Toxicity Ranking and Toxic Mode of Action Evaluation of Commonly Used Agricultural Adjuvants on the Basis of Bacterial Gene Expression Profiles

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Abstract

The omnipresent group of pesticide adjuvants are often referred to as “inert” ingredients, a rather misleading term since consumers associate this term with “safe”. The upcoming new EU regulation concerning the introduction of plant protection products on the market (EC1107/2009) includes for the first time the demand for information on the possible negative effects of not only the active ingredients but also the used adjuvants. This new regulation requires basic toxicological information that allows decisions on the use/ban or preference of use of available adjuvants. In this study we obtained toxicological relevant information through a multiple endpoint reporter assay for a broad selection of commonly used adjuvants including several solvents (e.g. isophorone) and non-ionic surfactants (e.g. ethoxylated alcohols). The used assay allows the toxicity screening in a mechanistic way, with direct measurement of specific toxicological responses (e.g. oxidative stress, DNA damage, membrane damage and general cell lesions). The results show that the selected solvents are less toxic than the surfactants, suggesting that solvents may have a preference of use, but further research on more compounds is needed to confirm this observation. The gene expression profiles of the selected surfactants reveal that a phenol (ethoxylated tristyrylphenol) and an organosilicone surfactant (ethoxylated trisiloxane) show little or no inductions at EC20 concentrations, making them preferred surfactants for use in different applications. The organosilicone surfactant shows little or no toxicity and good adjuvant properties. However, this study also illustrates possible genotoxicity (induction of the bacterial SOS response) for several surfactants (POEA, AE, tri-EO, EO FA and EO NP) and one solvent (gamma-butyrolactone). Although the number of compounds that were evaluated is rather limited (13), the results show that the used reporter assay is a promising tool to rank commonly used agricultural adjuvants based on toxicity and toxic mode of action data.

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Introduction

Adjuvants are compounds that modify the effects of other compounds without having any direct effects themselves. In most cases they are added to a pesticide formulation to increase the performance of the active ingredients or to make the formulation chemically more stable [1]. Depending on the usage, two different types of adjuvants are distinguished, spray adjuvants and formulation aditives. Spray adjuvants also called tank mix adjuvants are added in the spray tank along with the pesticide(s) just before application on the field. The second type of adjuvants called formulation additives or inert ingredients are part of the pesticide formulation [1,2].

Besides solvents, surfactants and especially non-ionic surfactants make up the largest group of adjuvants, a simplified overview of the most important chemical classes is listed in Figure 1. This large and heterogeneous group of chemicals is used in pesticides, detergents, personal care and many other products. Due to their variety in applications, adjuvants are the chemicals that are produced and consumed in the largest volumes in the world and most of them end up in detectable levels dispersed in different environmental compartments (soil, water, sediment) and in our food chain [3,4].

Nevertheless, there is a lack in current (pesticide) legislation concerning the use and allowable residue levels of adjuvants. Current regulation concerning the placing of plant protection products on the market, Directive 91/414/EEC, does not specifically deal with adjuvants. The upcoming new regulation (EG) 1107/2009 replaces the Directives 79/117/EEG and 91/414/EEG and will apply from June 2011. The new regulation acknowledges the need for more (eco)toxicological information regarding all the components of plant protection products and claims a better protection of human, animal and environmental health by applying the precautionary principle. Adjuvants will make part of future pesticide risk evaluations and a list of forbidden adjuvants for use in crop protection will be constructed...
when more information becomes available. Industry has to take responsibility to demonstrate that substances or products produced and placed on the market do not have any harmful effect on human or animal health or any unacceptable effects on the environment. Next to the legislation concerning the authorisation of pesticides, European regulations list the pesticide Maximum Residue Levels (MRLs) for different food products, but no such levels are set for adjuvants. Although adjuvants occur in large quantities in the environment only two products, nonylphenol and 4-nonylphenol, are listed as priority chemicals in the water framework directive [5]. This lack of regulation exists mainly because the applied adjuvants in a pesticide formulation are protected by industry are not disclosed to the public. Consequently, hardly any information on the toxicity, toxicological mode of action and environmental fate is available for authorities and the public. Furthermore, a lot of adjuvants are mixtures of different compounds and cause a lot of analytical challenges. Only very recently, US EPA considered requiring public disclosure of all ingredients of pesticide formulations [6,7].

