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Mycotoxins co-contamination: Methodological aspects and biological relevance of combined toxicity studies.

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Abstract

Mycotoxins are secondary fungal metabolites produced mainly by *Aspergillus*, *Penicillium* and *Fusarium*. As evidenced by large-scale surveys, humans and animals are simultaneously exposed to several mycotoxins. Simultaneous exposure could result in synergistic, additive or

antagonistic effects. However, most toxicity studies addressed the effects of mycotoxins separately.

We present the experimental designs and we discuss the conclusions drawn from *in vitro* experiments exploring toxicological interactions of mycotoxins.

We report more than 80 publications related to mycotoxin interactions. The studies explored combinations involving the regulated groups of mycotoxins, especially aflatoxins, ochratoxins, fumonisins, zearalenone and trichothecenes, but also the “emerging” mycotoxins beauvericin and enniatins. Over 50 publications are based on the arithmetic model of additivity. Few studies used the factorial designs or the theoretical biology-based models of additivity. The latter approaches are gaining increased attention. These analyses allow determination of the type of interaction and, optionally, its magnitude. The type of interaction reported for mycotoxin combinations depended on several factors, in particular cell models and the tested dose ranges. However, synergy among *Fusarium* toxins was highlighted in several studies. This review indicates that well-addressed *in vitro* studies remain valuable tools for the screening of interactive potential in mycotoxin mixtures.

Introduction

Mycotoxins are secondary fungal metabolites produced under specific environmental conditions by a variety of molds, mainly *Aspergillus*, *Penicillium* and *Fusarium spp.* As secondary metabolites, they are not essential for life, but may provide the fungus with an ecological advantage in certain environments. Some 300 compounds have been recognized as mycotoxins of which around thirty are considered as threat to human or animal health. Mycotoxin exposure via food and feed may result in many different adverse health effects such as carcinogenicity, immunotoxicity, reproductive toxicity, hepatotoxicity, nephrotoxicity, *etc.* (Bennett and Klich, 2003). Global surveys indicate that more than 70% of the samples of feed and feed raw materials are positive for at least one mycotoxin (Streit *et al.*, 2013a).

Human and animals are simultaneously exposed to several mycotoxins (Schothorst and van Egmond, 2004; Rodrigues and Naehrer, 2012; Streit *et al.*, 2013b); thus, there is a need for an update of the traditional single mycotoxin risk assessment approach (SCF, 2002). Indeed, in the field of toxicological evaluation of chemical mixtures, the consensus is that the customary chemical-by-chemical approach to risk assessment is in danger of underestimating the risk of chemicals to health (Kortenkamp *et al.*, 2009). Simultaneous exposure to different toxins could result in antagonistic, additive or synergistic effects. Although the demonstration of synergism would heighten concerns about health risks, the implications of additive combination effects have not received adequate attention. Sometimes the threshold dose for toxic effects may be exceeded in case of exposure to a mixture although the exposure to each single compound is unlikely to pose risk (Silva *et al.*, 2002). Therefore, an increasing number of mycotoxin studies

are devoted to their combined toxicity, especially to the exploration of the type of toxicological interactions.

The toxicity of a mixture is complex. The general principles for such analyses have been thoroughly reviewed elsewhere (ATSDR, 2004; Binderup, 2008). Testing for a possible interaction in mixture toxicity requires a comparison of the actual experimentally determined effects of the mixture with the theoretically expected no interaction effects. This prediction of no interaction, the null hypothesis, is done based on the toxicity of the individual compounds. Stronger-than-expected effects indicate synergism whereas lower-than-expected effects indicate antagonism. Several methods have been proposed but a generally agreed definition of zero interaction does not yet exist (Groten *et al.*, 2001). In this review we present the experimental designs and statistical aspects as well as the main conclusions drawn from experiments exploring interactions in combined toxicity of mycotoxins.

1. The reality of mycotoxins co-contamination

The reality of mycotoxins co-contamination is confirmed on the one hand by the co-occurrence of these toxins in food and feed stuff and on the other hand by co-exposure monitoring survey.

The co-occurrence of mycotoxins in food and feed is explained by three different reasons: (i) most fungi are able to simultaneously produce several mycotoxins, (ii) commodities can be contaminated by several fungi simultaneously or in quick succession, and (iii) the complete diet comprised different commodities. The recent application of LC-MS-MS based multi-mycotoxin methods revealed that food- and feed commodities almost always contain a cocktail of different

mycotoxins. In practice, the co-occurrence of mycotoxins represents the rule and not the exception.

In a three-year monitoring (2009 - 2011) on the worldwide occurrence of mycotoxins in feedstuffs and feed, Rodrigues and Naehrer (2012) showed that 48% of 7049 analyzed samples sourced in the Americas, Europe and Asia were contaminated with two or more of the tested mycotoxins (aflatoxins, zearalenone, deoxynivalenol, fumonisins and ochratoxin A). A literature review of European multi-mycotoxin contamination studies indicated that 75% to 100% of animal feed samples to contain more than one mycotoxin (Streit *et al.*, 2012), while the co-occurrence of more than two mycotoxins was reported in 95% of Spanish barley samples (Ibanez-Vea *et al.*, 2012). Analyzing 83 samples of maize, wheat, barley and silage from Europe, America and Australia by a multi-mycotoxin HPLC-MS/MS approach Streit *et al.* (2013) have shown that all the samples were co-contaminated by 7 to 69 mycotoxins or other potentially toxic secondary metabolites, mainly produced by the *Fusarium* genus. Moreover, combination of *Aspergillus* and *Fusarium* mycotoxins can be found in the same matrix. The co-occurrence aflatoxin B₁ (AFB₁), ochratoxin A (OTA) and deoxynivalenol (DON) accounted for 55% of the multi-contaminated Spanish barley samples (Ibanez-Vea *et al.*, 2012). The fact that unlike other foodborne toxins or microorganisms, most mycotoxins are resistant to milling, processing and heat treatments, increases the risk of their persistence in the food and feed chains and may participate to the co-contamination (Milicevic *et al.*, 2010, Streit *et al.*, 2012).

These trends depicted by food and feed monitoring for mycotoxins co-contamination are corroborated by exposure data collected in several human bio-monitoring studies. Simultaneous measurement of multiple mycotoxins using advanced LC-MS/MS technique for human exposure

assessment surveys in Germany, southern Italy and central Africa showed that 52% to 100% of urine samples contained biomarkers for two or more mycotoxins, and up to five mycotoxins were detected in a severe case of co-exposure (Abia *et al.*, 2013; Gerding *et al.*, 2014; Solfrizzo *et al.*, 2014). Moreover, as shown in Table 1, the exposure data highlight clearly that any kind of combinations involving mycotoxins irrespective of their producing fungi and their known geographical distribution could threaten consumer's health.

2. Experimental designs and statistical aspects to assess mycotoxin toxicological interactions

Interactions are inferred when a mixture of chemicals produces a biological response greater or lower than expected. Thus, the key question remains what is to be expected from a combination of contaminants. The application of Loewe's additivity equation or of Bliss' independence criterion, based on the dose-response curves of single compounds, enables the simulation of a theoretical response that represents the expected behavior of the mixture when interaction is excluded. Some reviews have been written about the subject and several aspects of the problem are still debated, with a particular regard to the biological plausibility of these two different theoretical approaches (Greco *et al.*, 1995; Chou, 2006; Goldoni and Johansson, 2007).

Classically, a two-step approach is recommended when analyzing the pharmacological or toxicological interactions between the different compounds of a mixture (Suhnel, 1996). First, the expected effects of the combination for the case of no interaction have to be predicted. This means a clear statement of what effect size can be expected if the compounds in the mixture do not interact. Then, the data on the effects of the experimental combination have to be compared

to the expected ones in order to classify the combination as additive (no interaction *i.e.* as expected), synergistic (*i.e.* interaction resulting in greater effect than expected) or antagonistic (*i.e.* interaction leading to lesser effect than expected). Several experimental designs that denote authors' point of view on this null case have been used in *in vitro* assessment of the combined effects of mycotoxins.

2.1. The arithmetic definition of additivity

In a number of studies, the expected mixture effect size was defined as equal to the arithmetic sum of the sizes of the effects for individual compounds when tested separately (Weber *et al.*, 2005; Kouadio *et al.*, 2007; Ribeiro *et al.*, 2010; Ficheux *et al.*, 2012; Klaric *et al.*, 2012). As an example for the null case, the expected size for the cytotoxic effect of a mixture could be defined as the sum of the cytotoxic effects induced by each mycotoxin alone in mono-exposure experiments, so:

$$\text{Cytotoxic effect (mycotoxin 1 + mycotoxin 2)} = \text{Cytotoxic effect (mycotoxin 1)} + \text{Cytotoxic effect (mycotoxin 2)}$$

When the measured cytotoxicity values are not significantly different or above or below the expected values the results are interpreted as additive, synergistic or antagonistic respectively.

Although intuitively plausible and very easy to handle, most researchers in the biomedical area seem to agree that combined effects do not simply equal the sum of single effects (Boedeker and Backhaus, 2010). The fallacy of this approach is better perceived when applying it to the combined effect of several doses of the same mycotoxin which by definition cannot behave

synergistically, nor antagonistically. As an illustration, dose-response experiments on the cytotoxic effects of DON and fumonisin B₁ (FB₁) were conducted in a study on the *in vitro* myelotoxicity induced by mixtures of *Fusarium* mycotoxins on human hematopoietic progenitors (Ficheux *et al.*, 2012). The authors concluded to an antagonistic effect of both *Fusarium* mycotoxins as the measured values were significantly lower than expected values. If we consider that the 2 μ M FB₁ dose can be seen as a 1+1 μ M FB₁ application, the predicted value for 2 μ M FB₁ using the arithmetic approach would be 60 \pm 8% cell viability, lower than the measured value (42 \pm 5%). Therefore the arithmetic sum model to which so many mycotoxin interaction studies referred to, does not provide a reasonable reference point.

2.2. Factorial design experiments

Factorial design experiments have been employed to assess interaction between mycotoxins (Tajima *et al.*, 2002; Heussner *et al.*, 2006; Lei *et al.*, 2013; Wan *et al.*, 2013a; Wan *et al.*, 2013b). When testing the effects of mixtures of varying combinations and the effects of each individual compound, the effect of any compound could be predicted by subtracting the mean of the groups not containing the compound from the mean of the other groups containing the compound (Groten *et al.*, 1996). In a full factorial design, each chemical in the mixture is studied at all dose levels of the other chemicals. This may require a large number of tested groups and can be very costly. Mycotoxin mixture studies favored the fractional factorials that enable more economy of experimentation because only part of the full factorial is run experimentally. A three-step study was proposed to detect interactions between five *Fusarium* mycotoxins inhibiting DNA synthesis *in vitro*. In stage 1 the combined action (additivity, or departure from

additivity) was assessed for the entire mixture, but not for specific pairs of mycotoxins. Stage 2 was specifically meant to economically screen for significant departure from additivity of specific (pairs of) mycotoxins using central composite designs, which allowed to finally apply full factorial design only to two-factor (two mycotoxins) interactions of particular interest (Groten *et al.*, 1998).

