and logGaussian) are available. However, future work will focus on extending the framework to  $n$ -component mixtures and more biphasic models.

An important application of the methodology is the possibility, by the first time, of predicting inducible whole cell biosensor responses to mixtures using single chemical experimental information only. As shown by the experiments with varying mixture ratios, this was true even when individual chemicals were present above or below the MPCs. This equals the potential applicability of inducible whole cell biosensors in mixture risk assessment with the same conditions and scope than those of monotonic toxicity tests, or *turn-off* whole cell biosensors such as Microtox<sup>23,24</sup>. This increases the opportunities of inducible whole cell biosensors to be included in future regulatory frameworks which will explicitly consider mixture risk assessment<sup>1,25</sup>. A complementary application of the methodology is the possibility of investigating departures from additivity (synergism and antagonism)<sup>3,4</sup>. In the presented case study with *Synechococcus elongatus* PCC 7942 pBG2120, we found that the response to the binary combinations of the different metals was quite complex when analyzing the two constitutive dimensions ( $D$  and  $E$ ), ranging from additivity to synergism and antagonism. In addition, we found that additivity, synergism or antagonism or any of their permutations, even of opposite effects occurred, for example a combination of synergism in ( $D$ ), and antagonism in ( $E$ ). In fact, the occurrence of opposite behaviours in the ( $D$ ) and ( $E$ ) dimensions was relatively common. One may wonder what the biological meaning of these patterns is. Most possibly, the answer will depend on the biological receptor used for the biosensor construction. In our case, the observed opposite behaviors in ( $D$ ) and ( $E$ ) dimensions for some metal mixtures may reflect the counteracting effects of the metals on the two independent but co-existing biological processes involved in the bioluminescence signal emission: On one hand, a synergistic effect of the metals on the induction of the *smtAB* promoter may result in lower metal concentration required to start to induce the system (synergism in  $D$ ). However, a synergism in the toxicity of the metals affecting cell metabolism may result in a lower FMNH<sub>2</sub> and ATP pool, inhibiting the bioluminescence signal (antagonism in  $E$ ). Similar arguments may hold for the different possible combinations of departures from additivity in the two dimensions for other biosensor systems. However, from a practical point of view, one may be more interested in finding out whether or not an overall deviation from additivity in the biosensor response may likely occur with a specific combination of analytes, rather than in a detailed analysis of deviations in the  $D$  and  $E$  dimensions and their causes. In such case,  $CI_w = CI_D \cdot CI_E$ , can be used as a measure of the overall fitness to additivity of the response of a biosensor to mixtures of analytes. Intuitively,  $CI_w$  will tend to zero when  $CI_D$  and  $CI_E$  have opposite values, reflecting the counteracting effects of the tendencies in both dimensions. In the other hand,  $CI_w$  would be magnified (either in synergism or antagonism) when both  $CI_D$  and  $CI_E$  show the same kind of interaction. For example, if  $CI_D = 0.5$  and  $CI_E = 0.5$ ,  $CI_w = 0.25$ , this results in increased synergism in the overall response of the biosensor (lower concentration and higher induction than expected to reach the same  $p$  level). Therefore,  $CI_w$  will allow to easily detect those metal combinations which may result in evident deviations from additivity which may potentially be problematic from a practical view-point. The analysis of  $CI_w$  of the biosensor response for the 15 binary metal combinations revealed that the weighted response of *Synechococcus elongatus* PCC 7942 pBG2120 could be considered nearly additive for the mixtures including Cu, Cd, Zn. However, important departures from additivity occurred when Hg, Co and to a lesser extent Ag were present in the mixtures. It is interesting to note that Hg and Co are the weakest inducers of the *smtAB* promoter alone<sup>16</sup>, however their impact on the analytical performance of the biosensor would be important. A similar analysis can be applied to any other biosensor strain whose robustness to interaction effects with detectable analytes would be addressed. Despite in the present work we have not performed any correction of the biosensor signal with a cell viability control which may serve to increase the dynamic range or bioanalytical performance of a whole-cell biosensor<sup>17,18</sup>, the presented analytical method (additivity formulations and