Most studies regarding adjuvants focus on the efficacy and only few research papers focus on toxicity and environmental fate. Nevertheless, there is an urgent need for information concerning the toxic mode of action, residue levels and the environmental fate of adjuvants for correct risk assessment and estimation of threshold levels [8]. Information on the toxic mode of action of compounds is important to develop a solid scientific basis for risk assessment [9,10]. The use of appropriate alternative in vitro systems, can provide relevant information to facilitate regulatory decision-making. Moreover, the use of non-animal tests is promoted by the new EU crop protection regulation (1107/2009). The European OSIRIS project (Optimised strategies for risk assessment of industrial chemicals through integration of non-test and test information), proposes that a good way to improve the evaluation of chemicals may be by categorisation in modes of toxic action [11]. In this way, priorities for the evaluation of compounds can be set based on the toxic modes of action like for example the genotoxic potential of a compound. The in vitro assay used in this study is an example of such a test system. The multiple endpoint bacterial reporter assay is based on the induction of specific signalling pathways (oxidative stress, DNA damage, membrane damage and general cell lesions) that are universal in the living cell and hence the assay is able to combine the detection of toxic compounds and at the same time provide information on a number of universal mechanisms of toxicity [12].

In the present study we applied the bacterial multiple endpoint reporter assay to evaluate different adjuvants at the toxicity (growth inhibition) and toxic mode of action level. In a first step, bacterial growth inhibition (IC₅₀, NOEC and LOEC) is quantified and compared between the different adjuvants. Second, new information regarding different mechanisms of toxic action, i.e. DNA damage, oxidative stress, membrane damage and general cell lesions is obtained and these results are applied to categorise the adjuvants according to the mechanisms of toxic action. The toxicological results (acute toxicity and toxic mode of action) of this study are applied to select adjuvants that have a preference of use.

**Material and Methods**

**Selection of compounds**

The different adjuvants were selected based on their high frequency of use in pesticides in Belgium (consumption data 2003). A broad selection was made containing compounds from the major adjuvant categories (Figure 1). To this selection of adjuvants, toxicological model compounds were added, i.e. mitomycin C (MytC) and methyl methane sulphonate (MMS).
for DNA damage, hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}) and paraquat (PQ) as model compounds for oxidative stress, and pentachlorophenol (PCP) and lindane (L) for membrane damage and general cell lesions. The different solvents and model compounds were of analytical quality and obtained from Sigma (Sigma-Aldrich, Bornem, Belgium).

### Bacterial strains

All bacterial strains used, except SfiA, are based on an *Escherichia coli* K-12 derivative SF1 containing the mutations lac\_4169 deleting the entire lac operon, and sfiA. All the different lac\_Z fusions are present as single chromosomal inserts [13]. A selected list of strains from the publication by Orser et al, were used responding to different types of stress like DNA damage, oxidative stress, protein denaturation, membrane damage, osmotic stress, general cellular stress and heavy metal presence (Table 1). The SfiA strain is part of the SOS chromotest derived from *E. coli* GC4436 with a deletion in the lac operon carrying a sfiA::lacZ fusion so that responses to DNA-damaging agents can be measured [14].

### Toxicity evaluation of the selected adjuvants

The growth inhibition test was performed with the *E. coli* ClpB strain. The ClpB strain is a growth inhibition sensitive strain for a broad range of chemicals [12]. Pre-cultures were grown overnight at 37°C and 250 rpm in Luria Bertani (LB) broth medium (Sigma-Aldrich, Bornem, Belgium). Subsequently bacteria were exposed in 96-well plates for 90 minutes at 37°C and 200 rpm (detailed protocol described in [12]). Exposure experiments were carried out in 96-well plates in a linear ½ dilution series containing seven nominal concentrations (Table 2). Six replicates were performed for each exposure experiment and control (received only growth medium) and solvent control (growth medium and pure water) were included. Growth inhibition was calculated as the ratio of exposed versus non-exposed cell yield, expressed by the measured pre- and post-exposure optical density at 600 nm.

The standard to evaluate toxicity of a compound is based on the comparison of LC\textsubscript{50}, EC\textsubscript{50}, or IC\textsubscript{50} values (concentration at which 50% of the test species die (L), are immobile (E) or stop growing (I)) obtained after exposure of the test species to a serial dilution of the selected compounds. This single value is not enough to characterise toxicity if the obtained dose response curves show differences in slopes, as compounds can be equitoxic based on IC\textsubscript{50} values but the dose-response and hence slopes can be different (Figure 2). A supplementary value characterising toxicity at lower concentrations gives additional information, i.e. the NOEC and LOEC (no and lowest observed effect concentration).

For each compound, IC\textsubscript{50} values were calculated using the logistic 4 parameter regression curve (GraphPad Prism). Lowest observed effect concentrations (LOEC) and NOEC at the level of growth inhibition were statistically derived using ANOVA and post hoc Dunnett’s test (p < 0.05).

