We investigated the toxicity of soil samples derived from a former municipal landfill site in the South of the Netherlands, where a bioremediation project is running aiming at reusing the site for recreation. Both an organic soil extract and the original soil sample was investigated using the ISO standardized Folsomia soil ecotoxicological testing and gene expression analysis. The 28 day survival/reproduction test revealed that the ecologically more relevant original soil sample was more toxic than the organic soil extract. Microarray analysis showed that the more toxic soil samples induced gene regulatory changes in twice as less genes compared to the soil extract. Consequently gene regulatory changes were highly dependent on sample type, and were to a lesser extent caused by exposure level. An important biological process shared among the two sample types was the detoxification pathway for xenobiotics (biotransformation I, II, and III) suggesting a link between compound type and observed adverse effects. Finally, we were able to retrieve a selected group of genes that show highly significant dose-dependent gene expression and thus were tightly linked with adverse effects on reproduction. Expression of four cytochrome P450 genes showed highest correlation values with reproduction, and maybe promising genetic markers for soil quality. However, a more elaborate set of environmental soil samples is needed to validate the correlation between gene expression induction and adverse phenotypic effects.

Keywords: microarray, Folsomia candida, biotransformation, cytochrome P450

INTRODUCTION

The focus of ecotoxicological research is aimed at understanding toxicological phenomena in a variety of biota (Fent, 2004). Much emphasis is placed on lab-controlled testing of single compounds to address regulatory issues of chemical registration. However, testing the toxicity of complex environmental samples such as fresh water, river sediments, and natural soils remains challenging for several reasons. One of the major problems is that chemical analysis of pollutants in ecosystems often reveals an extensive list of toxicants that are potentially hazardous. Although legislation is based on the concept of concentration thresholds that must not be exceeded to ensure that the site is safe, such analysis cannot provide evidence for the real toxicological consequences of complicated mixtures. One of the valuable tools to assess ecotoxicological consequences of complex environmental mixtures is to apply bioassays. In such assays, survival and reproduction is studied of model organisms (validated in international standard tests), exposed to samples from the environment under controlled conditions. However, identification of the compound(s) causing the adverse effects among the potential list of compounds in a mixture is challenging, due to the fact that the endpoints survival, growth, and reproduction are not specific to the type of stress exerted on the test animals. Traditional bioassays do not allow conclusions on the nature of the chemicals causing the effects.

Transcriptional profiling seems to have important advantages over traditional bioassays. Several recent studies provided evidence that transcriptome profiles bear a signature of the type of pollution (Owen et al., 2008; Nota et al., 2010). If combined with traditional endpoints, genomics analysis of exposed animals can link adverse effects at the organismal level to mechanistic explanation. However, up to now this is only exemplified for single-compound exposures (van Straalen and Roelofs, 2008).

To simulate more realistic ecotoxicological scenarios, some recent studies have investigated toxic effects at the gene regulatory level of compounds presented in binary mixtures. For instance, trinitrotoluene (TNT) mixed with an additional explosive trinitrotiazacyclohexane (RDX) radically altered the gene expression profile of the ecotoxicological model organism Eisenia fetida when compared to single TNT exposure (Gong et al., 2007). While TNT alone regulated 321 genes, the mixture decreased the count to only three genes. These results implied a strong antagonistic effect of RDX on gene expression induced by TNT. In contrast, mixture toxicity studies with compounds proposed to have comparable modes of action should generate comparable transcriptional responses. Indeed, when Daphnia magna was exposed to two polycyclic aromatic hydrocarbons (fluoranthene and pyrene) no clear distinction could be made between the compounds, suggesting similar molecular modes of action (Vandenbrouck et al., 2010).
Furthermore, cluster analysis with both the single compounds and the binary mixture treatments resulted in a separation of treatments based on differences in toxic ratios rather than component differences. However, the results were highly dependent on the composition of the binary mixture. In any case, these lab-controlled experiments suggest that transcriptomics may prove valuable in determining the most toxic substances among complex environmental samples.

Only few studies (restricted to aquatic samples and river sediments) have addressed gene regulatory consequences of exposure to complex environmental samples. Menzel et al. (2009) studied exposure of nematodes to polluted and clean river sediments and showed that several biological processes, such as oxidative phosphorylation, xenobiotics, and development in dependence of exposure to the most polluted samples. This demonstrates that ecotoxicogenomics can be used to distinguish pollution levels in river sediments. To our knowledge, such an approach has not yet been applied to assess soil quality.