Despite the fact that interaction is definitely revealed by such statistical methods, the nature of interaction with regard to additivity, synergism or antagonism is not clearly explored and has to be inferred indirectly (Bhat and Ahangar, 2007). Wan *et al.* (2013a) applied a factorial design approach to elucidate the interactions in the combined cytotoxic effects of DON, nivalenol (NIV), zearalenone (ZEA) and FB₁ in swine jejunal epithelial cells. They first conducted dose-response experiments for each mycotoxin individually to select the range limits for subsequent interaction analysis. Then, a central composite design including a fractional factorial part was applied with four factors, i.e. DON, NIV, ZEA and FB₁, in order to minimize the number of possible toxin combinations (44 possible combinations of every concentration of each toxin). Nonetheless, 16 more data-points (in addition to the individual dose-response experiments) were required for interaction analysis. Univariate analysis of variance conducted on such data revealed non-additive interactions in all mixtures except DON-ZEA-FB₁, though the type of non-additive interactions (synergy or antagonism) still remained to be established. The factorial design approach could also just point out “a potential for interactive (synergistic) effects of citrinin and ochratoxin A and possibly other mycotoxins in cells of renal origin” (Heussner *et al.*, 2006).

2.3. *The theoretical biology models-based definitions of additivity*

The most commonly used theoretical biology models-based definitions of zero interaction are Bliss' independent criterion also known as response addition, Loewe's additivity model also named concentration or dose addition (Goldoni and Johansson, 2007) and the median effect principle of the mass action law (Chou, 2006).

Bliss' independence criterion and Loewe's additivity model

The main assumption for Bliss' independent criterion is that the chemical agents act independently from one another. In other words, the mode and possibly the site of action of the compounds in the mixture differ. When no interaction occurs for a combination, the Bliss independent criterion for two toxic agents can be expressed by the following equation:

$$E(x, y) = E(x) + E(y) - E(x) * E(y)$$

where E is the fractional effect (between 0 and 1), and x and y are the doses of two compounds in a combination experiment.

Loewe's additivity model relies on the assumption that the toxic agents in the mixture of concern act on the same biological sites, by the same mechanisms of action and differ only in their potency. Relatively simple Loewe's additivity model extensions are the isobolographic method and its algebraic variant, the interaction index, particularly useful when assessing two toxic substances *in vitro*. The interaction index can be expressed as (Berenbaum, 1981):

$$I = c_1 / EC_{x,1} + c_2 / EC_{x,2}$$

with c_i denoting the applied concentrations (of agent 1 and 2, respectively) and EC_x their individual concentrations that provoke a certain effect x , *e.g.* the effect concentration 50% (EC_{50}). $I < 1$, $I > 1$ and $I = 1$ mean the agents interact synergistically, antagonistically, or are additive. This index has been applied to mycotoxin pairs association (McKean *et al.*, 2006a; McKean *et al.*, 2006b).

Some papers simultaneously tested both Loewe's additivity and Bliss' independence criterion models of zero interaction for mycotoxins combined effects because there is no final agreement on the biological plausibility of these concepts (Tammer *et al.*, 2007; Mueller *et al.*, 2013). Both Bliss' independence criterion and Loewe's additivity models were used to analyze the inhibition of interferon gamma ($IFN\gamma$) production induced by co-exposure to mycotoxins, patulin, gliotoxin, citrinin and ochratoxin A on the human peripheral blood mononuclear cells (Tammer *et al.*, 2007). Dose-response data for the individual inhibition of $IFN\gamma$ production by each mycotoxin and the mixture inhibition were generated. The dose-response relationships of the individual substances and the mixture were biometrically modelled by fitting the Hill-model to the experimental data set using the best-fit approach. The dose-response functions of individual mycotoxins allowed predictions of the additive responses based either on Bliss' independence criterion or Loewe's additivity models. For the combined effect of the doses of mycotoxin that individually induced a 20% inhibition of $IFN\gamma$ production the predicted values for additivity were 59% for Bliss independence criterion model and 79% for Loewe's additivity model. Compared to the 69% inhibition of $IFN\gamma$ production that was actually induced for the co-exposure, the conclusion is that the combined effect for the four mycotoxins appeared synergistic based on Bliss' independence criterion model and antagonistic based on Loewe's additivity

model. There are still ongoing debates on which is the “better” or even the “correct” concept (Boedeker and Backhaus, 2010); however, Loewe’s additivity model is slightly preferred because of an overall higher biological plausibility (Goldoni and Johansson, 2007; Kortenkamp *et al.*, 2009). It is likely that most toxic substances exert actions that are not completely different and independent from those of other toxicants due to converging signaling pathways and inter-linked subsystems.

The isobolograms

The isobolographic method can be considered the graphical variant of the interaction index method (Tallarida, 2011). In a Cartesian coordinate system, doses of each agent that give a specified effect e.g. EC₅₀ are represented on the x- and y-axes. The straight line connecting the equally effective doses of the agents is assumed to represent the set of dose pairs that give the specified effect in a situation of no interaction (additivity). Actual dose pairs that give the specified effect are then experimentally determined and reported on the graph. Experimental dose pairs lie below the additivity line in synergistic associations, and above in antagonistic associations.

Isobolograms were drawn to analyze *in vitro* the interactions for *Penicillium* mycotoxins and *Fusarium* mycotoxins as well (Bernhoft *et al.*, 2004; Luongo *et al.*, 2006; Luongo *et al.*, 2008). However, the isobolographic method fails to take into account the variability of the data, and there is a need for further development of statistical methods to characterize accurately the interaction of combination of agents (Gennings *et al.*, 1990). In a study investigating the type of interaction for the combined effects of four mycotoxins (fumonisin B₁, α -zearalenol, nivalenol and deoxynivalenol) on porcine whole-blood cellular proliferation, dose-response data were

generated for a range of six different doses of each mycotoxin individually and for six doses of their binary mixtures at fixed ratios (Luongo *et al.*, 2008). For the analysis of each binary combination, the IC₅₀ value with its confidence interval for each mycotoxin alone was determined and represented on the x- or y-axis of an isobologram. The dose of mixture that corresponded to a 50% inhibition of cell proliferation was also estimated and subsequently, the corresponding doses for each mycotoxin and their confidence intervals were reported on the isobologram. An additivity line was drawn to connect the x- and y-axis at the levels of the individual IC₅₀s. However the study could not indicate the confidence band that was associated to the additivity line though the uncertainties for the doses of each mycotoxin in the mixture were represented. Hence, strong conclusions could not be drawn concerning the position of the mixture point regarding the additivity line, especially to exclude additivity.

The Median Effect Principle of the Mass action law

Another concept that is independent of the mode of action and just considers both the potency (EC₅₀) and the shape of the dose-effect curve for each chemical agent and their mixture has been proposed (Chou, 2011). In the so-called Chou-Talalay method, the mass-action law allows a computer simulation of the individual dose-effect curves and the “no interaction” response that could be expected from the combined effect of several agents (Chou, 2006). Individual agents and their mixtures dose-effect relationship are biometrically modelled using the median-effect equation of the mass action law that is:

$$f_a / f_u = (D / D_m)^m$$

Where D is the dose of the agent (*e.g.* a cytotoxic mycotoxin), f_a is the fraction affected by D (*e.g.* percentage of viability inhibition/100), and f_u is the fraction unaffected (*i.e.* $f_u = 1 - f_a$). D_m is the median-effect dose (*e.g.* IC_{50}), and m is the coefficient signifying the shape of the dose-effect relationship ($m = 1$, $m > 1$, and $m < 1$ indicate hyperbolic, sigmoidal, and flat sigmoidal dose-effect curves, respectively).

Then interactions can be analyzed by a combination index- isobologram method derived from the median-effect equation. The combination index (CI) is calculated according to (Chou, 2011):

$${}^n(CI)_x = \sum_{j=1}^n \frac{(D)_j}{(D_x)_j}$$

Where ${}^n(CI)_x$ is the combination index for n agents at $x\%$ inhibition, $(D)_j$ are the doses of n agents that exerts $x\%$ inhibition in combination, $(D_x)_j$ are the doses of each of n agents alone that exerts $x\%$ inhibition. $CI = 0.9-1$, $CI < 0.9$, and $CI > 1.1$ indicate an additive effect, a synergism, and an antagonism, respectively, regardless of the mechanisms or the units of the agents. Besides indicating the type of interaction (additivity, synergy or antagonism), this index allows a quantitative assessment of the magnitude of the interaction.

The Combination index-Isobologram method also known as the Chou-Talalay method that was tentatively introduced several years ago in the field of mycotoxin mixture assessment, is gaining the interest of an increasing number of researchers (Koshinsky and Khachatourians, 1992; Jones *et al.*, 1995; Ruiz *et al.*, 2011b; Lu *et al.*, 2013; Tatay *et al.*, 2014; Wang *et al.*, 2014). We used this approach to analyze the interactions for the combined toxicity of *Fusarium* mycotoxins DON, NIV and their acetylated derivatives 3- and 15-acetyldeoxynivalenol (3- and 15-ADON) and fusarenon-X (FX) in human intestinal epithelial cells (Alassane-Kpembi *et al.*, 2013). Dose-

response data for individual mycotoxins and their mixtures were generated and dose-response relationships were biometrically modeled using the Median-Effect Equation of the Mass-Action Law. The combination index values were then calculated over the range of the cytotoxicity observed. Binary or ternary mixtures of type B trichothecenes (DON, NIV, and their acetylated derivatives) demonstrated mainly synergistic cytotoxicity at low mycotoxin concentrations (cytotoxic effect between 10 and 30-40 %). At higher concentrations (cytotoxic effect around 50 %), the combinations had an additive or nearly additive effect. The magnitude of the synergistic interaction for 10% cytotoxicity was evaluated to range from 2 to 9.

3. Combined toxicity of mycotoxins

Using either of the methodological approaches described above, several teams have thoroughly examined the combined toxicity of mycotoxins *in vitro* and *in vivo*. The *in vivo* experiments have been reviewed elsewhere (Grenier and Oswald, 2011) and will not be discussed in this review. *In vitro* bioassays have obvious limitations; nonetheless they are less restrictive in the number of test groups which makes the assessment of complex mycotoxin mixtures easier. In particular, *in vitro* experiments allow for dose-response analysis of the individual contaminants and the mycotoxin mixtures. We will now review the *in vitro* experiments investigating the combined toxicity of mycotoxins. In most combined toxicity studies, the mycotoxins tested were grouped based on (i) a shared community in chemical structures (i.e aflatoxins or type B trichothecenes); (ii) toxicological modes of action (i.e mutagenic mycotoxins or carcinogenic mycotoxins), or (iii) their simultaneous production by a given fungi (i.e *Fusarium* mycotoxins or *Aspergillus* mycotoxins).