### Toxic mode of action evaluation of the selected adjuvants

The toxic mode of action of the different selected compounds was evaluated with a bacterial multiple endpoint reporter assay (Table 1). Concentrations for the toxic mode of action studies were based on the results from the growth inhibition experiments, i.e. highest test concentrations chosen were IC\textsubscript{50} values. The bacterial reporter assay was performed as previously described [12,15]. The assay was performed in triplicate in 96 well plates, column 2 till 11 received a uniform amount of the different overnight *Escherichia coli* cultures diluted in Luria Bertani (LB) medium, column one was used as a blank and only received LB. Optical density was measured at 600 nm to check uniformity. After 90 minutes of resuscitation (37°C and 200 rpm) the plates received the compound to be tested at different concentrations, optical density (600 nm) was measured before and after dosing. Columns 5 to 11 received an increasing concentration of the compound in a ½ serial dilution, columns 2 to 4 were negative controls. After 90 minutes of exposure (37°C and 200 rpm) optical density (600 nm) was measured again and the cells were lysed for β-galactosidase measurement. The reduction of ONPG (O-nitrophenyl- β-D-galactopyranoside) (colorless) to ONP (O-nitrophe-

### Table 1. Stress gene promoters fused to the LacZ gene and their functional grouping (modified from Dardenne et al., 2007 and Orser et al., 1995).

| Type of stress response | Promoter | Gene product/Function | Responsive to |
|------------------------|----------|-----------------------|---------------|
| Oxidative stress       | KatG     | Hydrogen peroxidase I | Oxidative stress |
|                        | Zwf      | Glucose-6-phosphate dehydrogenase | Oxidative stress |
|                        | Soi2B    | Superoxide inducible gene | Superoxide radical generating agents |
|                        | Nfo      | Endonuclease IV | Ss and dsDNA breaks, oxidative DNA damage |
| Membrane damage        | MicF     | Antisense RNA to 5’ OmpF | Membrane integrity, osmotic stress |
|                        | OsmY     | Periplasmic Protein | Osmotic stress |
| General cell lesions   | UspA     | Universal stress protein | Growth arrest |
|                        | ClpB     | Proteolytic activation of ClpP | Protein perturbation |
| Heavy metal stress     | MerR     | Regulation of the mercury resistance operon (mer) | Heavy metals |
| DNA Damage             | Nfo      | Endonuclease IV | Ss and ds DNA breaks, oxidative DNA damage |
|                        | RecA     | General recombination and DNA repair | SOS response |
|                        | UmuDC    | DNA repair | Radiation and/or chemically induced DNA damage |
|                        | Ada      | Adaptive response to alkylatation | DNA damage, mainly methyl adducts |
|                        | SfiA     | Inhibitor of cell division | SOS response |
|                        | DinD     | Unknown function within the DNA damage inducible response | DNA damage |

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Toxic Mode of Action of Agricultural Adjuvants

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nol) (yellow) by β-galactosidase was measured spectrophotometrically at 420 nm and was used as a measure for activity of the promoters. Activity of the promoter was calculated taking into account the growth inhibition of the used strain. The results are presented as fold inductions at a given dose $i$, relative to the control values and were calculated through a set of formulas as given below [12]:

$$\text{Activity}_i = \frac{OD_{PE}^{420\text{ nm}} - OD_{SE}^{420\text{ nm}}}{OD_{PD}^{600\text{ nm}} \times 90\text{ min}} \times \frac{OD_{PE}^{600\text{ nm}} - OD_{SE}^{600\text{ nm}}}{90\text{ min}/2}$$

**Formula 1** Activity at a given dose $i$. OD: optical density, PE: Post exposure, SE: start exposure (= post dose), PD: pre-dose.

$$\text{Fold Induction}_i = \frac{\text{Activity}_i}{\text{Average Activity}_{negative\ controls}}$$

**Formula 2** Fold Induction at a given dose $i$.

The presented fold inductions are the mean of three independent replicates. Fold inductions were considered significant when the following criteria were met: (a) presence of a concentration response relationship ($R^2 > 0.5$, significant at $p < 0.05$ for six degrees of freedom) and a positive slope different from 0 ($p < 0.05$) in a linear model, (b) signal statistically significantly higher than the blank (Dunnett’s test $p < 0.05$) [16]. If fold inductions were not significant they were set to 1 to enhance readability of the data and to reduce noise.

### Toxicity and toxic mode of action classification

An initial ranking of the selected adjuvants was made based on the obtained toxicity and toxic mode of action data. Toxicity of compounds was characterised by IC50 and statistically derived NOEC and LOEC values. To characterise the toxic mode of action, the different stress responses were grouped into four major classes (Table 1), heavy metal response was left out. The promoter MerR was not considered for further analysis since it strongly and specifically reacts to specific heavy metal ions i.e. mercury and cadmium, and no such inductions were observed in the dataset.

In classical mortality tests 100% lethality can always be achieved if solubility of the test compound is not a limitation, however this is not the case at the gene expression level. The maximum induction level of a gene is not known and depends on the regulatory mechanism of the gene and the nature of the inducing compound, hence ECx and toxic units have no direct biological meaning in this case. Consequently, a different approach was used to quantify the information at the gene expression level.