In the present study we investigated the toxicity of soil samples derived from a former municipal landfill site in the South of the Netherlands, where a bioremediation project is running aiming at reusing the site for recreation. Very recently, Legler et al. (2011) investigated this complex environmental sample to study the substances that cause toxicity using effect-directed analysis. They identified the presence of compounds (11H-benzo[j]fluorene, 9-methylacridine, 4-azapyrene, and 2-phenylquinoline) with previously unknown teratogenic toxicity in zebrafish. They concluded these compounds may have been missed by current soil chemical quality assessment.

Here we present toxicogenomic data using the soil ecotoxicological model organism Folsomia candida. We asked the question whether an original soil sample exerts comparable toxic responses in our soil ecotoxicological model when compared to the toxic responses exerted upon exposure to the organic extract from that soil (Legler et al., 2011). If this is the case, analysis of extracts will have predictive power to estimate adverse effects in the field (Fent, 2004). To that end, the arthropods were exposed to an organic soil extract and the original Vlagheide soil sample. Results from a 28 day survival/reproduction test revealed differences in toxicity between the organic extract and the ecologically more relevant original soil sample. The more toxic soil samples induced gene regulatory changes in twice as less genes compared to the soil extract. Despite these differences several gene categories (biological processes) were shared among the two samples. In addition, a substantial number of genes were dependent on sample type (soil or extract), potentially explaining the difference in toxicity. Our results show that bioassays deploying functional genomics can reveal crucial information on the nature of the toxicants. Furthermore, we argue that it is essential to include ecologically relevant test organisms in order to properly assess the risk of environmental samples.

**MATERIALS AND METHODS**

**STUDY SITE**

The Vlagheide municipal waste landfill site is located about 10 miles South-East from ’s-Hertogenbosch, the Netherlands. Haskoning B.V. sampled the site in October 2005 at depths varying from 3 to 18 m. In total seven soil samples were taken and pooled, sieved (mesh size 250 μm), homogenized, and freeze-dried to end up as a composite sample of approximately 1 kg dry weight. In order to retrieve an extract the sample was subjected to pressurized liquid extraction an accelerated solvent extraction (ASE) apparatus (Dionex, ASE200, Sunnyvale, CA, USA) with a mixture of acetone and dichloromethane in a 1:3 ratio. The extract was then subjected to gel permeation chromatography (GPC) cleanup with dichloromethane (Legler et al., 2011). An overview of concentrations of persistent organic pollutants such as polychlorinated biphenyls, polycyclic aromatic hydrocarbons, brominated flame retardants, and organochlorine pesticides in the composite sample have been published elsewhere (Legler et al., 2011). Several metals were measured in three samples and were present at variable levels depending on the depth of sampling. Mean concentrations of lead, cadmium, zinc, and copper were 1751, 29, 3060, and 761 mg/kg soil respectively. A full overview of metal measurements and soil parameters is given in datasheet Table S1 in Supplementary Material.

**ECOTOXICITY TEST**

Treatments consisted of two separate dilution series, the first being a series of diluted whole environmental Vlagheide soil, and the second being a series of Lufa 2.2 reference soil spiked with the extract derived from composite Vlagheide soil sample according to Droge et al. (2006) using acetone as solvent.

For the 100% extract treatment, an extract derived from 100 g (d.w.) Vlagheide soil was spiked-in 100 g (d.w.) Lufa 2.2 soil. For the 100% soil treatment, we used sieved and freeze-dried Vlagheide soil. Using Lufa 2.2 soil for dilution, we employed a dilution factor of 2.5 to prepare five additional treatments within each series, resulting in a 40, 16, 6.4, 2.56, and 0% dilution of both undiluted extract and soil treatments. The control (0%) sample was 100% Lufa 2.2 soil in case of the Soil sample dilution series, whereas the control for the Extract dilutions was a solvent control consisting of Lufa 2.2 soil including acetone (the solvent) in an identical amount as was used for the extract dilution series. In this way we were able to normalize for the effect of the solvent during the spike-in procedure of the extract samples (Droge et al., 2006).

Preparation of the test soils and experimental set-up was done following the standard ISO protocol 11267 (ISO, 1999), with four biologically replicated test jars per treatment. The standard ISO test procedure for inhibition of reproduction after 28 days was followed. In parallel, we exposed 10 animals in each of four biological replicate jars per treatment, for 4 days on top of a compressed layer of test soil. These arthropods were snap frozen in liquid nitrogen immediately after exposure so that total RNA could be isolated (see below) for subsequent Microarray and QPCR analysis.