analysis of departures from additivity) is equally applicable when the biosensor signal is corrected by such viability control. Finally, since our solution ultimately relies on a multivariate extension of the effective dose notation, the method may be generalized to other additivity formulations, such as *Bliss independence*<sup>7</sup>. *Bliss independence* may be a functional additivity formulation for stimuli with dissimilar mode of action but whose effects converge in the activation of a common biological effect<sup>26</sup>. Similarly, the bi-dimensional *Loewe additivity* may be generalized to any number of  $k$  extra dimensions which may be required to describe any biological response. For example, one important dimension, which may be approached with the same multivariate extension, is the exposure time. It may be done defining a three-dimensional *Loewe additivity* formulation, and an extension of the effective dose notation in the form  $ED_p = (D_{(p)}, E_{(p)}, t_{(p)})$ . This method may have important applications in the formulation of additivity hypothesis in studies which may explicitly consider toxicokinetics and toxicodynamics, such as transcriptomics<sup>27</sup>.

## Conclusions

A multivariate extension of the *effective dose* ( $ED_p$ ) notation in order to take into account the occurrence of differential maximal effects and signal inhibition beyond the MPCs in biphasic dose-response curves has been developed. This allows a multivariate extension of *Loewe additivity* enabling its direct application in a biphasic dose-response framework. The proposed additivity definition has been experimentally validated using sham mixtures, finding excellent agreement between experimental and predicted dose-response patterns. The method was applied to study the response of the cyanobacterial self-luminescent metallothionein-based whole-cell biosensor *Synechococcus elongatus* PCC 7942 pBG2120 to binary mixtures of 6 heavy metals (Zn, Cu, Cd, Ag, Co and Hg). The response of *Synechococcus* to the mixtures can be considered nearly additive except when Hg, Co and to lesser extent Ag were present in the mixtures which resulted in important departures from additivity. The method has different applications and is a useful contribution for the entire whole-cell biosensors field and related areas allowing to perform sound mixture research in non-monotonic dose-response frameworks.

## Material and Methods

**Chemicals.** Chemicals were of analytical grade, culture media and heavy metals stock solutions were prepared in MilliQ water. Heavy metal salts  $ZnCl_2$ ,  $CdCl_2$ ,  $AgSO_4$ ,  $CuSO_4$ ,  $HgCl_2$ ,  $CoCl_2$ ,  $PbNO_3$ ,  $MgCl_2$ ,  $NiCl_2$ ,  $FeCl_2$ ,  $BaCl_2$  and  $SrCl_2$  were from Sigma-Aldrich (Germany). Concentrated metal salts solutions (1000 mg/L) were prepared in deionized water (Millipore) and stored at 4 °C in opaque bottles. Dilutions and mixtures were freshly prepared in MilliQ water before the experiments.

**Heavy metal mixture exposure experiments.** Culture conditions and heavy metal exposure of self-luminescent *Synechococcus elongatus* PCC 7942 pBG2120 were as previously described<sup>16</sup>. Briefly, exposure was performed in transparent 24 well microtiter plates in 1.5 mL final volume BG11 medium (without Co, Ni, Cu, Zn). Plates were incubated at 28 °C in the light, Ca. 40  $\mu\text{mol photons m}^{-2} \text{s}^{-1}$  during 4 h. Luminescence measurements were performed in a Centro LB 960 luminometer in opaque white 96 well microtiter plates. The biosensor signal was expressed as luminescence induction factor (BIF) as previously reported<sup>16,28</sup>.

Mixture exposure design was adapted from<sup>3,29</sup>. Basically, *Synechococcus elongatus* PCC 7942 pBG2120 was exposed to Zn, Cu, Cd, Ag, Co and Hg, and to their 15 possible binary combinations. Binary mixtures were prepared according to a constant ratio design (1:1) based on the individual  $D_{-50}$  (the dose required to get half of the MPC of each individual metal). 7 to 9 serial dilutions (factor 2) of each individual metal and binary combination were tested at the same time<sup>4</sup>. At least three independent experiments were performed.