### Table 2. Bacterial growth inhibition of adjuvants with used abbreviations throughout the study, concentration range tested (g/L), respective CAS number, IC50 values (concentration at which 50% of the bacteria stopped growing) with confidence intervals (CI), NOEC and LOEC (no and lowest observed effect concentration) at the growth inhibition level.

| SURFACTANTS | abbreviation | concentration range (g/L) | CAS-number | IC50 (CI) g/L | LOEC g/L | NOEC g/L |
|-------------|--------------|---------------------------|------------|---------------|---------|---------|
| Ethoxylated tallow alkyl amine | POEA | 0.010–0.070 | CAS 68478-96-6 | 0.019 (0.018–0.021) | 0.010 | <0.010 |
| Ethoxylated fatty alcohol (AE7) | AE | 0.00156–0.1 | CAS 68002-97-1 | 0.039 (0.029–0.052) | 0.013 | 0.006 |
| Trisiloxaan ethoxy-propoxyten side | Tri EO-PO | 0.0156–1 | CAS 134180-76-0 | 0.082 (0.060–0.11) | 0.031 | 0.016 |
| Ethoxylated phosphate ester (isotridecanol) | Eo PE | 0.078–5 | CAS 9046-01-9 | 0.775 (0.69–0.86) | 0.156 | 0.078 |
| Ethoxylated fatty acid (isotridecanol) | Eo FA | 0.312–20 | CAS 9043-30-5 | 2.02 (1.67–2.45) | 0.531 | <0.531 |
| Trisiloxaan ethoxylate tenside | Tri EO | 0.023–1.5 | CAS 27306-78-1 | >1.5 (p) | 0.468 | 0.234 |
| Ethoxylated tristyrylphenol | Eo TP | 0.14–9 | CAS 99734-09-5 | >0.63 (s) | >0.63 | ≥0.63 |
| Ethoxylated nonylphenol | Eo NP | 0.0078–0.5 | CAS 9016-45-9 | >0.5 (p) | 0.015 | 0.008 |

**SOLVENTS**

| Solvent | abbreviation | concentration range (g/L) | CAS-number | IC50 (CI) g/L | LOEC g/L | NOEC g/L |
|---------|--------------|---------------------------|------------|---------------|---------|---------|
| Isophorone | Is | 0.562–36 | CAS 78-59-1 | 3.98 (3.40–4.618) | 0.563 | <0.562 |
| N-methyl-2-pyrolidone | Pyr | 0.662–42.4 | CAS 872-50-4 | 11.84 (9.32–15.03) | 0.66 | <0.66 |
| γ-butyrolactone | But | 0.6–44 | CAS 96-48-0 | >44 (s) | >44 | ≥44 |
| Dichloromethane | Di | 0.0001–0.0015 | CAS 75-09-2 | >0.0015 (s) | >0.0015 | ≥0.0015 |
| Isopropanol | Isp | 1.2–100 | CAS 67-63-0 | >100 (s) | >100 | ≥100 |

If IC50 could not be calculated, the reason was mentioned (p = precipitation, s = solubility).

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Figure 2. Description of differences in dose-respons curves, IC50 concentrations are equal but LOEC values differ due to differences in slope.

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![Figure 2. Description of differences in dose-respons curves, IC50 concentrations are equal but LOEC values differ due to differences in slope.](https://example.com/figure2.png)
expression level. If fold inductions were significant (criteria see above), the response for each gene was characterized by: 1) the fold induction scores (FIS) at the IC20 level, defined as the ratio of the measured FI to the reference compound FI (set to 100%) and 2) the LOEC at the gene expression level.

Principal component analysis (PCA) was performed using SIMCA-p v11.5 software, (Umetrics AB, Umeå, Sweden) to assess similarities between cases. This multivariate approach allows the visualization of (combination of) mode(s) of action to which the adjuvants belong since the reference compounds were included in the dataset. The FIS dataset was used for PCA analysis, if inductions were not significant, FIS was set to 1, reference gene inductions of the model compounds were set to 100%.

Results

Escherichia coli growth inhibition (Table 2)

Lowest IC50 values are found for ethoxylated tallow alkyl amine (19 mg/L), ethoxylated fatty alcohol (39 mg/L) and trisiloxaan ethoxy-propoxyline tenside (32 mg/L) (Table 2). Due to solubility and precipitation problems, IC50 values cannot be calculated for 3 surfactants (trisiloxaan ethoxy tenside, ethoxylated tristeryl-phenol and ethoxylated nonylphenol) and 3 solvents (gamma-butyrolactone, dichloromethane and isopropanol). Nevertheless, LOEC and NOEC values could be calculated for ethoxylated nonylphenol (15 mg/L and 7.5 mg/L respectively) and trisiloxaan ethoxyline tenside (468 mg/L and 234 mg/L respectively). On the basis of the IC50 values one would conclude that ethoxylated nonylphenol is one of the non-toxic surfactants, but NOEC-LOEC calculations show that already at low concentrations growth inhibition (20%) is observed.