3.1. Aflatoxins and other mycotoxins

The aflatoxins are a group of closely related highly substituted coumarins containing a fused dihydrofurofuran moiety. Four aflatoxins may occur naturally: the two blue fluorescent toxins (B₁, B₂) that are characterized by fusion of a cyclopentenone ring to the lactone ring of the coumarin moiety, and the two greenish yellow fluorescent toxins (G₁, and G₂) that contain a fused lactone ring. AFB₂ and AFG₂ are considered relatively nontoxic unless they are first metabolically oxidized to AFB₁ and AFG₁ *in vivo*. The metabolism of aflatoxin B₁ and B₂ in the mammalian body may result in two metabolites M₁ and M₂ as hydroxylated derivatives of the parent compound. Aflatoxins are hepatocarcinogenic agents in numerous animal species and have been implicated in the etiology of human hepatocellular carcinoma (Wild and Montesano, 2009).

In association with other mycotoxins, the mutagenic and cell viability effects of aflatoxins have been frequently questioned. Nearly all the papers addressing mutagenic activity of these mycotoxin combinations referred to the well-known Ames test using *Salmonella* Typhimurium strains TA 100 and TA 98 (Sedmikova *et al.*, 2001; Kuilman-Wahls *et al.*, 2002; Vilar *et al.*, 2003). However, a bioluminescence test using the marine bacterium *Photobacterium phosphoreum* strain NCMB 844 was also proposed (Yates *et al.*, 1987). Mutagenicity was analyzed with a dark mutant of this organism whose reversion to the bioluminescent condition is stimulated by compounds attacking guanine sites in desoxyribonucleic acids. Aflatoxin combinations have been assessed for their cytotoxic and genotoxic effects mainly in human and animal primary hepatocytes or transformed cell lines (Friedman *et al.*, 1997; He *et al.*, 2010; Ribeiro *et al.*, 2010; Corcuera *et al.*, 2011). Aflatoxin combinations are also considered potentially immunotoxic, thus they have been evaluated for their combined effects on the

viability and functionality of immune system cells (Theumer *et al.*, 2003; Herzog-Soares and Freire, 2004; Russo *et al.*, 2010; Theumer *et al.*, 2010; Russo *et al.*, 2011).

Publications related to mycotoxin mixtures involving aflatoxins have been grouped in studies presenting the combined effects of (i) the different aflatoxins, (ii) aflatoxin B₁ and other possibly carcinogenic mycotoxins, (iii) aflatoxin B₁ and other mycotoxins from *Aspergillus* and (iv) aflatoxin B₁ and mycotoxins from *Fusarium*.

Combined toxicity of the different aflatoxins

Arithmetic definition of additivity was used in a large part of the combined toxicity studies for the different aflatoxins that are presented in Table 2. Cell viability as an endpoint showed synergy for the combined toxicity of AFB₁ and AFB₂ in human umbilical vein endothelial cells while additivity was observed for the same endpoint in human lung fibroblast and in human ovarian cancer cell line A 2780 (Braicu *et al.*, 2010). No interaction was reported between AFB₁ and AFB₂ for RNA synthesis and membrane integrity in rat hepatocytes primary culture, while an undetermined interaction was revealed between AFB₁ and AFG₁ (Friedman *et al.*, 1997). The immunotoxic interactions between aflatoxin metabolites AFM₁ and AFM₂ excreted in milk and between these metabolites and their parent-compounds AFB₁ and AFB₂ have been investigated (Russo *et al.*, 2010; Russo *et al.*, 2011; Bianco *et al.*, 2012b). No interaction could be detected when macrophages were co-exposed to AFM₁ and AFM₂, while their combinations with the parent-compounds AFB₁ and AFB₂ resulted in stronger toxicity compared to individual toxins, suggesting a synergism. Naturally-occurring mixtures of aflatoxins, i.e. aflatoxins B₁, B₂, G₁ and G₂ have been rated as carcinogenic to humans (group 1) and the metabolite AFM₁ possibly

carcinogenic to humans (group 2B) by the International Agency for Research on Cancer (IARC). However we reported no publication analyzing *in vitro* the combined genotoxicity of the aflatoxins.

Combined toxicity of aflatoxins and other possibly carcinogenic mycotoxins

Besides aflatoxins, OTA and FB₁ are the only other mycotoxins that have been demonstrated to be carcinogenic in laboratory animals and rated as possible human carcinogens (group 2B). Co-exposure to these carcinogenic mycotoxins is not uncommon. Dietary exposure assessment in several villages in Tanzania showed that up to 82% of children tested were positive for blood AFB₁-albumin adducts and urinary FB₁ (Shirima *et al.*, 2013). Co-occurrence of AFB₁, OTA and FB₁ was also detected in 20% of randomly collected cereal and feed samples from households of an endemic nephropathy area in Croatia (Klaric *et al.*, 2009). This explains the interest to investigate the interaction between aflatoxins and these mycotoxins especially in term of carcinogenic effect (Table 3).

Arithmetic definition of additivity was mainly used in these combined toxicity studies. Antagonistic cytotoxicity in human hepatoma cells HepG2 has been strongly demonstrated for AFB₁ and FB₁ by the calculation of their interaction index (McKean *et al.*, 2006b). Conflicting conclusions for the clastogenic effect of AFB₁-OTA association exist. A quantitative analysis of DNA fragmentation in monkey kidney Vero cells exposed to both mycotoxins simultaneously, suggested an additive effect (El Golli-Bennour *et al.*, 2010). This conclusion was made based on the calculation of a ratio of expected to observed IC₅₀ values for the mycotoxin mixture. However, the authors did not specify how the expected IC₅₀ value was obtained. On the contrary,

OTA was shown to reduce the DNA damage caused by AFB₁ alone in HepG2 cell line, while an increase of the mutagenic effect of AFB₁ in presence of OTA was reported using the *S. Typhimurium* mutagenicity test (Sedmikova *et al.*, 2001; Corcuera *et al.*, 2011). The authors speculated that AFB₁ and OTA could compete for the same CYP enzymes that represent a bio-activation route for AFB₁, and a higher affinity of OTA for the CYPs involved could result in less AFB₁ bio-activated molecules (AFB₁-epoxide) to attack and damage DNA.

No enhancement of the clastogenic effect has been noted when combining FB₁ and AFB₁ in rat primary hepatocyte and spleen mononuclear cell culture, whereas biomarkers of oxidative stress were lowered by the mixture compared to the individual AFB₁ effect (Ribeiro *et al.*, 2010; Theumer *et al.*, 2010).

With respect to immunotoxic effects, the AFB₁- FB₁ mixture was more effective in reducing the mitogenic response and cytokine production of mononuclear cells on the one hand and H₂O₂ release of adherent peritoneal cells on the other, compared to the individual mycotoxins (Theumer *et al.*, 2003).

Combined toxicity of aflatoxins and other mycotoxins from Aspergillus species

Citrinin (CIT) and cyclopiazonic acid (CPA) are mycotoxins produced by *Aspergillus* and/or *Penicillium* strains that have been frequently associated to AFB₁ for mixture toxicity studies (Table 4). A number of *Aspergillus* strains that produce B- and G-type aflatoxins may also produce CPA (Lee and Hagler, 1991; Pildain *et al.*, 2008). As a consequence CPA and aflatoxins often co-contaminate crops (Urano *et al.*, 1992; Chang *et al.*, 2009). Likewise, aflatoxins and

citric acid have been simultaneously detected in various food and feed commodities (Kpodo *et al.*, 1996; Garon *et al.*, 2006; Nguyen *et al.*, 2007; Richard *et al.*, 2009).

All the studies reported for the combined toxicity of aflatoxins and CPA or CIT defined their reference point using the arithmetic definition of additivity. Following metabolic activation by either human S-9 mix or rat S-9 mix, the mutagenic activity of AFB₁ and CPA combination assessed by reverse mutation of *S. Typhimurium* TA 98 and TA 100 strains constantly resulted in a reduction compared to AFB₁ individual effect (Kuilman-Wahls *et al.*, 2002; Vilar *et al.*, 2003). This reduction of the AFB₁ mutagenicity by CPA was attributed to the inhibitory effect of CPA on cytochrome P450 (CYP450) 3A4 activity. On the opposite, the marine bacterium *P. phosphoreum* reverse mutation test revealed an enhanced genotoxic effect for AFB₁ in mixture with CPA (Yates *et al.*, 1987).

Combined toxicity of aflatoxin B₁ and mycotoxins from Fusarium species

The simultaneous spoilage of food commodities by *Aspergillus* and *Fusarium* strains is not uncommon and may be associated to natural co-occurrence of aflatoxins and various *Fusarium* mycotoxins, including DON, NIV and ZEA (Ali *et al.*, 1998; Almeida *et al.*, 2012). Data on the combined toxicity of aflatoxins and these fusariotoxins are presented in Table 5.

Interactive cytotoxicity between AFB₁ and *Fusarium* toxins ZEA and DON has been demonstrated at low doses and high doses in porcine kidney cells using the factorial design approach (Lei *et al.*, 2013). Synergy for cytotoxicity has been previously shown between AFB₁ and T-2 toxin by calculation of their interaction index (McKean *et al.*, 2006a). It is noteworthy that the type of toxic interaction in cell viability between AFB₁ and *Fusarium* toxins, especially

trichothecenes, may depend on the cell model as additivity was reported in fish primary hepatocytes and human hepatoma cells (HepG2) while synergy was observed in human bronchial epithelial cells (BEAS-2B) (McKean *et al.*, 2006a; He *et al.*, 2010).

Surprisingly, the mutagenic activity of AFB₁ was significantly enhanced by the trichothecene mycotoxins DON and T-2 toxin which demonstrated no individual effect by their own in the *Salmonella* prokaryote mutagenicity test (Smerak *et al.*, 2001). However, the authors reported a significant clastogenic effect for the trichothecene mycotoxins that may explain the enhanced mutagenic outcomes of the activity of AFB₁ in presence of either or both trichothecenes.

3.2. Ochratoxins and other mycotoxins

Ochratoxins are produced by several species belonging to both *Aspergillus* and *Penicillium* genera. Ochratoxin A (OTA) is toxic to several organs, especially the kidney, whereas its dechloro-analogue ochratoxin B only displays limited toxicity (Roth *et al.*, 1989; Heussner *et al.*, 2006). Studies addressing the toxicity of ochratoxins in association with other mycotoxins mainly concern OTA.

As already mentioned, OTA is a nephrotoxic compound, and as a consequence, most of the studies involved renal cell lines or renal primary cells cultures. Cytotoxicity is the main endpoint explored for the mycotoxin combinations. Mycotoxins associations including ochratoxins have also been screened for genotoxicity via DNA damages, clastogenic effects and mutagenic activity (Knasmuller *et al.*, 2004). As far as the immune system is concerned, mitogen-induced lymphocyte proliferation and cytokine production were the main endpoints for papers addressing the combined toxicity of ochratoxin (Table 6 and Table 7).