A logistic model was fitted to estimate toxicity end points no-effect concentration (NOEC) and a significant sublethal decrease in reproduction (DiR) for extract samples and soil samples of 50% (further deduced as DiR).

**RNA PREPARATION, AMPLIFICATION, LABELING, AND HYBRIDIZATION**

Samples (Lufa control, NOEC; sublethal effects DiR) were subjected to RNA extraction using the SV Total RNA Isolation System (Promega) according to the manufacturer’s instructions. Agilent
RNA Spike-In Kit (Agilent Technologies) was used to prepare Spike A Mix and Spike B Mix in order to normalize the hybridization measurements. Approximately, 50 ng input of total RNA was used for amplification and labeling with Agilent’s Low Input Quick Amp Labeling Kit. The RNA was reverse transcribed into cDNA and treated with a T7 RNA polymerase to incorporate cyanine 3- or cyanine 5-labeled CTPs in the synthesized cRNA, which was purified with RNeasy (Qiagen) and quality controlled using spectrophotometric measurements on a NanoDrop 2000 (Thermo scientific). Hybridization of 8 × 15 K format microarray slides was performed with 300 ng cyanine 3-labeled cRNA and 300 ng cyanine 5-labeled cRNA according to manufacturer’s protocol (Agilent). The design of the microarray is described by Nota et al. (2009) and details can be found under Gene Expression Omnibus (GEO) platform number GPL7150. A replicate reference design was used (Figure 1) so that each treatment sample was competitively hybridized against a control sample (Lufa 2.2). The design included dye-swapped biological replicates. Two slides of the 8 × 15 K Agilent microarray platform, containing 5069 unique gene probes in triplicate, was used for soil samples and extract samples (Figure 1). Hybridization was performed at 65°C for 17 h rotating at 10 rpm in an incubator. Following the hybridization, the slides were washed using Gene Expression wash buffers and scanned on an Agilent DNA Microarray Scanner. The microarray scan images were preprocessed with Feature Extraction software (version 10.5.1.1.) and the obtained Fold changes were subjected to further statistical analyses. The data was submitted to NCBI’s GEO and can be retrieved under accession number GSE37154.

METHODOLOGY

MICROARRAY DATA ANALYSIS

Statistical analysis of microarray data was performed using the Limma package in R environment (version 2.13.0, Wittenhall and Smyth, 2004). The data were normalized to account for dye bias with the global loess method and the significance of gene expression was verified for each of the soil and extract dataset by a modified t-test using Bayesian statistics two-way analysis of variance (ANOVA) with factors sample type and treatment as main factors. All calculated probabilities were corrected for multiple testing using Benjamini and Hochberg’s false discovery rate procedure at the level of p < 0.05 (Benjamini and Hochberg, 1995). Each probe was assigned a mean log2 expression ratio and an adjusted p-value. Gene annotation was performed in R using a Blast2go script. Subsequently, a GO term Enrichment Analysis was performed by applying the TopGO algorithm on significant gene lists (Alexa et al., 2006; de Boer et al., 2011a) to assess which biological processes, molecular functions, and cellular components were mostly affected. The TIGR MultiExperiment Viewer (TIGR Mev version 4.6.2; Saeed et al., 2006) was used to perform cluster analysis in order to define groups of genes that share common patterns of expression. Hierarchical clustering was done using Euclidean distance and average linkage method. Heat maps were used to represent the data. A general linear model was used to investigate the interaction between factors affecting the variability in the data with the factors Sample type (Extract or Soil), treatment (NOEC or DiR), and the Sample X Treatment interaction according to de Boer et al. (2011a). Finally, a Principal Component Analysis (PCA) in TIGR Mev allowed the identification of factors that most contribute to the variability in the data.

QUANTITATIVE PCR ANALYSIS

Quantitative PCR (QPCR) was performed using a selected group of genes according to de Boer et al. (2011b) on a Biomark HD system (Fluidigm). Information on gene description and PCR primer sequence can be found in Table 2 of the results section. Quantitative analyses of cycle threshold (Ct) values were performed with the software package Genex Light 4.3.5 (Multi ID analysis) according to de Boer et al. (2011b). First, the three technical replicates were averaged over each sample. Efficiency corrections were applied on the mean Ct values using PCR efficiency values previously established and published by de Boer et al. (2011b). Subsequently, gene expression values were assessed relative to an internal reference by normalization with the geometric mean of the housekeeping genes SDHA and YWAZ (de Boer et al., 2009). Finally, the log2 transformed normalized gene expression values were subjected to statistical analysis in SPSS version 17 (IBM). Linear regression was applied to assess dose-dependency. The residuals were tested for normality using a Kolmogorov–Smirnov test. A one-way ANOVA was applied to test whether expression values were significantly different between sample type (Extract or Soil).