Toxic mode of action

Next to toxicity data for the selected adjuvants more information regarding their toxic mode of action was obtained through a bacterial reporter assay with 14 different toxicologically relevant stress genes. The dose response profile after exposure to ethoxylated nonylphenol (Figure 3A) showed clear concentration responses for 10 stress genes, a detailed figure of the significantly induced genes with standard error is given in Figure 3C. The significantly induced genes belong to different toxic modes of action, oxidative damage (KatG, Zuf, Soi28 and Nfo), DNA damage (RecA, DinD and SoiA), membrane damage (OumF) and cellular stress (OspaB and Uspa). The induced genes show a 3 fold induction at IC20 concentrations for SoiA and Uspa and a 2.5 fold induction for Zuf, DinD and OumF. Compared to the induction profile of ethoxylated nonylphenol the bacterial gene expression profile after exposure to the reference compound paraquat, induced a specific oxidative stress response (Zuf, Soi28, Nfo and SoiA) and the fold inductions are much higher i.e. up to 10-fold inductions (Figure 3B).

As mentioned above for gene expression data the maximum fold induction is not known, hence relative values are used, i.e. fold induction scores (FIS) (Table 3). These values can be compared since they represent gene expression at equitoxic concentrations. The individual fold inductions are given as supporting information (Table S1). To characterise the results of the dose response curves at lower concentrations, LOEC values are calculated (Table 4).

Two important groups of adjuvants are evaluated in this study, solvents and non-ionic surfactants, the results show that in general much lower inductions are found for solvents than for surfactants. The observed LOEC values for the tested solvents are much higher (g/L range) than for surfactants (mg/L range), illustrating that the effect concentrations are much higher for solvents than for surfactants (Table 4).

All tested surfactants, except EO TP, exceeded the 100% level for one or more genes indicating that they provoke higher inductions than the reference compounds. For the selected solvents only Pyr exceeded the 100% level for KatG. It is clear from the FIS at IC20 values that POEA and EA provoke far more stress responses than the other surfactants and solvents. The related LOEC values illustrate that the effects at the gene expression level appear at low concentrations, ranging from 20–80 mg/L for POEA and from 1.6–25 mg/L for EA (Table 4).

The markers for membrane damage are not induced after exposure to the selected solvents. The SOS response related genes RecA, UmuDC and SfiA are induced after exposure to Pyr and But, mild SOS responses for EO FA, tri EO and EO NP and severe SOS responses, RecA and UmuDC inductions, after exposure to POEA and AE (Table 3).

Categorization into toxic mode of action

Compared to the reference compounds which show a principal mode of action, i.e. the reason why they are considered model compounds, the adjuvants show “mixed” toxic modes of action (Table 3, 4 and Figure 4). The toxic mode of action of POEA and EA is complex with inductions of all classes of stress genes making it impossible to assign one or more principal mechanisms of action to these compounds.

Principal component analysis on the FIS dataset illustrates that POEA and AE are grouped separately from all the other compounds and the software labeled them as possible outliers (Figure 4). In the obtained model (R2 = 0.66 ) the first principal component (PC1) explains the majority of the variance (41%) and describes the difference in the SoxRS mediated oxidative stress response on one hand and the OxyR oxidative stress response and membrane damage response on the other hand. The second component (24%) separates DNA damage markers from oxidative damage and membrane damage markers. The data points that are grouped together are isoforin, isopropanol dichloromethane, pentachlorophenol, hydrogen peroxide and ethoxylated tristeryl-phenol, for these compounds the FIS profiles show low inductions. Ethoxylated fatty acid is grouped together with DNA damage inducers MMS and MyrC, mostly because the Ada response is induced, yet the FIS show main inductions for membrane damage related genes.

Discussion

Toxicity and toxic mode of action of adjuvants

Adjuvants comprise of three major groups: surfactants, solvents and synergists and are often referred to as “inert ingredients”. A consumer survey performed by US EPA learned that many consumers are mislead by the term “inert ingredient”, believing it to mean harmless [18]. This certainly is not the case and in fact they can be toxic to humans, may have biological activity of its own [18,19]. Nevertheless, up till recently adjuvants were not taken into account for the risk evaluation of pesticides. The upcoming new EU regulation concerning the placing of plant protection products on the market (EC1107/2009) includes for the first time the demand for information on the possible negative effects of not only the active ingredients but also the used adjuvants. This new regulation requires basic toxicological information that is used to decide on the use, ban or preferential use of available adjuvants [8].