**Experimental validation of the additivity definition by sham mixtures.** Validation of the additivity definition was performed according to the *sham* mixture procedure<sup>30</sup>. *sham* mixtures are *false* mixtures in which the two components are exactly the same substance, and therefore, their fractional combination should be perfect additivity<sup>30</sup>. If an additivity formulation is correct, experimental and predicted dose-response pattern of the *sham* mixtures should perfectly overlap. *sham* mixtures of Cu:Cu, Zn:Zn and Cd:Cd (ratio 1:1) were prepared and tested as described in theory.

**Analysis of results.** *Fitting biphasic dose-response profiles and obtaining  $ED_p$  vectors.* The entire dose-response profiles of the response of *Synechococcus elongatus* PCC 7942 pBG2120 to the different individual heavy metal cations (Zn, Cu, Cd, Ag, Co and Hg) and their binary combinations were fitted using the non-linear functions described in Theory, with and without the use of variance-stabilizing Box-Cox transform-both-sides approach<sup>20</sup>. Best fit models were selected based on the minimum of the residual sum of squares<sup>20</sup>. *Effective doses* ( $ED_p$ ) = ( $D_{(p)}, E_{(p)}$ ) for the different metal cations and combinations were calculated from the fitted *v-shaped* nonlinear functions.

*Additivity predictions and departures from additivity.* Additive *dose-effect* profiles for the different heavy metal combinations were predicted by solving equation (7) and (9). The molar fractions ( $j_i$ ) of

the mixture components were fixed according to the experimental mixture metal fractions described in Theory. Two-dimensional Combination Index  $CI_{(D, Ep)} = (CI_{Dp}, CI_{Ep})$  and weighted  $CI_{wp}$  values were computed accordingly to equations 5, 6 and 11, respectively. An illustrative example of the whole procedure of fitting dose-response curves, additivity formulation and analysis of departures from additivity can be found in Supplementary Material SM3.

**Criteria for quantification of departures from additivity.** In the present work, two criteria for the definition of departures from additivity are considered. The “pharmacological criterion” considers departures from additivity when  $H_0: CI = 1$ , is rejected (two-tailed Student’s t-Test), and is equivalent to the criterion used in human pharmacology<sup>4</sup>. The second criterion considered is the “risk management criterion (singular)” which defines departure from additivity as significant based on pragmatic thresholds:  $Exp < 0.5 \cdot CA$  (synergism) and  $Exp > 2 \cdot CA$  (antagonism)<sup>31,32</sup>. The equivalent thresholds based on CI are:  $0.5 < CI > 2$ . Departures from additivity for  $CI_D$  and  $CI_E$  were analyzed based on the pharmacological criterion, while analysis of departures from additivity in  $CI_w$  were based on the risk management (for consistency) criterion.