RESULTS

ECOTOXICITY TEST

The effects of the original soil from the landfill and its extract on reproduction of Folsomia candida were assessed in an ISO standardized 28 days toxicity test. Figure 2 shows the dose-response curves resulting from exposure to Soil (blue) and Extract (red) samples. The different shapes of the lines clearly indicate a difference in toxicity between the two kinds of samples. The original soil samples show a clearer dose-dependence and appear to be more toxic compared to the organic extracts.
The NOEC and the 50% DiR were deduced for each sample type and a spike-in concentration of 6.4% was taken as NOEC for both Extract and Soil exposures. Furthermore, DiR was observed at 100% spike-in Extract, whereas 16% of spiked-in Soil concentration did not significantly deviate from the 50% DiR estimated by the logistic model for Soil toxicity and was thus taken for further investigations at the molecular level.

**MICROARRAY ANALYSIS**

**Figure 3** shows the number of genes that were differentially expressed in response to toxic exposure in Soil and Extract samples. Regarding the Soil exposure, 109 genes were significantly regulated in response to both NOEC and DiR concentration, of which 76 were up-regulated and 33 were down-regulated. Moreover, 354 genes were only regulated at the NOEC level and 521 genes were only regulated at the DiR level. Intriguingly, exposure to Extracts caused differential expression of an increased amount of genes as compared to Soil exposure. As much as 1581 genes were significantly regulated in response to both NOEC and DiR, of which 747 were up-regulated and 834 down-regulated. Moreover, 613 genes were only regulated at NOEC level and 536 exclusively regulated at DiR level in the Extracts.

Subsequently, the two-way ANOVA with factors Sample type (Extract or Soil) and Treatment (NOEC or DiR) was applied to identify genes only affected by Sample type or Treatment, and to assess whether genes exerted a Sample X Treatment interaction. In total 1929 were affected only by Sample type (Soil, Extract), while 396 genes were exclusively regulated by Treatment (NOEC, DiR). In addition, 400 genes showed a significant Sample x Treatment interaction. Heat maps of significantly regulated genes for these three factors are represented in Supplementary data.

**Table 1** shows the Gene Ontology terms for biological processes enriched in the significant gene lists for treatment, sample type, and the interaction.

Some of the biological processes that were differentially affected by different treatments (low and high level of exposure) are lipid metabolism, response to nutrient level, response to chemical stimulus. The exposure to contaminated samples had a strong impact on 368 genes in both kinds of samples (soil and extract). Glucosyl glucuronosyl transferases (Fcc00734, biological process “lipid metabolism”) and superoxide dismutase (Fcc01344, biological process “response to nutrient level”), for example, were both affected by the exposure, being differentially expressed in response to different toxic levels.

On the other side, other biological processes were found to be affected mainly by the difference in sample type: 1929 genes showed different responses between soil and extract samples. These genes are referable to biological processes such as RNA processing, biosynthetic processes, translation, and translational elongation. For instance, isopenicillin-N-synthetase (Fcc00057), translation initiation factor (Fcc00057), and translation elongation factor (Fcc00062), and ribosomal proteins (Fcc00498, Fcc00410) were differentially affected by the exposure in soil and extract samples.

**Figure 4** shows an acyclic graphs resulting from the enrichment analysis referring to biological processes affected in the context of the interaction between sample type and level of exposure. The most significant ones are lipid metabolic processes (such as fatty acid metabolism and fatty acid oxidation), cellular biosynthetic processes, organic acid metabolic processes, regulation of body fluid, vascular development, and response to wounding.

Some significant genes were found to respond to the toxic exposure in either kind of sample type, at both high and low levels of exposure. Up-regulation of heat-shock proteins (Fcc05793), ubiquitins (Fcc02887), biotransformation enzymes (Fcc01651, Fcc04073, Fcc05526), and components of the antibiotic biosynthetic pathway (Fcc00057, Fcc00170, Fcc05568) was observed. On the contrary, hedgehog, antimicrobial genes, and molecular chaperones were down-regulated.