Combined toxicity of ochratoxins and other mycotoxins from Aspergillus or Penicillium

Ochratoxins may co-occur with other mycotoxins produced by species from *Aspergillus* and *Penicillium* genera. Among these mycotoxins, citrinin (CIT) is the most frequently associated with OTA, as illustrated by several studies undertaken in Bulgaria, Croatia and Serbia that showed that populations in endemic nephropathy regions were more frequently exposed to OTA and CIT due to microclimatic conditions (Klaric *et al.*, 2013). The combined toxicity of ochratoxins with patulin (PAT), cyclopiazonic acid (CPA), gliotoxin (GLIO), roquefortin (ROQ), penicillic acid (PA) and sterigmatocystin (STER) was also studied (Bernhoft *et al.*, 2004; Heussner *et al.*, 2006; Tammer *et al.*, 2007; Anninou *et al.*, 2014). The publications related to ochratoxins and other mycotoxins from *Aspergillus* and *Penicillium* are presented in Table 6.

OTA and CIT have mostly been reported to act in a synergistic manner for their cytotoxic (Roth *et al.*, 1989; Bouslimi *et al.*, 2008a; Bouslimi *et al.*, 2008b; Klaric *et al.*, 2012) and their genotoxic effects (Knasmuller *et al.*, 2004). The co-exposure of human kidney cells (HK-2) with both mycotoxins increased DNA adduction and CYP 450 and peroxydase enzymes expression (Manderville and Pfohl-Leszkowicz, 2008; Pfohl-Leszkowicz *et al.*, 2008). However, both mycotoxins failed to induce reverse mutation in the Ames test, and showed antagonism in porcine kidney PK15 epithelial cells (Wurgler *et al.*, 1991; Klaric *et al.*, 2012). Considering immunotoxicity endpoints a synergy between OTA and CIT was observed for mitogen-induced lymphocyte proliferation, while an additive effect was observed for the inhibition of IFN- γ production by peripheral blood mononuclear cells (Bernhoft *et al.*, 2004; Tammer *et al.*, 2007). Except CPA that showed an antagonistic immunotoxicity, the combinations of OTA with other

Aspergillus and *Penicillium* mycotoxins were reported as additive. However, most of these studies rely on an arithmetic definition of additivity.

Combined toxicity of ochratoxins and mycotoxins from Fusarium species

Publications related to the combined toxicity of ochratoxins and mycotoxins produced by *Fusarium* species mainly concern OTA and fumonisin B₁ (FB₁), and in a lesser extent OTA and the emerging mycotoxin beauvericin (BEA) (Table 7). All of these studies considered the arithmetic definition of additivity as the reference point. Conflicting conclusions have been reported for the interaction between OTA and FB₁ for cytotoxicity including synergism (Creppy *et al.*, 2004; Carratu *et al.*, 2005; Mwanza *et al.*, 2009) and addition (Klaric *et al.*, 2007; Klaric *et al.*, 2008b). Genotoxic potential for binary combinations of OTA, FB₁ and BEA were mainly depicted as additive (Klaric *et al.*, 2007; Klaric *et al.*, 2008a).

3.3. *Fusarium* mycotoxins

Fusarium species can produce a wide variety of mycotoxins. The most common *Fusarium* mycotoxins that occur at biologically significant concentrations in food chain are fumonisins, zearalenone and trichothecenes (Placinta *et al.*, 1999). The mycotoxin association patterns involved the "major" mycotoxins from *Fusarium*, the trichothecenes, FB₁ and ZEA, although increasing attention is being paid to combinations including the emerging mycotoxins beauvericin (BEA) and enniatins (ENN) (Ruiz *et al.*, 2011a; Ruiz *et al.*, 2011b; Ficheux *et al.*, 2012; Kolf-Clauw *et al.*, 2013; Lu *et al.*, 2013). Combined toxicity studies mainly investigated

the cytotoxicity or immunotoxicity of *Fusarium* mycotoxins. Due to its estrogenic activity, mixtures including ZEA have also been tested on the reproductive system cells (Malekinejad *et al.*, 2007; Ranzenigo *et al.*, 2008). Among publications concerning the combined effect of *Fusarium* mycotoxins, we have separately considered (i) those concerning the combined effects of trichothecene mycotoxins, (ii) those concerning the major mycotoxins from *Fusarium*, and (iii) other studies on combined toxicity of mycotoxins from *Fusarium*.

Combined toxicity of trichothecenes

Publications analyzing the combined toxicity of trichothecenes are presented in Table 8. Not all the trichothecenes involved in association studies are *Fusarium* mycotoxins. The non-macrocylic trichothecenes produced by *Fusarium* species have been combined to the macrocylic trichothecenes roridin A and verrucarin A produced by *Myrothecium* species (Koshinsky and Khachatourians, 1992; Jones *et al.*, 1995).

Combination of the type B trichothecenes DON or NIV to the type A T-2 toxin or DAS resulted in additive or antagonistic response either for the cytotoxic or the immunotoxic endpoints (Thompson and Wannemacher, 1986; Thuvander *et al.*, 1999; Ruiz *et al.*, 2011a; Ruiz *et al.*, 2011b). However, the striking fact of the combined toxicity of this group of mycotoxins is that the type and intensity of interactions vary accordingly with the tested doses and the combination ratios. Using human and porcine intestinal cells as well, we observed a synergistic cytotoxicity when combining DON, NIV and their acetyl derivatives at low doses while the interaction was additive to antagonistic for higher mycotoxin doses (Alassane-Kpembi *et al.*, 2013; Alassane-Kpembi *et al.*, 2015). Likewise, the interactive immunotoxicity of DON and NIV mixture is

thought to be limited to low doses (Severino *et al.*, 2006). Earlier, it has also been shown that the interaction between the type A trichothecenes T-2 toxin and HT-2 toxin and the type D trichothecene roridin A changes from antagonistic to synergistic for graded toxicity levels towards the yeast *Kluyveromyces marxianus* (Koshinsky and Khachatourians, 1992).

Except factorial designs, all kinds of methodological approaches have been used for the elucidation of the type of interaction for mixtures involving trichothecenes. However, unlike early discussed mycotoxin groups, a number of studies in this group can be considered reliable enough for their conclusions since they are not built on mistaken interaction analysis approaches.

Combined toxicity of the "major" mycotoxins from Fusarium

Most joint toxicity studies are related to simultaneous contamination by type B trichothecenes (DON and or NIV), FB₁ and ZEA or its alcohol metabolite α -zearalenol (Table 9). Using the factorial designs the combination between the main *Fusarium* mycotoxins was shown to act additively on the porcine Ipec J2 cell viability reduction while interaction occurred for pro-inflammatory cytokines mRNA expression and the modulation of the expression of β -defensins 1 and 2 (Wan *et al.*, 2013a; Wan *et al.*, 2013b; Wan *et al.*, 2013c). Their toxicity was also found additive for the inhibition of DNA synthesis in mouse fibroblast by the same methodological approach (Groten *et al.*, 1998; Tajima *et al.*, 2002). In binary association, synergy was reported between ZEA or its alcohol metabolite α -zearalenol and FB₁ for various endpoints and cell systems (Groten *et al.*, 1998; Tajima *et al.*, 2002; Luongo *et al.*, 2006; Kouadio *et al.*, 2007; Luongo *et al.*, 2008). The synergy may also exist for the combined anti-proliferative effect of

ZEA and DON on porcine granulosa cell but it was not confirmed for other endpoints in pig reproductive toxicology (Malekinejad *et al.*, 2007; Ranzenigo *et al.*, 2008).

Other studies on combined toxicity of mycotoxins from Fusarium

The emerging *Fusarium* toxins beauvericin (BEA) and enniatins (ENN) have been involved in mycotoxins combined effects studies for their cytotoxic and genotoxic potential (Table 10). Binary and ternary mixtures of ENN A, A₁, B and B₁ clearly exerted synergistic cytotoxicity on ovarian cells and intestinal cells (Lu *et al.*, 2013; Prosperini *et al.*, 2014). On the contrary, the toxicity of T-2 toxin was down-modulated by ENN B₁ in pig intestinal epithelial cells and explants culture (Kolf-Clauw *et al.*, 2013). Assuming the arithmetic definition of additivity, no interaction could be detected in combined myelotoxicity for ENN B and BEA, while synergy was shown in BEA and DON mixture (Ficheux *et al.*, 2012). In other cell lines and by means of the Chou-Talalay method antagonism was observed for the latter association (Ruiz *et al.*, 2011a; Ruiz *et al.*, 2011b). This cell line-related discrepancy was also noted for the combined toxicity of BEA and the type A trichothecene T-2 toxin (Ruiz *et al.*, 2011a; Ruiz *et al.*, 2011b). The combination of BEA to FB₁ led to an additive induction of apoptosis in mononuclear cells (Dombrink-Kurtzman, 2003).

Conclusion

For the main mycotoxin groups, reference doses for regulatory purpose already exist. Exposure below these levels is usually considered safe. Whether the consumer is also protected against combined exposure to mycotoxins if each component is present below its individual threshold

dose is gaining increasing interest. The present review analyzed the methodological aspects and main conclusions for the publications related to the toxicological interactions of mycotoxins.

More than eighty publications have been dedicated to the combined toxicity of mycotoxins, especially *Fusarium* toxins. Besides the regulated mycotoxins, an increasing number of studies are paying attention to mixtures involving the “emerging” ones. Considering the increasing attention given to modified mycotoxins; we can anticipate that their combined toxicity will be studied (Alassane-Kpembé *et al.*, 2015; Pierron *et al.*, 2015). Many methodological approaches have been used to explore the interactions in combined toxicity of mycotoxins. The main approaches are (i) the arithmetic definition of additivity, (ii) the factorial designs and (iii) the theoretical biology-based Combination index-isobologram method. A crucial issue for toxicodynamic interaction analysis is the statement of the non-interaction response. Factorial designs allow a reliable detection of departure from the additive response, while the Combination index-isobologram method makes it possible to determine the type of the interaction and to optionally quantify its magnitude. Only a few papers used these approaches for mycotoxin interaction analysis and most of them concern the combined toxicity of *Fusarium* toxins. Out of 35 publications only 13 used the isobologram approach and 4 used factorial designs.

Many biological models with different metabolic abilities along with various mycotoxin association patterns have been used. The biological models include human or animal primary cells or non-transformed or immortalized cell lines as well as prokaryote models. This review gathered the mycotoxins according to their producing fungi and indicates that *Fusarium* mycotoxins were the most studied. However, other mycotoxin combination strategies could be considered, as the mycotoxin co-occurrence patterns in commodities and the co-exposure

patterns reported in bio-monitoring studies indicate that humans and animals are exposed to a wide variety of mycotoxin combinations in real life.