## References

- Altenburger, R. *et al.* Future water quality monitoring—adapting tools to deal with mixtures of pollutants in water resource management. *Sci Total Environ* **512–513**, 540–551, doi: 10.1016/j.scitotenv.2014.12.057 (2015).
- Kortenkamp, A., Backhaus, T. & Faust, M. State of the art report on mixture toxicity. Final Report to the European Commission under Contract Number 070307/2007/485103/ETU/D.1., (European Commission, Brussels, Belgium, 2009).
- Rodea-Palomares, I. *et al.* Application of the combination index (CI)-isobologram equation to study the toxicological interactions of lipid regulators in two aquatic bioluminescent organisms. *Water Res* **44**, 427–438, doi: 10.1016/j.watres.2009.07.026 (2010).
- Chou, T. C. Theoretical basis, experimental design, and computerized simulation of synergism and antagonism in drug combination studies. *Pharmacol Rev* **58**, 621–681, doi: 10.1124/pr.58.3.10 (2006).
- Scholze, M., Silva, E. & Kortenkamp, A. Extending the applicability of the dose addition model to the assessment of chemical mixtures of partial agonists by using a novel toxic unit extrapolation method. *PLoS one* **9**, e88808, doi: 10.1371/journal.pone.0088808 (2014).
- Loewe, S. The problem of synergism and antagonism of combined drugs. *Arzneimittelforschung* **3**, 285–290 (1953).
- Bliss, C. I. The toxicity of poisons applied jointly. *Ann Appl Biol* **26**, 585–615, doi: 10.1111/j.1744-7348.1939.tb06990.x (1939).
- Greco, W. R., Bravo, G. & Parsons, J. C. The search for synergy: a critical review from a response surface perspective. *Pharmacol Rev* **47**, 331–385 (1995).
- Chou, T. C. & Talalay, P. Quantitative analysis of dose-effect relationships: the combined effects of multiple drugs or enzyme inhibitors. *Adv Enzyme Regul* **22**, 27–55 (1984).
- Calabrese, E. J. Hormesis: why it is important to toxicology and toxicologists. *Environ Toxicol Chem* **27**, 1451–1474, doi: 10.1897/07-541 (2008).
- Hecker, M. *et al.* Human adrenocarcinoma (H295R) cells for rapid *in vitro* determination of effects on steroidogenesis: hormone production. *Toxicol Appl Pharmacol* **217**, 114–124, doi: 10.1016/j.taap.2006.07.007 (2006).
- McMahon, T. A. *et al.* The fungicide chlorothalonil is nonlinearly associated with corticosterone levels, immunity, and mortality in amphibians. *Environ Health Perspect* **119**, 1098–1103, doi: 10.1289/ehp.1002956 (2011).
- Silva, E., Scholze, M. & Kortenkamp, A. Activity of xenoestrogens at nanomolar concentrations in the e-screen assay. *Environ Health Perspect* **115**, 91–97, doi: 10.1289/ehp.9363 (2007).
- Belz, R. G., Cedergreen, N. & Sorensen, H. Hormesis in mixtures — Can it be predicted? *Sci Total Environ* **404**, 77–87, doi: http://dx.doi.org/10.1016/j.scitotenv.2008.06.008 (2008).
- Ohlsson, A., Cedergreen, N., Oskarsson, A. & Ulleras, E. Mixture effects of imidazole fungicides on cortisol and aldosterone secretion in human adrenocortical H295R cells. *Toxicology* **275**, 21–28, doi: 10.1016/j.tox.2010.05.013 (2010).
- Martin-Betancor, K., Rodea-Palomares, I., Munoz-Martin, M. A., Leganes, F. & Fernandez-Pinas, F. Construction of a self-luminescent cyanobacterial bioreporter that detects a broad range of bioavailable heavy metals in aquatic environments. *Front Microbiol* **6**, 186, doi: 10.3389/fmicb.2015.00186 (2015).
- van der Meer, J. R. & Belkin, S. Where microbiology meets microengineering: design and applications of reporter bacteria. *Nat Rev Microbiol* **8**, 511–522, doi: 10.1038/nrmicro2392 (2010).
- Roda, A. *et al.* Analytical strategies for improving the robustness and reproducibility of bioluminescent microbial bioreporters. *Anal Bioanal Chem* **401**, 201–211, doi: 10.1007/s00216-011-5091-3 (2011).
- Yagi, K. Applications of whole-cell bacterial sensors in biotechnology and environmental science. *Appl Microbiol Biotechnol* **73**, 1251–1258, doi: 10.1007/s00253-006-0718-6 (2007).
- Ritz, C. & Streibig, J. C. Bioassay Analysis using R. *J Stat Softw* **12**, 17 (2005).
- Faust, M. *et al.* Predicting the joint algal toxicity of multi-component s-triazine mixtures at low-effect concentrations of individual toxicants. *Aquat Toxicol* **56**, 13–32, doi: http://dx.doi.org/10.1016/S0166-445X(01)00187-4 (2001).
- Gonzalez-Pleiter, M. *et al.* Toxicity of five antibiotics and their mixtures towards photosynthetic aquatic organisms: implications for environmental risk assessment. *Water Res* **47**, 2050–2064, doi: 10.1016/j.watres.2013.01.020 (2013).
- Fernandez-Pinas, F., Rodea-Palomares, I., Leganes, F., Gonzalez-Pleiter, M. & Angeles Munoz-Martin, M. Evaluation of the ecotoxicity of pollutants with bioluminescent microorganisms. *Adv Biochem Eng Biotechnol* **145**, 65–135, doi: 10.1007/978-3-662-43619-6\_3 (2014).
- Wernersson, A.-S. *et al.* The European technical report on aquatic effect-based monitoring tools under the water framework directive. *Env Sci Eur* **27**, 7 (2015).
- Brack, W. *et al.* The SOLUTIONS project: challenges and responses for present and future emerging pollutants in land and water resources management. *Sci Total Environ* **503–504**, 22–31, doi: 10.1016/j.scitotenv.2014.05.143 (2015).
- Junghans, M., Backhaus, T., Faust, M., Scholze, M. & Grimme, L. H. Application and validation of approaches for the predictive hazard assessment of realistic pesticide mixtures. *Aquat Toxicol* **76**, 93–110, doi: http://dx.doi.org/10.1016/j.aquatox.2005.10.001 (2006).
- Altenburger, R., Scholz, S., Schmitt-Jansen, M., Busch, W. & Escher, B. I. Mixture Toxicity Revisited from a Toxicogenomic Perspective. *Environ Sci Technol* **46**, 2508–2522, doi: 10.1021/es2038036 (2012).