A PCA was then performed in order to explore how the two factors affect each other and to identify prevalent expression profiles among samples. **Figure 5** is the output of the PCA and shows that the 58% of the variation in the data can be explained by the two factors sample type and exposure level.

**QUANTITATIVE PCR ANALYSIS**

From the microarray analysis it became apparent that some genes show a dose-dependent regulation in response to both soil...
Table 1 | Gene Ontology (GO) terms for biological processes that are over represented in the lists of significant transcripts and their P-values as obtained using the R package topGO (Alexa et al., 2006).

| GO ID | GO term                                | p Value | # in GO term | # Significant |
|-------|----------------------------------------|---------|--------------|--------------|
| GO:0006629 | Lipid metabolic process                | 1.78e-06 | 173          | 30           |
| GO:0009605 | Response to external stimulus          | 3.80e-07 | 103          | 23           |
| GO:0009991 | Response to extracellular stimulus     | 5.02e-07 | 36           | 13           |
| GO:0031667 | Response to nutrient level             | 9.71e-08 | 32           | 13           |
| GO:0042221 | Response to chemical stimulus          | 0.00052  | 207          | 28           |
| GO:007584  | Response to nutrients                 | 1.57e-06 | 23           | 10           |

| GO:006996 | Organelle organization                | 0.150994 | 292          | 126          |
| GO:0042254 | Ribosome biogenesis                   | 0.000193 | 56           | 36           |
| GO:0006364 | RNA processing                        | 0.000367 | 38           | 26           |
| GO:0009058 | Biosynthetic process                  | 0.002262 | 363          | 170          |
| GO:0009059 | Macromolecule biosynthetic process    | 0.003222 | 180          | 90           |
| GO:006412  | Translation                            | 4.86e-05 | 128          | 73           |
| GO:006414  | Translational elongation              | 6.04e-06 | 28           | 23           |

| GO:0044249  | Cellular biosynthetic process         | 0.001781 | 210          | 24           |
| GO:0019395  | Fatty acid oxidation                  | 0.001658 | 15           | 5            |
| GO:0006629  | Lipid metabolic process               | 2.77e-05 | 173          | 25           |
| GO:0001944  | Vasculature development               | 0.000117 | 26           | 8            |

**TREATMENT**

**SAMPLE TYPE**

**INTERACTION**

| GO ID | GO term                                | p Value | # in GO term | # Significant |
|-------|----------------------------------------|---------|--------------|--------------|
| GO:0009605 | Response to external stimulus          | 3.80e-07 | 103          | 23           |
| GO:0009991 | Response to extracellular stimulus     | 5.02e-07 | 36           | 13           |
| GO:0031667 | Response to nutrient level             | 9.71e-08 | 32           | 13           |
| GO:0042221 | Response to chemical stimulus          | 0.00052  | 207          | 28           |
| GO:007584  | Response to nutrients                 | 1.57e-06 | 23           | 10           |

| GO:006996 | Organelle organization                | 0.150994 | 292          | 126          |
| GO:0042254 | Ribosome biogenesis                   | 0.000193 | 56           | 36           |
| GO:0006364 | RNA processing                        | 0.000367 | 38           | 26           |
| GO:0009058 | Biosynthetic process                  | 0.002262 | 363          | 170          |
| GO:0009059 | Macromolecule biosynthetic process    | 0.003222 | 180          | 90           |
| GO:006412  | Translation                            | 4.86e-05 | 128          | 73           |
| GO:006414  | Translational elongation              | 6.04e-06 | 28           | 23           |

| GO:0044249  | Cellular biosynthetic process         | 0.001781 | 210          | 24           |
| GO:0019395  | Fatty acid oxidation                  | 0.001658 | 15           | 5            |
| GO:0006629  | Lipid metabolic process               | 2.77e-05 | 173          | 25           |
| GO:0001944  | Vasculature development               | 0.000117 | 26           | 8            |

**DISCUSSION**

Here we presented a full ecotoxicogenomics assessment of a complex environmental sample and identified large differences between the unprocessed, ecologically more relevant, raw soil, and an organic total extract. The extract samples showed a clearly different level of toxicity when compared with the original soil sample. As such, extract samples may have only weak predictive power to estimate the actual toxicity status in the field. The soil sample is highly contaminated with organic micropollutants such as polychlorinated biphenyls, organochlorine pesticides, and one brominated flame retardant (Legler et al., 2011). Heavy metals...
Table 2 | QPCR assays and results of statistical analysis.