The main conclusion from all these studies is that very few studies used a robust methodological approach for the analysis of the combined effect of mycotoxins, and the type of interaction in terms of additivity, synergy or antagonism varies accordingly with the mycotoxin combinations, and even with the concentrations tested. More studies employing the isobologram approach are needed to feed a reliable database for the interactions between mycotoxins. Several publications reported synergy, especially for *Fusarium* toxins, using the Combination index-isobologram method. These *in vitro* synergistic interactions should be confirmed *in vivo*.

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Table 1: Selected mycotoxins' co-exposure/co-occurrence patterns reported worldwide

Co-exposure/ Co-contamination patterns	Sampling	Methodological approach	References
DON-ENNB-ZEA ; DON-CIT-T2 ; DON-CIT ; DON-ZEA ; DON-ENNB	Urine samples from 101 German adult volunteers	LC-MS/MS urinary multi-biomarker approach	(Gerding <i>et al.</i>, 2014)
DON-CIT-OTA-FB₁; DON-CIT-OTA; DON-OTA-ENNB CIT-OTA-ENNB ; CIT-OTA-FB₁ ; AFM₁-CIT-OTA ; AFM₁-CIT-DON ; ENNB-OTA ; DON-OTA ; CIT-OTA ; CIT-FB₁ ; CIT-ENNB AFM₁-CIT	Urine samples from adult volunteers:		

-95 Bangladeshis -50 Germans -142 Haitians	LC-MS/MS urinary multi-biomarker approach	(Gerding <i>et al.</i>, 2015)	
DON-ZEA-FB₁-OTA-AFB₁	Urine samples from 52 Italian adult volunteers	LC-MS/MS urinary multi-biomarker approach	(Solfrizzo <i>et al.</i>, 2014)
AFB₁-FB₁-DON	Blood and urine samples from 148 Tanzanian children aged 12-22 months	Albumin ELISA, HPLC and HPLC/MS	(Shirima <i>et al.</i>, 2013; Srey <i>et al.</i>, 2014)
AFM₁-FB₁-OTA-DON-NIV ; FB₁-FB₂-OTA-NIV ; FB₁-DON-NIV ; DON-ZEA-NIV ; OTA-NIV	Urine samples from 175 Cameroonian HIV-positive and HIV-negative adult volunteers	LC-MS/MS urinary multi-biomarker approach	(Abia <i>et al.</i>, 2013)
AFB₁-AFB₂-AFG₁-AFG₂-FB₁-FB₂-...-BEA-DON-NIV-ZEA-CIT-FA-ENNB₁; AFB₁-AFB₂-...-FB₁-...-OTA-BEA-STER-ZEA	122 maize, millet, infant food and feed samples from Burkina Faso and Mozambique	LC-MS/MS multi-toxin method	(Warth <i>et al.</i>, 2012)
FBs-DON-ZEA-AFs-OTA	92 commercial compound feeds from South Africa	LC-MS/MS multi-toxin method	(Njobeh <i>et al.</i>, 2012)

AFs-ZEA; AFs-OTA; OTA-ZEA; FBs-ZEA	37 randomly collected cereal and feed samples from households in endemic nephropathy areas (Croatia)	ELISA and Thin-layer chromatography	(Klaric <i>et al.</i>, 2009)
DON-NIV-BEA-ENNs	93 oat samples collected in 2010 and 2011 from field trials and grain delivery stations in central and southern Sweden	HPLC/ESI-MS/MS	(Fredlund <i>et al.</i>, 2013)

*Abbreviations used: AFs= aflatoxins, BEA= beauvericin, CIT= citrinin, DON= deoxynivalenol, ENN= enniatin, FA= fusaric acid, FBs= fumonisins, NIV= nivalenol, OTA= ochratoxin A, STER= sterigmatocystin, ZEA= zearalenone

Table 2: Interactions between aflatoxins

Mycotoxin association*	Interaction model	Cell model	Study design	Endpoint	Combined effect	Reference
AFB₁-AFB₂-AFG₁	Arithmetic definition of additivity	J774A.1 murine macrophages	Comparison of combination treatment with 0.01 ng/mL AFB ₁ + 0.01 ng/mL AFB ₂ to treatment with AFB ₁ or AFB ₂ alone	Cytokine secretion	Alleged synergy AFB ₁ - AFB ₂ at the lower dose for increase in IL-6 secretion	(Bruneau et al., 2012)
AFB₁-AFB₂-AFG₁-AFG₂	Arithmetic definition of additivity	Human umbilical vein endothelial cells (HUVEC), human lung	Comparison of IC ₅₀ values for mixtures and indi	Cell viability	Alleged synergy AFB ₁ - AFB ₂ in HUVEC cells, additivity (AFB	(Braicu et al., 2010)

		fibroblasts (HFL), and A2780	vidual toxins		¹⁷ AFB ₂ - AFG ₁ - AFG ₂) in HFL and A2780	
AFB₁-AFB₂-AFG₁	Two-way ANOVA	Rat hepatocytes, rat liver slices	Comparison of the effects of serial dilutions of AFB ₁ (0-480 ng/mL) in presence or absence of AFB ₂ at 120 ng/mL, and comparison of the effects of serial dilut	RNA synthesis, membrane integrity	No interaction AFB ₁ - AFB ₂ , undetermined interaction AFB ₁ - AFG ₁	(Friedman <i>et al.</i>, 1997)

			ions of AFB ₁ (0-480 ng/mL) in presence or absence of AFG ₁ at 120 ng/mL or 240 ng/mL			
AFB₁-AF B₂-AF M₁-AF M₂	Arithmetic definition of additivity	J7741.A murine macrophages	Comparison of the effects of serial dilutions of individual toxins and their combination	Cell viability, activation of macrophagic functions (Nitric oxide production)	Stronger effects of mixtures compared to individual toxins suggesting interactions	(Russo <i>et al.</i>, 2011)
AFB₁-AF B₂-AF M₁-AF M₂	Arithmetic definition of additivity	J774A.1 murine macrophages	Comparison of IC ₃₀ values for viability	Cell viability, apoptosis, inhibition of nitric	Stronger effects of mixtures compared to indivi	(Bianco <i>et al.</i>, 2012b)

			reduction for individual toxins and mixtures, comparison of nitric oxide production inhibition by graded levels of individual toxins and mixtures	oxidation	dual toxins suggesting interactions	
AFM₁-AFM₂	Arithmetic definition of additivity	J7741.A murine macrophages	Comparison of the effects of serial dilutions of	Cell viability, activation of macrophagic functions	No interaction	(Russo <i>et al.</i>, 2010)

			indi vidu al toxi ns and their com binat ion	(Nitr ic oxid e prod uctio n)		
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*Abbreviations used: AFB₁= aflatoxin B₁, AFB₂= aflatoxin B₂, AFG₁= aflatoxin G₁, AFG₂=aflatoxin G₂,
AFM₁=aflatoxin M₁, AFM₂=aflatoxin M₂

Table 3: Interactions between Aflatoxin B₁ and carcinogenic or possibly carcinogenic mycotoxins

Mycotoxin association*	Interaction model	Cell model	Study design	Endpoint*	Combined effect	Reference
AFB₁-FB₁	Interaction index metric	Human hepatoma cells (Hep G2), Human bronchial epithelial cells (BEAS-2B)	Dose-response curves and determination of IC ₅₀ values for individual toxins and their mixture	Cell viability	Demonstrated additivity BEAS-2B, antagonism Hep G2	(McKeenan <i>et al.</i>, 2006b)
AFB₁-FB₁	Arithmetic definition of additivity	Rat primary hepatocytes culture	Comparison of the toxic effects of individual mycotoxin and mixture doses	Cell viability, DNA fragmentation and apoptosis	No toxicity enhancement	(Ribeiro <i>et al.</i>, 2010)
AFB₁-FB₁	Arithmetic	Rat Wistar	Comparison	Mitogenic response and	Differences in	(Theumer <i>et</i>

	definition of additivity	spleen mononuclear cells (SMC) and adherent peritoneal cells (APC)	effect of 20 $\mu\text{mol/L}$ AFB ₁ and 10 $\mu\text{mol/L}$ FB ₁ , to the effects of a mixture 20 $\mu\text{mol/L}$ AFB ₁ + 10 $\mu\text{mol/L}$ FB ₁	cytokines (IL-2, IL-4, IL-10) production of SMC and H ₂ O ₂ release of APC	the effects produced by a mixture of mycotoxins in comparison to the individual action of the same toxins	<i>al.</i> , 2003)
AFB₁-FB₁	Arithmetic definition of additivity	Rat Wistar spleen mononuclear cells (SMC)	Comparison of the individual effects of 20 $\mu\text{g/mL}$ FB ₁ and 10 $\mu\text{g/mL}$ AFB ₁ to	Genotoxicity (alkaline comet assay and micronuclei assay) and oxidative stress (malondialdehyde (MDA) levels, catalase	No difference in DNA injury, no difference in MDA level, higher CAT and	(Theumer et al., 2010)

			the effects of a mixture of 20 µg/mL FB ₁ + 10 µg/mL AFB ₁	(CAT) and superoxide dismutase (SOD) activities)	SOD activities in AFB ₁ individual treatment compared to FB ₁ , and the mixture	
AFB₁-OTA	Arithmetic definition of additivity	Human hepatoma Hep G2 cells	Comparison of IC ₅₀ values for mixtures and individual toxins	Cytotoxicity and genotoxicity	Alleged additive effect for cytotoxicity, antagonism for genotoxicity	(Corcuera <i>et al.</i> , 2011)
AFB₁-OTA	Brown Interaction index	Monkey kidney Vero cells	Calculation of a ratio of expected to observed IC ₅₀ for the mixture	Cytotoxicity and genotoxicity	Alleged additivity	(El Golli-Bennour <i>et al.</i> , 2010)

<p>AFB₁- OT A</p>	<p>Arith- m- eti- c- de- fin- iti- on- of ad- dit- ivi- ty</p>	<p><i>Salmonella</i> Typ- him- uriu- m strai- ns TA 100 and TA 98</p>	<p>Compara- ison of the muta- genic activi- ty for serial diluti- ons of indiv- idual toxins and their mixt- ure.</p>	<p>Mutagenic activity</p>	<p>Significa- nt incre- ase of the muta- geni- c activ- ity of AFB₁</p>	<p>(Sedmi- kov a <i>et</i> <i>al.</i>, 200 1)</p>
<p>AFB₁- OT A- FB 1</p>	<p>Arith- m- eti- c- de- fin- iti- on- of ad- dit- ivi- ty</p>	<p>Madin- Darby Bovine Kidney (MD BK) cells</p>	<p>Compara- ison of the toxic effec- ts of mixt- ures to the sums of the toxic effec- ts of indiv- idual comp- ound s at their conc- entra- tions in the mixt</p>	<p>Cell viabilit- y MTT, NR</p>	<p>Alleged addit- ivity for AFB₁ and AFB₁- OTA . Alle- ged syne- rgy for AFB₁- OTA terna- ry mixt- ure</p>	<p>(Clarke <i>et</i> <i>al.</i>, 201 4)</p>