This study provides information on the toxicity and toxic mode of action of the selected compounds. The ranking of the adjuvants based on their toxicity (growth inhibition) showed that the surfactants are far more toxic than the selected solvents in the
assay. Ethoxylated tallow alkyl amine is the most toxic compound tested. High toxicity after exposure to ethoxylated tallow alkyl amine was already reported for several species e.g. tadpoles and green algae [20–23]. Within the group of surfactants toxicity varies by three orders of a magnitude, with ethoxylated fatty acid (isotridecanol) and trisiloxaan ethoxylate tenside as the least toxic compounds. The toxicity results illustrate the importance of reporting toxicity in different ways (here IC50 and NOEC-LOEC) to characterise the toxicity of a compound. If only IC50 values are determined EO NP would be regarded as a non-toxic compound while growth inhibition already occurs at low concentrations. For several compounds IC50 and LOEC values could not be calculated due to limited water solubility. We preferred not to use other solvents than water since in realistic conditions (sprays and tank-mixes) water is used as a diluent or solvent.

Organosilicone surfactants, a fairly new class of non-ionic wetting agents, do not act like classical surfactants through the membranes but they provide a faster penetration of the pesticide in the plant through a specific mode of action i.e. by facilitating stomatal infiltration of solutions [24]. They are considered as promising compounds since improved spreading of the active ingredient can lead to a reduction of the latter in formulations. Two organosilicone surfactants were tested in this study, i.e. trisiloxane ethoxylate tenside (tri EO) and trisiloxane ethoxypropoxylate tenside (tri EO-PO). Both compounds increase the uptake and efficacy of pesticides in a similar way [25], though this study demonstrates that they differ by one order of magnitude at the toxicity level. Stark and Walthall (2003) investigated the acute toxicity of several agricultural adjuvants, including organosilicone surfactants, with Daphnia pulex. They found different LC50 values for different organosilicone surfactants: Silwet L-77® 3 mg/L and Kinetic® 111 mg/L. The results from our study at the gene expression level confirm that the main mode of action of the tested organosilicone surfactants is not through membrane damage (MicF, OsmY and ClpB) since these genes are not significantly induced. The toxic mode of action of organosilicone surfactants is mainly oxidative damage through part of the SoxRS pathway (Zwf and So28). Both compounds are not grouped together with

Figure 3. Bacterial dose response profile after exposure to an adjuvant (ethoxylated nonylphenol) and a reference compound (paraquat). Figure 3a) ethoxylated nonylphenol, and 3b) paraquat. The y-axis denotes the induction at any given dose, the x-axis shows the different stress genes and the z-axis shows the applied concentrations in a 1/2 serial dilution. All data are means of three replicates (n = 3), c) Detailed results for significantly induced genes after exposure to ethoxylated nonylphenol meeting the criteria as mentioned in Material and Methods; bars indicate standard error. *Significantly different from solvent control (one-way ANOVA, Dunnett’s test, p<0.05).

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### Table 3. Significantly induced effects at the gene expression level after exposure to the selected adjuvants.

| Oxidative damage | DNA damage | Membrane damage | General cell lesions |
|------------------|------------|-----------------|---------------------|
| Kat G Zwf Soi 28 Nfo Rec A Umu DC Ada DinD SfiA Mic F Osm Y UspA Clp B | | | |
| **ADJUVANTS** | | | |
| POEA 128 68 281 300 58 106 20 54 - | 240 275 101 75 | | |
| AE 213 90 57 15 56 14 52 - | 253 52 319 231 441 | | |
| Tri EO-PO - - 121 40 - | 136 - - - - | | |
| Eo PE 55 19 - - | 81 - 47 - | | |
| Eo FA - - 67 34 48 - 19 58 - | 169 149 - | | |
| Tri EO - 47 78 - | 136 - - - - | | |
| Eo TP - - - - - | 54 - - - - | | |
| Eo NP 71 33 44 15 35 - | 146 - 72 159 89 | | |
| Is - - - - - | - - - - - | | |
| Pyr 166 21 - 16 - | 83 - | | |
| But - 22 37 16 30 11 12 16 | - - - - | | |
| Di - - | 69 | 56 | |
| Isp | | | |

Results are expressed as fold induction scores (FIS) (%), calculated as the ratio of the measured fold induction (FI) at IC20 level to the reference compound FI at IC20 level. – not significantly induced.
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### Table 4. Statistically derived no observed effect concentrations (NOEC) (g/L) at the level of gene expression for significantly induced genes (ANOVA, post hoc Dunnett’s test p<0.05).