28. Jouanneau, S., Durand, M. J., Courcoux, P., Blusseau, T. & Thouand, G. Improvement of the identification of four heavy metals in environmental samples by using predictive decision tree models coupled with a set of five bioluminescent bacteria. *Environ Sci Technol* **45**, 2925–2931, doi: 10.1021/es1031757 (2011).
29. Rodea-Palomares, I. F.-P., F. González-García, C. & Leganes, F. in *Nova Science Publishers, Inc. New York. USA.* (2009).
30. Dawson, D. A., Allen, J. L., Schultz, T. W. & Pösch, G. Time-dependence in mixture toxicity with soft-electrophiles: 2. Effects of relative reactivity level on time-dependent toxicity and combined effects for selected Michael acceptors. *J Environ Sci Health A* **43**, 43–52, doi: 10.1080/10934520701750371 (2007).
31. Altenburger, R., Backhaus, T., Boedeker, W., Faust, M. & Scholze, M. Simplifying complexity: Mixture toxicity assessment in the last 20 years. *Environ Toxicol Chem* **32**, 1685–1687, doi: 10.1002/etc.2294 (2013).
32. Cedergreen, N. Quantifying Synergy: A Systematic Review of Mixture Toxicity Studies within Environmental Toxicology. *PLoS one* **9**, e96580, doi: 10.1371/journal.pone.0096580 (2014).

## Acknowledgements

This research was supported by MINECO grants CGL2010-15675 and CTM2013-45775-C2-2-R.

## Author Contributions

All authors have made essential contributions to this study. Experimental design: I.R., K.M., F.L. and F.F.P. Programation of R scripts C.R., I.R. and K.M. Experimental execution: K.M., F.L. and I.R. Data analysis: K.M. and I.R. Manuscript drafting: I.R., F.F.P. and C.R. All authors critically read and approved the final manuscript.

## Additional Information

**Supplementary information** accompanies this paper at <http://www.nature.com/srep>

**Accession codes:** Data used in the present manuscript is publicly available at: Doi: <http://dx.doi.org/10.6084/m9.figshare.1476176>

**Competing financial interests:** The authors declare no competing financial interests.

**How to cite this article:** Martin-Betancor, K. *et al.* Defining an additivity framework for mixture research in inducible whole-cell biosensors. *Sci. Rep.* **5**, 17200; doi: 10.1038/srep17200 (2015).



This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>