<p>AFB₁- OT A- FB 1</p>	<p>Arith m eti c de fin iti on of ad dit ivi ty</p>	<p>Madin- Darb y Bovi ne Kidn ey (MD BK) cells</p>	<p>ures Com paris on of the toxic effec ts of mixt ures to the sums of the toxic effec ts of indiv idual comp ound s at their conc entra tions in the mixt ures</p>	<p>Cell viabilit y: high content analysis , MTT, NR</p>	<p>Alleged addit ivity for AFB 1- FB₁a nd AFB 1- OTA . Alle ged syne rgy for AFB 1- FB₁- OTA terna ry mixt ure</p>	<p>(Clarke <i>et al.</i>, 201 5)</p>
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*Abbreviations used: AFB₁= aflatoxin B₁, FB₁= fumonisin B₁, OTA= ochratoxin A, MTT= (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) tetrazolium, NR= Neutral Red

Table 4: Interactions between Aflatoxin B₁ and other mycotoxins from *Aspergillus* species

Mycotoxin association*	Interaction model	Cell model	Study design	Endpoint	Combined effect	Reference
AFB₁-CIT	Arithmetic definition of additivity	Mice macrophage	Comparison of infectivity and proliferation in control, individual and combined toxin groups	Infectivity and proliferation of <i>Toxoplasma gondii</i>	Increased infectivity and proliferation of <i>T. gondii</i> for the combined exposure compared to control	(Herzog-Soarens and Freire, 2004)
AFB₁-CPA	Arithmetic definition of additivity	<i>Salmonella</i> Typhimurium strains TA 100 and TA 98	Comparison of the mutagenic activity for serial dilution	Mutagenic activity following metabolic activation by rat S-9 mix	Reduction of AFB ₁ mutagenic activity	(Kuilmann-Wahls et al., 2002)

			ions of individual toxins and their mixture			
AFB₁-CPA	Arithmetic definition of additivity	<i>Salmonella</i> Typhimurium strains TA 100 and TA 98	Comparison of the mutagenic activity for serial dilutions of individual toxins and their mixture	Mutagenic activity following metabolic activation by human S-9 mix	Reduction of the mutagenic activity of AFB ₁ .	(Vilar <i>et al.</i> , 2003)
AFB₁-CPA	Arithmetic definition of additivity	Marine bacterium <i>Photobacterium phosphoreum</i> strain NCMB 844	Comparison of the effects for individual	Genotoxicity and cytotoxicity	Enhanced genotoxic effect of AFB ₁ by CPA	(Yates <i>et al.</i> , 1987)

		and strain NRRL B 1177	al toxi n dose s and dose pairs in mixt ure			
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*Abbreviations used: AFB₁= aflatoxin B₁, CIT= citrinin, CPA= cyclopiazonic acid

Table 5: Interaction between aflatoxin B₁ and mycotoxins from *Fusarium* species

Mycotoxin association*	Interaction model	Cell model	Study design	Endpoint*	Combined effect	Reference
AFB ₁ -DON	One-way ANOVA	<i>Cyprinus carpio</i> primary hepatocytes	Comparison of the effects of the mixture and the effects of individual toxins	Cell viability (MTT test), enzyme (Aspartate aminotransferase AST, Alanine transferase ALT, Lactate dehydrogenase LDH) activity in cell supernatant	Alleged additivity	(He <i>et al.</i> , 2010)
AFB ₁ -T-2 toxin	Interaction in dex metric	Human hepatoma Hep G2 cells, Human BEA S-2B bronchial epithelial cells	Dose-response curves and determination of IC ₅₀ values for individual	Cell viability	Demonstrated synergy in BEA S-2B, additivity in Hep G2	(McK <i>et al.</i> , 2006a)

			toxins and their mixture			
AFB ₁ -DO N-T-2 toxin	Arithmetic definition of additivity	Prokaryote model (<i>Salmonella</i> Typhimurium, strains TA98 and TA100)	Comparison of the effects of individual toxins and their mixtures	Mutagenic activity	Significant enhancement of the mutagenic effect of AFB ₁ (no activity for T-2 and DON alone, but greater activity for the combinations with AFB ₁)	(Smerak <i>et al.</i> , 2001)
AFB ₁ -DO N-ZEA	Factorial design	Immortalized BRL 3A rat liver cells	Central composite design for binary and ternary mixt	Cell viability (MTT)	Demonstrated interactive cytotoxicity. Alleged synergy	(Sun <i>et al.</i> , 2015)

			ures expe rime nts, with the IC ₃₀ of cell viabi lity of each myc otoxi n chos en as the cente r point		for AFB 1- ZEA and AFB 1- DON .	
AFB ₁ - DO N- ZE A	Factori al de sig n	Porcine Kidn ey PK15 cells	Central com posit e desig n for binar y and terna ry mixt ures expe rime nts, with the IC ₃₀ of cell viabi lity of each myc	Cell viabilit y (MTT), membr ane damage (LDH), apoptos is and oxidati ve stress	Demonst rated inter activ e cytot oxici ty. Alleg ed syner gism for AFB 1- ZEA, AFB 1- DON , low dose antag onis m and high	(Lei et al. , 20 13)

			otoxin chosen as the center point		dose synergism ZEA - AFB ₁ , and alleged synergism DON - AFB ₁ for oxidative damage	
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*Abbreviations used: AFB₁= aflatoxin B₁, DON= deoxynivalenol, ZEA= zearalenone, MTT= (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) tetrazolium, LDH= lactate deshydrogenase

Table 6: Interactions between ochratoxins and other mycotoxins from *Aspergillus* and *Penicillium*

Mycotoxin association*	Interaction model	Cell model	Study design*	Endpoint	Combined effect	Reference
OTA-CIT	Arithmetic definition of additivity	Human Hep G2 hepatoma cells	Comparison of the toxic effect of the mixture at a dose of 20% of the IC ₅₀ of each toxin to that produced by either of the toxins at its IC ₅₀ .	Cell viability	Alleged synergy	(Gayathri et al., 2015)
OTA-CIT	Arithmetic definition	Porcine PK15kid	Comparison of	Cell viability, apoptosis	Alleged additive	(Klaric et al.,

	nitro of add itivity	ney epit heli al cell s	the toxi c effe ct of the mixt ure to the sum of the toxi c effe cts of indi vidu al com pou nds at their conc entr atio n in the mixt ure	s, necrosis, genotoxi city	effe ct for cell viab ility , syn ergy for apo ptos is and necr osis , anta gon ism for gen oto xici ty	201 2)
OTA- CI T	Arithm etic defi niti on of add itivity	Monkey kidn ey Ver o cell s	Cytotoxi city com pari son of IC ₅₀ valu es for indi vidu al toxi ns	Cell prolifera tion, DNA damage	Alleged syn ergy	(Bousli mi et al., 200 8a)

			and their mixture, DNA damage comparison of the effects for several concentrations of individual toxins and their mixture			
OTA-CIT	Arithmetic definition of additivity	Monkey kidney Verocells	Cytotoxicity comparison of IC ₅₀ values for individual toxins and	Cell proliferation, oxidative cell damage	Alleged synergy	(Bouslimi et al., 2008b)

			their mixture, DNA damage comparison of the effects for several concentrations of individual toxins and their mixture			
OTA-CIT	Arithmetic definition of additivity	Human HK 2 kidney cells	Comparison of the toxic effects of the mixture to the toxic effects	DNA adduction, expression of CYP3A4, COX and LOX	Two-fold increase of the OTA-related DNA adduction and sign	(Mandriville and Pfohl-Leszkowicz, 2008)

			of individual compounds		significant increase of the expression of COX and LOX	
OTA-CIT	Arithmetic definition of additivity	Human proximal tubule-derived cells (HK2 cells)	Comparison of the toxic effect of the mixture to the sum of the toxic effects of individual compounds at their concentration in	Cytotoxicity, apoptosis	Alleged antagonism for apoptosis at concentration of 2.5-5 μ M CIT, additivity at concentration 7.5-15 μ M	(Knecht <i>et al.</i>, 2005)

			the mixture			
Mycotoxin association*	Interaction model	Cell model*	Study design*	Endpoint	Combined effect	Reference
OTA-CIT	Theoretical biology model-based definition of additivity	Pig renal cortical cubes	Logistic function analysis of the dose-response curve for the individual compounds and their mixture	Protein synthesis, organic ions tetraethylammonium (TEA) and paraaminohippurate (PAH) transport	Demonstrate synergy and additivity for TEA and PAH ions transport and protein synthesis	(Braunber <i>et al.</i>, 1994)
OTA-OTB-CIT	Arithmetic definition of additivity	Hepatic tissue culture cells	Comparison of the toxic effects of mixtures	Protein synthesis	Alleged slight synergy	(Roth <i>et al.</i>, 1989)

			to the sums of the toxic effects of individual compounds at their concentrations in the mixtures			
OTA- O T B- CI T- P A T	Factorial design	Porcine LL C-PK1 renal cell line	Step-wise approach: Full factorial design, then inscribed central composite design	Cell viability	Demonstrated potential synergy OTA CIT	(Heusser <i>et al.</i>, 2006)
OTA- CI T-	Loewe additivity	Peripheral blood	Compa	Cell viability, cytokine	Demonstrated	(Tammer <i>et</i>

P A T- G L I O	ty and Blis s ind epe nde nce crit erio n mo dels	d mon onu clea r cell s	riso n of actu al mixt ure toxi city data to pred icte d ones base d on conc entr atio n addi tion and resp onse addi tion conc epts	producti on	addi tivity	al., 200 7)
OTA- CI T- P A T- C P A- R O Q- Pe n Ac	Isobolo gram met hod	Piglet Lymphocytes	Dose- response curves and isobologram drawing at IC ₂₀	Mitogen- induced lymphocyte proliferation	Demonstrated synergy OT A- CIT , additivity OT A- CP A, Pen Ac- RQ,	(Bernhoff <i>et al.</i> , 2004)

					PA T- RQ, PA T- Pen Ac, anta gon ism CIT - CP A	
OTA- CI T- ST E R	Arithm etic defi niti on of add itivi ty	Human Hep 3B hep atoc ellul ar cell line	Co mpa riso n of the expe cted and the obse rved effe cts for myc otox in mixt ures and calc ulati on of the Coe ffici ents of Dru g Inter acti	Cytotoxicity, cytostati city and genotoxi city	Alleged addi tive to anta gon istic cyto toxi c and gen oto xic effe cts	(Annin ou et al., 201 4)

			on			
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*Abbreviations used: CIT= citrinin, COX= cyclooxygenase, CPA= cyclopiazonic acid, CYP3A4= cytochrome P450 3A4, GLIO= gliotoxin, IC₂₀-IC₅₀= inhibitory concentration 20-50%, LOX= lipoxygenase, OTA= ochratoxin A, OTB= ochratoxin B, PAT= patulin, Pen Ac= penicillic acid, ROQ= roquefortin, STER= sterigmatocystin