| Oxidative damage | DNA damage | Membrane damage | General cell lesions |
|------------------|------------|-----------------|---------------------|
| Kat G Zwf Soi 28 Nfo Rec A Umu DC Ada DinD SfiA Mic F Osm Y UspA Clp B | | | |
| **ADJUVANTS** | | | |
| POEA 8.0E-05 8.0E-05 8.0E-05 4.0E-05 2.0E-05 2.0E-05 1.6E-04 | 2.0E-05 8.0E-05 2.0E-05 8.0E-05 8.0E-05 8.0E-05 | | |
| AE 3.0E-02 1.6E-03 | 3.1E-03 2.5E-02 2.5E-02 1.6E-03 1.6E-03 1.6E-03 | | |
| Tri EO-PO - | 3.0E-03 2.0E-02 - - - | | |
| Eo PE 6.3E-01 1.3E+00 | 1.6E-01 1.6E-01 - 1.6E-01 1.6E-01 | | |
| Eo FA - | 1.3E+00 1.3E+00 | 1.3E+00 | 1.3E+00 | 1.6E-01 | 1.6E-01 | 1.6E-01 | 3.1E-01 |
| Tri EO - 2.5E-03 4.0E-02 | 2.0E-02 1.3E-03 | | | |
| Eo TP - | 5.6E-01 | | | | | |
| Eo NP 1.6E-02 1.6E-02 6.3E-02 1.3E-01 1.6E-02 | 1.6E-02 2.0E-03 | 8.0E-03 | 1.6E-02 | 1.6E-02 | |
| Is 1.3E+00 | | | | | | | | |
| Pyr 6.3E-01 2.5E+00 | 5.0E+00 5.0E+00 1.3E+00 | 2.5E+00 | | 5.0E+00 | |
| But - | 5.0E+00 2.5E+00 2.5E+00 5.0E+00 1.0E+01 | 1.0E+01 | 1.0E+01 | 1.0E+01 | 5.0E+00 | |
| Di - 1.5E-03 | 1.5E-03 | | | |
| Isp - | 5.0E-01 | | | | | |

**REFERENCE COMPOUNDS**

| MytC 6.3E-04 | | | |
| MMS | | | |
| PQ 1.3E-02 1.6E-03 | 2.5E-02 5.0E-02 | 5.0E-02 | 1.6E-03 | 2.5E-02 | 5.0E-02 | |
| H2O2 1.0E-04 | 2.1E-03 | 2.1E-03 | | | | | | |
| PCP - 1.5E-03 | 1.5E-03 | 2.0E-05 | | | | |
| Li 1.3E-03 2.5E-03 | 2.5E-03 | | 2.5E-03 | | 2.5E-03 | |

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paraquat, the model compound for SoxRS mediated oxidative damage, in the PCA analysis the reason for this is that not the whole SoxRS pathway is induced as can be observed from the FIS dataset.

In a mode of action and QSAR (quantitative structure activity relationships) context, non-ionic surfactants are described as compounds that provoke toxicity through non-specific mechanisms, the toxic potency of these compounds correlates well with their hydrophobicity. Such a mode of action is defined as narcosis, one of the four mode of action categories (narcotics, non-polar narcotics, reactive chemicals and specifically acting reactive chemicals) in the Verhaar classification scheme (Verhaar et al., 2000). Exposure to narcotics typically results in disruption of the biological membrane integrity [26,27]. Several of the non-ionic surfactants included in this study induced membrane damage (MicF and OsmY) and general cell lesions (UspA and ClpB). Narcosis was already described for several of the adjuvants tested, dichlormethane, ethoxylated nonylphenol, ethoxylated alcohol [20,28]. The results in our study confirmed these results for EA and EO NP and also revealed membrane damage after exposure to POEA, EO PE and gamma-butyrolactone. In our study, no membrane damage is found after exposure to dichlormethane, but the test concentrations were low due to the limited solubility.

Membrane damage and general cell lesions were not the only pathways affected after exposure to these compounds, DNA damage and oxidative stress are induced as well. The induced DNA damage markers are part of the SOS response, a well described repair mechanism in bacteria [29]. Valuable markers for the SOS response are RecA, UmuDC and SfiA, they can be considered as indicators for potential genotoxic compounds like the model compound methylmethane sulphonate (MMS) [12–14]. SfiA, is also part of the validated SOS chromotest [14]. The observed DNA damage (both FIS and NOEC) demonstrated that in the reporter assay the SOS response pathway is induced as described in literature, mild SOS response only RecA induction and severe SOS response both RecA and UmuDC inductions [29]. Previous studies already pointed out that the induction of SfiA could be related to oxidative DNA damage [12]. This is also the case in our study since together with the high induction of the oxidative damage markers the induction of SfiA was observed.