Table 7: Interactions between ochratoxins and *Fusarium* mycotoxins

Mycotoxin association *	Interaction model	Cell model	Study design	Endpoint*	Combined effect	Reference
OTA-FB₁	Arithmetic definition of additivity	Madin-Darby Bovine Kidney (MDBK) cells	Comparison of the toxic effects of mixtures to the sums of the toxic effects of individual compounds at their concentrations in the mixtures	Cell viability: high content analysis, MTT, NR	Alleged synergy	(Clarke <i>et al.</i>, 2015)
OTA-FB₁	Arithmetic definition of additivity	Madin-Darby Bovine Kidney (MDBK) cells	Comparison of the toxic effects of mixtures to the sums of the toxic effects of individual compounds at their concentrations in the mixtures	Cell viability: MTT, NR	Alleged additivity	(Clarke <i>et al.</i>, 2014)
OTA-FB₁	Arithmetic definition of additivity	Pig lymphocytes, human lymphocytes	Comparison of the toxic effect of the mixture to the sum of the toxic effects of individual compounds at their concentration in the mixture	Cell viability	Alleged synergy	(Mwanza <i>et al.</i>, 2009)
OTA-FB₁	Arithmetic definition of additivity	Porcine PK15 kidney epithelial cells	Combination of equal concentrations of two or all three mycotoxins	Clastogenic effect	Alleged additivity for presence of micro nuclei and for presence of	(Klaric <i>et al.</i>, 2008a)

					nucleoplasmic bridges	
OTA-FB₁	Arithmetic definition of additivity	Human intestinal Caco-2 cells	Comparison of the toxic effects of mixtures to the sums of the toxic effects of individual compounds at their concentrations in the mixtures	Protein synthesis inhibition	Alleged synergy	(Carratu <i>et al.</i>, 2005)
OTA-FB₁	Arithmetic definition of additivity	Monkey kidney Vero cells, human intestinal caco-2 cells, rat C6 glioma cells	Comparison of the toxic effects of mixtures to the sums of the toxic effects of individual compounds at their concentrations in the mixtures	Cell viability	Alleged synergy	(Creppy <i>et al.</i>, 2004)
OTA-FB₁-CIT	Arithmetic definition of additivity	Human peripheral blood mononuclear cells	Comparison of the toxic effect of the mixture to each of the toxic effects of individual compounds at their concentration in the mixture	Mitogen-induced cell proliferation, cell viability MTT	Stronger effect of the mixture compared to any individual compound	(Stoev <i>et al.</i>, 2009)
OTA-FB₁-BEA	Arithmetic definition of additivity	Porcine PK15 kidney epithelial cells	Comparison of the toxic effects of individual mycotoxin and binary and ternary mixture of equal	Cell viability, apoptosis	Alleged additivity for cell viability, additivity and synergy for apoptosis	(Klaric <i>et al.</i>, 2008b)

			concentrations of the toxins			
OTA-FB₁-BEA	Arithmetic definition of additivity	Porcine PK15 kidney epithelial cells	Combination of equal concentrations of two or all three mycotoxins	Cell viability, lipid peroxidation (TBARS) and GSH depletion	Alleged additivity, possibly synergy and antagonism	(Klaric <i>et al.</i>, 2007)
OTA-BEA	Arithmetic definition of additivity	Porcine PK15 kidney epithelial cells, Human leukocytes (HL)	Combination of two concentrations	Genotoxic potential	Alleged additivity and synergy in PK15, additivity in HL	(Klaric <i>et al.</i>, 2010)
OTA-ZEA	Loewe additivity and Bliss independence criterion models	Human HepG2 hepatoma cells and Immortalized murine ovarian granular KK-1 cells	Comparison of actual mixture toxicity data to predicted ones based on concentration addition and response addition concepts	Cell viability and intracellular ROS production	Demonstrated additivity for cell viability, departure from additivity for ROS production	(Li <i>et al.</i>, 2014)
OTA-ZEA-α-ZOL	CI-isobologram method	Human HepG2 hepatoma cells	Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept	Cell viability MTT	Demonstrated antagonism for OTA-ZEA, OTA- α -ZOL and OTA-ZEA- α -ZOL mixtures	(Wang <i>et al.</i>, 2014)

Table 8: Interaction between trichothecenes

Mycotoxin association*	Interaction Model	Cell model	Study design	Endpoint*	Combined effect	Reference
DON-NIV	Arithmetic definition of additivity	Rat IEC-6 intestinal epithelial cells	Incubation with graded levels of DON or NIV alone or in combination	Cell viability, apoptosis, cell migration	No additive or synergistic effects	(Bianco <i>et al.</i>, 2012a)
DON-NIV	Arithmetic definition of additivity	Murine J7741. A macrophages	Comparison of the IC ₅₀ values of the toxins	Cell viability, Proapoptotic activity	No synergy	(Marzocco <i>et al.</i>, 2009)

	ty		and the ir mi xtu re			
DON-NIV	Arith m eti c de fi ni ti on of ad di ti vi ty	Human Jurk at T cell s	Incuba tio n wit h gra de d lev els of D O N or N I V alo ne or in co mb ina tio n rat io 1:1 and 10: 1 for D O N: N I V	Lympho cyte proli ferati on and cyto kines expr essio n	Alleged inter activ e effe ct for lym phoc yte proli ferat ion, and inter activ e effe cts at lowe r conc entr atio ns (0.0 6- 4μ M) for IFN gam ma and IL-2 mR NA trans cript ion	(Severin o et al., 2006)

<p>DON-NIV-3-DON-15-ADON-FX</p>	<p>CI-Isobologram method</p>	<p>Porcine IPE C-1 intestinal epithelial cells</p>	<p>Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept</p>	<p>Cytotoxicity MTT</p>	<p>Demonstrated synergy for all binary combinations, excepted for NIV-FX (additivity), and for DON-FX (antagonism)</p>	<p>(Alassane-Kpe mbi <i>et al.</i>, 2015)</p>
<p>DON-NIV-3-DON-15-ADON-FX</p>	<p>CI-Isobologram method</p>	<p>Human intestinal Caco-2 cells</p>	<p>Comparison of actual mixture toxicity data</p>	<p>Cytotoxicity NR MTT</p>	<p>Demonstrated low-dose synergies, antagonism for ternary</p>	<p>(Alassane-Kpe mbi <i>et al.</i>, 2013)</p>

			to predicted on es base d on the M ass act ion la w co nc ept		mixt ure DO N- NIV -FX	
DON-NIV-T-2 toxin-DAS	Arith m eti c de fi ni ti on of ad di ti vi ty	Human lym pho cyte s	Compa ris on of the mi xt ure tox icit y and the ind ivi du al tox icit y of 2× the co nc ent rat	Mitogen- indu ced lym phocy te proli ferati on Cell viabi lity, Imm unog lobul in prod uctio n	Alleged addi tivit y (NIV - T2, NIV - DA S, NIV - DO N), and sligh t anta goni sm (DO N - T2, DO N - DA S)	(Thuvan der <i>et al.</i>, 1999)

			ion of tox in us ed at the co mb ine d ex po sur e		for lym phoc ytes proli ferat ion	
Mycotoxin associat ion*	Intera cti on m od el	Cell mo del	Study de sig n	Endpoint	Combin ed effe ct*	Referenc e
T-2 toxin- Verruc arin A	CI- Is ob ol og ra m	Yeast <i>Klu yver omy ces mar xian us</i>	Compa ris on of act ual tox icit y dat a for bin ary mi xtu re at var iou s rat ios to pre dic	Growth inhib ition	The type and inte nsity of inter actio ns vari ed with the com bina tion ratio s, gro wth perc ent inhi bitio n and the	(Jones <i>et al.</i>, 1995)

			ted on es ba se d on the M ass act ion la w co nc ept		gro wth med ium: In a rich med ium syne rgy over a 2- 3 log valu e conc entr atio n rang e for a 1.0 µg/ ml T-2 toxi n:0. 75µ g/ml verr ucar in A ratio	
T-2 toxin- HT-2 toxin- Roridin A	CI- Is ob ol og ra m	Yeast <i>Klu yver omy ces mar xian us</i>	Compa ris on of act ual tox icit y dat a for bin ary	Growth inhib ition	The type of inter actio n vari ed acco rdin gly to the mixt ure	(Koshins ky and Khac hato urian s, 1992)

			mi xtu re at var iou s rat ios to pre dic ted on es ba se d on the M ass act ion la w co nc ept		ratio s and the perc ent of inhi bitio n of gro wth: from anta goni sm to syne rgy for incr easi ng perc ent inhi bitio n of yeas t gro wth	
12-13 epoxytr ichothe cenes	Arith m eti c de fi ni ti on of ad di ti vi ty	Monkey Ver o cell s, rat sple en lym pho cyte s	Compa ris on of IC 50 val ue s of dif fer ent rat ios bin	Protein synt hesis inhib ition	Alleged addi tivity	(Thomps on and Wan nema cher, 1986)

			ary mi xtu res to the IC 50 val ue s of the ind ivi du al tox ins			
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*Abbreviations used: DAS= diacetoxyscirpenol, DON= deoxynivalenol, FX= fusarenon-X, IC₅₀=inhibitory concentration 50%, IFN= interferon, IL-2= interleukin 2, MTT= (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) tetrazolium, NIV= nivalenol, NR= neutral red, 3-ADON= 3-acetyldeoxynivalenol, 15-ADON= 15-acetyldeoxynivalenol

Table 9: Interactions between the “major” mycotoxins from *Fusarium*

Mycotoxin association*	Interaction model	Cell model	Study design	Endpoint*	Combined effect	Reference
DON-ZEA	Arithmetic definition of additivity	Human HCT 116 colon carcinoma cells	Comparison of the effects of 100 µM DON, 40 µM ZEA and 100+40 µM DON+ZEA	Cell viability, cell cycle, transmembrane potential and permeability transition pore opening	Alleged sub-additive response	(Bensaïssi et al., 2014)
DON-ZEA	Arithmetic definition of additivity	Porcine granulosa cells	Comparison of the effects of 30 ng/mL of DON, 30 ng/	Granulosa cell proliferation, steroidogenesis, gene expression	Alleged possibility synergism for alteration of GC proliferation	(Ranzig et al., 2008)

			mL of ZE A and bot h		atio n	
DON- Z E A	Arith m eti c de fin iti on of ad dit ivi ty	Pig oocytes	Compar ison of the effe cts of vari ous rati os lead ing to 3.1 2 μM mix ture DO N- ZE A to the effe cts of 3.1 2 μM of eac h toxi n alo ne of DO N	Abnormali ties in format ion of the meioti c spindl e, Inhibit ion of oocyte matura tion, develo pment al compe tence of mature d oocyte s after <i>in vitro</i> fertiliz ation	Alleged no syn erg y	(Malek inej ad et al., 200 7)