Several of the surfactants (POEA, AE, tri-EO, EO FA and EO NP) and one solvent (gamma-butyrolactone) that were tested showed significant inductions for the SOS response pathway. The FIS showed for several compounds inductions of up to 50% of the MMS signal for RecA, indicating that POEA, AE, EO FA and tri EO are half as potent as MMS to induce RecA. These results were observed at mg/L range for POEA, EA and tri EO and in g/L range for EO FA. Environmental concentrations of the selected compounds are not routinely monitored so little data are available, Belanger and colleagues found concentrations of AE (sum of all) in European effluents of 6.8 μg/L, far below the LOEC at the gene expression level, nevertheless further research on the potential genotoxic effects of these compounds is needed as there are no threshold levels for genotoxic compounds [30].

The most recent US-EPA classification of adjuvants lists gamma butyrolactone as harmless and the usage in pesticides is unlimited, though the report lists genotoxic effects at high concentrations [31]. The concentrations that were tested in this study are very high and unlikely to occur in the environment or food chain. Information on possible genotoxic potential of the other tested compounds is not found in literature.

Figure 4. Principal component analysis of FIS (fold induction score) dataset. The first two components (PC1 and PC2) are shown. Individual points represent the gene expression pattern. This plot shows the possible presence of outliers, groups, similarities and other patterns in the data. Observations situated outside the ellipse are outliers. Blue dots: solvents, red dots: surfactants, green dots: reference compounds. doi:10.1371/journal.pone.0024139.g004
Ethyoxylated fatty alcohol is considered as an alternative for the endocrine disruptor ethoxylated nonylphenol which is banned in Europe. Nevertheless, toxicity results in this study show that NOEC-LOEC values are comparable [20]. At the gene expression level, both compounds induce several stress genes, the LOEC at the gene expression level is even lower for ethoxylated fatty alcohol than for ethoxylated nonylphenol. Based on the results from this study other surfactants seem more appropriate to replace ethoxylated nonylphenol i.e. organosilicone surfactants or ethoxylated tristyryl phenol. The results show a first ranking based on toxicity and toxic mode of action of adjuvants, but additional information concerning other relevant endpoints like endocrine disruption potential is needed.

Future perspectives of toxic mode of action studies for ranking of chemicals

Information on environmental concentrations of surfactants is very scarce, moreover for most adjuvants the persistence, bioaccumulation rates and effects in aquatic and terrestrial systems are not known. However, this information is necessary for correct risk assessment. The results from this study provide important information on the effects (toxicity and toxic mode of action) of environmentally important adjuvants. Nevertheless, this study also illustrates that most compounds do not trigger the induction of one specific mode of action, but a combination of several pathways. The interpretation of such results requires expert judgment since the categorization into toxic modes of action is difficult with mixed modes of action, e.g. a compound can be genotoxic and cause membrane damage. In this case the genotoxic properties are more important for the environment and human population, but other combinations of modes of action are possible as well, a compound can provoke narcosis (membrane damage) and have endocrine disrupting potential. Powerful clustering and multivariate statistics are necessary to interpret such complex information and these are important challenges for the use of mechanistic information and categorization into toxic modes of action.

Conclusions

In this study a bacterial multiple endpoint reporter assay with universally stress related endpoints was used to obtain more information on the toxicity and toxic mode of action of several agricultural adjuvants. The results show that the selected solvents are less toxic than the surfactants, suggesting that solvents may have a preference of use, but further research on more compounds is needed to confirm this observation. The gene expression profiles of the selected surfactants reveal that a phenol (ethoxylated tristyrylphenol) and an organosilicone surfactant (ethoxylated tristyryl) show little or no inductions at EC50 concentrations, making them preferred surfactants for use in different applications. The organosilicone surfactant is a fairly new compound that looks very promising, with little or no toxicity and good adjuvant properties.

However, this study also illustrates severe effects at the level of DNA damage with the induction of the bacterial SOS response indicating possible genotoxicity for several of the surfactants (POEA, AE, tri-EO, EO FA and EO NP) and one solvent (gamma-butyrolactone). For several compounds the FIS show inductions of up to 50% of the MMS signal for RecA indicating that POEA, AE, EO FA and tri EO are half as potent as MMS to induce RecA.

Using the information at the gene expression level, we attempted to assign a principal mode of action to the selected adjuvants using multivariate statistics. The principal component analysis revealed that most compounds show a mixed mode of action and AE and POEA show such high inductions for several stress genes that they are allocated as outliers. The technique that was applied shows promising perspectives for the classification of compounds and classification will improve as the dataset expands.

Supporting Information

Table S1 Significant gene inductions after exposure to the selected adjuvants and reference compounds. Results are expressed as fold induction (FI) at IC20 level, non induced genes are set to 1.

Author Contributions

Conceived and designed the experiments: IN PS GH JR RB. Performed the experiments: IN. Analyzed the data: IN. Contributed reagents/materials/analysis tools: IN PS GH. Wrote the paper: IN PS GH JR RB.

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