			ZE A mix ture			
DON- Z E A- FB 1	Arith m eti c de fin iti on of ad dit ivi ty	Human intestina l cell line Caco-2	Compar ison of the effe cts of Bin ary and terti ary mix ture s to the sum of the effe cts of eac h toxi n alo ne	Malonedia ldehyd e (MDA) produc tion, DNA and protei n synthe sis inhibit ion, DNA methyl ation, DNA fragm entatio n, cell viabili ty, lipid peroxi dation	Alleged ant ago nis m FB ₁ - ZE A for cell via bilit y, syn erg y ZE A- FB ₁ and ZE A- DO N for lipid per oxi dati on, far less tha n add itivi ty in DN A syn thes	(Kouad io et al., 200 7)

					is inhibition for binary mixtures of DON, FB ₁ and ZEA	
DON-ZEA-FB₁	Arithmetic definition of additivity	Brewing yeast strains (<i>Saccharomyces cerevisiae</i> , <i>Kluyveromyces marxianus</i> , <i>Pichia membranifaciens</i> , <i>Hansenula anomala</i> , and <i>Schizosaccharomyces pombe</i>)	Comparison of the growth with inhibition for combinations and the sum of the inhibition for toxins alone	Yeast growth	Alleged interaction at high concentration: synergism or antagonism depending on toxin combination ratios	(Boeira et al., 2000)
Mycot	Interac	Cell model	Study	Endpoint*	Combin	Referen

oxi n ass oci ati on *	tio n m od el		desi gn		ed effe ct	ce
DON- ZE A- FB r- NI V	Factor ial de sig n	Porcine intestina l Ipec J- 2 cells	Inscribe d cent ral co mp osit e desi gn wit h four toxi ns and two con cent rati ons for eac h toxi n	Cell viabili ty	Demon stra ted non - add itiv e inte ract ion s exc ept DO N- FB ₁ - ZE A	(Wan et al., 201 3a)
DON- ZE A- FB r- NI V	Factor ial de sig n	Porcine intestina l Ipec J- 2 cells	Inscribe d cent ral co mp osit e desi gn wit h four toxi ns	Pro- inflam mator y cytoki nes IL-1 α , IL-1 β , IL-6, IL-8, TNF- α and MCP- 1 mRN	Demon stra ted non - add itiv e inte ract ion s	(Wan et al., 201 3b)

			and two concentrations for each toxin	A expression		
DON-ZE A-FB ₁ -NI ₂ V	Factorial design	Porcine intestinal Ipec J-2 cells	Inscribed central composite design with four toxins and two concentrations for each toxin	Modulation of the expression of beta-defensin in 1 & 2	Alleged non-additive interactions	(Wan <i>et al.</i> , 2013c)
DON-ZE A-FB ₁ -NI ₂ V	Arithmetic definition of additive	Human Jurkat cells and porcine lymphocytes	Increasing concentrations of FB ₁ with con	Mitogen-activated lymphocyte proliferation	Alleged interaction	(Severino <i>et al.</i> , 2008)

	ivity		stant concentrations of ZEA or co-incubation with DON and NIV			
DON-ZEA-FB ₁ -NIV-T ₂ toxin	Factorial design	L929 mouse fibroblasts	Central composite design with five toxins and five concentrations for each toxin, the full fact	DNA synthesis inhibition	Demonstrated mainly additive combinations and synergy for ZEA-FB ₁ , NIV-T ₂	(Tajima <i>et al.</i> , 2002)

			oria l desi gn for two - fact or inte ract ions of part icul ar inte rest			
DON- ZE A- FB r- NI V- T- 2 tox in	Factor ial de sig n	L929 mouse fibroblas ts	Central co mp osit e desi gn wit h five toxi ns and five con cent rati ons for eac h toxi n, the n full fact oria l desi	DNA synthe sis inhibit ion	Demon stra ted mai nly add itiv e co mbi nati ons and syn erg y for ZE A- FB ₁ , NI V- T2	(Grote n <i>et</i> <i>al.</i> , 199 8)

Mycotoxin association *	Interaction model	Cell model	Study design	Endpoint *	Combined effect	Reference
DON- α -ZOL-FB ₁	Arithmetic definition of additivity	Porcine granulosa cells	Comparison of the effects of 3.4 μ M DON, 9.4 μ M α -ZOL, and 10 μ M FB ₁ to 3.4 +10 μ M DO	Proliferation of granulosa cells, and their steroid (progesterone and estradiol) production	Alleged significant interaction for progesterone production, no significant interaction for	(Cortinovis <i>et al.</i> , 2014)

			N+ FB ₁ and 9.4 +10 μM α- ZO L +F B ₁		cell proliferation and estradiol production	
DON- α- Z O L- FB r- NI V	Isobologram method	Swine whole-blood culture	Analysis of the effects for serial dilutions of the mycotoxins	Cell proliferation	Demonstrated synergism (FB ₁ α-ZO L), no interaction (DON NI V)	(Luongo <i>et al.</i> , 2008)
ZEA- T- 2 to xin	Arithmetic definition of additivity	Monkey kidney Vero cells	Combination of toxins at equimolar concentration	Cytotoxicity MTT ROS production, and expression of Heat shock protein HSP70	Increased toxicity compared to each toxin alone	(Bouaziz <i>et al.</i> , 2013)
ZEA- α-	CI-iso	Human HepG2	Comparison	Cell viability	Demonstration	(Wang <i>et</i>

Z O L	bo lo gr a m m e t h o d	hepatom a cells	of actu al mix ture toxi city data to pre dict ed one s bas ed on the Ma ss acti on law con cept	ty MTT	ted ant ago nis m for low dos es, add itivi ty for me diu m dos es and syn erg y at high dos es	<i>al.</i> , 201 4)
ZEA- α - Z O L- β - Z O L	CI- Is ob ol og ra m m e t h o d	Hamster ovarian cells CHO- K1	Compar ison of actu al mix ture toxi city data to pre dict ed one s bas ed on the Ma ss	Cytotoxici ty MTT	Demon stra ted add itiv e effe cts for ZE A- α - ZO L and ZE A- β - ZO L, ant ago	(Tatay <i>et</i> <i>al.</i> , 201 4)

			action law concept		nis m at high con cen trati on for α - ZO L- β - ZO L and syn erg y at low con cen trati on for ZE A- α - ZO L- β - ZO L	
FB₁-α- Z O L	Isobol og ra m m et hod	Human Jurkat T cells	Analysi s of the effe cts for seri al dilu tion s of the my cot	Lymphocy te prolife ration, cytoki ne (IL-2 and INF- gamm a) expres sion	Demon stra ted syn erg y for lym pho cyt e prol ifer atio	(Luong o <i>et</i> <i>al.</i> , 200 6)

			oxi ns		n, inte ract ive effe ct cyt oki ne exp ress ion	
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*Abbreviations used: DON= deoxynivalenol, FB₁= fumonisin B₁, IL= interleukin 2, IFN= interferon, NIV= nivalenol, MCP-1= monocyte chemoattractant protein-1, MTT= (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) tetrazolium, NIV= nivalenol, ROS= reactive oxygen species, TNF- α = tumor necrosis factor- α , ZEA= zearalenone, α -ZOL= α -zearalenol, β -ZOL= β -zearalenol

Table 10: Interactions involving other mycotoxins from *Fusarium*

Mycotoxin association *	Interaction Model	Cell model	Study design	Endpoint*	Combined effect	Reference
BEA-DON-T-2 toxin	CI-Isobologram method	Chinese hamster ovarian CHO-K1 cells	Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept	Cell viability MTT and NR	Demonstrated antagonism DON-BEA, DON-T2 antagonism, BEA-T2 synergism, DON-BEA-T2 synergism and low dose antagonism	(Ruiz et al., 2011a)
BEA-DON-T-2 toxin	CI-Isobologram method	Monkey kidney Vero cells	Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept	Cell viability NR	Demonstrated antagonism DON-BEA, T2-BEA antagonism, DON-T-2 antagonism, DON-BEA-T-2 antagonism	(Ruiz et al., 2011b)
BEA-DON-T-2 toxin	Arithmetic definition of additivity	Chinese hamster ovary CHO-K1 cells, monkey kidney Vero cells	Comparison of the tested and predicted toxicities for mycotoxin mixtures (simple additive)	Cell viability NR	Potential of interactive effects	(Font et al., 2009)
BEA-DON-T-2 toxin-ZEA-	Arithmetic definition of	Human Colony Forming Unit-	Comparison of the toxic effect of mixtures to	Myelotoxicity	Alleged synergy DON-BEA, antagonism	(Ficheux et al., 2012)

ENN	additivity	Granulocyte and Macrophage (CFU-GM)	the sum of the toxic effects of individual compounds at their concentration in the mixture		DON-FB ₁ , synergy or additivity DON-T-2, additivity DON-ZEA, T2-ZEA, BEA-ENNB,	
BEA-FB ₁	Arithmetic definition of additivity	Turkey peripheral blood mononuclear cells	Comparison of 8 μM FB ₁ , 8 μM BEA, 8+8 μM FB ₁ & BEA	Apoptosis assessed by nuclear DNA fragmentation	Alleged slightly additive effect	(Dombrink - Kurtzman, 2003)
ENN A-A ₁ -B-B ₁	CI-Isobologram method	Caco-2 cells	Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept	Cell viability MTT	Demonstrated synergy for ENNB -- ENNA ₁ , ENNB ₁ -- ENNA ₁ , ENNA -- ENNA ₁ -- ENNB ; antagonism for ENNB -- ENNB ₁ ; additivity for all other combinations	(Prosperini et al., 2014)
ENN A-A ₁ -B-B ₁	CI-Isobologram method	Hamster ovarian cells CHO-K1	Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept	Cell viability MTT	Demonstrated synergistic effect of combined ENs A+A ₁ , A+B, A ₁ +B ₁ , A+A ₁ +B, A+A ₁ +B ₁ , A+B+B ₁ and	(Lu et al., 2013)

					A ₁ +B+B ₁	
ENN B₁-T-2 toxin	CI-Isobologram method	Porcine intestinal IPEC 1 cells and porcine intestinal tissue explants	Comparison of actual mixture toxicity data to predicted ones based on the Mass action law concept	Cytotoxicity and morphological and histopathological scoring	Demonstrated less than additivity with decreasing concentrations of toxins	<u>(Kolf-Clauw et al., 2013)</u>
FA-DON-FB₁	Arithmetic definition of additivity	Pineal cell cultures	Comparison of 1 μM FA, 1 μM DON, 1 μM FB ₁ , 1+1 μM DON+FA, and 1+1 μM FA+FB ₁	Levels of pineal 5-HT and 5-HTP	Alleged possibly synergy or antagonism	<u>(Rimando and Porter, 1999)</u>

*Abbreviations used: BEA= beauvericin, DON= deoxynivalenol, ENN= enniatin, FA= fusaric acid, FB₁= fumonisin B₁, MTT= (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) tetrazolium, NR= neutral red, 5HT= 5-hydroxytryptamine, 5-HTP= 5-hydroxy-l-tryptophan, ZEA= zearalenone