| Code   | Collembase | Function                     | Forward                        | Reverse                        | Efficiency | Linear regression$ | p Value ANOVA$$ |
|--------|------------|------------------------------|--------------------------------|--------------------------------|------------|---------------------|-----------------|
| PHE1   | Fcc05839   | Alcohol dehydrogenase       | GCGCGTAATCGATGCTAGTGT          | GCCAGGCTTCTTTAATGAAA           | 2.2        | 0.75***             | 0.83***         | 0.001           |
| PHE13  | Fcc06009   | Putative deoxynucleoside kinase | TGTCCGGAGTGGTCAGAAGTGC | CACGGCACCCTGCTAGCATC          | 1.9        | 0.47*               | 0.65***         | 0.008           |
| PHE21  | Fcc05124   | CCCTC-binding factor-like    | TGCACTCACTACCTGCCGAGG         | CTCGTGAGTTCCCTTAGC             | 2.0        | 0.68***             | 0.83***         | 0.235           |
| PHE23  | Fcc04316   | Phosphoserine aminotransferase | GCAGCTGATATACCTCAGG           | GGTGCCGCTTTCATGATTGC           | 2.1        | 0.89***             | 0.94***         | 0.195           |
| CYP5   | Fcc00015   | CYP96                        | TCTGATCATTCTTGGACGGG          | TCATGCTCTTCCACCTCTGTCG         | 1.9        | 0.68***             | 0.58**          | 0.035           |
| CYP11  | Fcc01780   | CYP6N3v2                     | GCTTAAAGAGCGAGCAGA            | GCGATATCAGCTGCTGACTT           | 1.9        | 0.99***             | 0.96***         | 0.235           |
| CYP19  | Fcc00390   | CYP2P3                       | CATTGTCTTACATGGACGAAATCC      | CGAATCATTACATGGTCTG            | 1.9        | 0.93***             | 0.89***         | 0.116           |
| CYP23  | Fcc01651   | CYP6N4v1                     | TCCGACAGCTACTACATCAT          | CGGAAACACAAAGATCCGTC           | 1.9        | 0.89***             | 0.92***         | 0.387           |
| CD49   | Fcc00343   | Halocyst dehalogenase-like hydrolase | TATTATACCCGCACTGGGTCT | AATAATGCGCTCCTGAGGTC           | 2.0        | 0.77***             | 0.56**          | 0.519           |
| PHE5   | Fcc04834   | Crossveinless-2 BMP binding  | AAGACGGTGCCGAGTAGATCATC       | GAGATCAGACCTTCCACCTG           | 2.0        | NS                  | NS              | <0.001          |
| PHE7   | Fcc03839   | LMBR1 domain containing 2   | AAAAAAAGCAGAGACAGCCAGA        | TCCCAAACATCGTCCGTTCA           | 2.0        | NS                  | NS              | <0.001          |
| PHE11  | Fcc06002   | ABC transporter              | TGGTGCGAGAGGTCAGAAGTGC        | CACGGCAATTACCTACCTACCT         | 1.9        | 0.48*               | 0.74**          | <0.001          |
| CFP3   | Fcc06377   | Exportin 7                   | CAGTTGTTGGAATCAGGAAAGC         | CACGGCAATTACCTACCTACCT         | 1.9        | 0.49*               | NS              | <0.001          |
| CFP5   | Fcc06380   | Ubiquitin ligase E3 alpha    | TACCACAGACAGATTCGATTCCAGAG    | CACGGTCAAAACAGAGGACG           | 1.9        | NS                  | NS              | 0.004           |
| CFP13  | Fcc06453   | NADH dehydrogenase subunit 5 | TGGTGCCAGACCCAGAATCCCTA       | TTAATATCGACGGCACCACCAT         | 1.9        | NS                  | 0.47**          | 0.012           |
| CFP27  | Fcc06524   | 60S ribosomal protein L30   | ATCGGTGCGAGTCGTTCCG           | GCCGGAGATCATGATAGG             | 1.9        | 0.78***             | 0.71***         | <0.001          |
| CFP51  | Fcc03128   | No significant hit           | AATCATTGGAATACCTCCCAAATG      | AATAGCGGAAATACCTCAGGCT         | 1.9        | 0.47*               | NS              | 0.02            |
| CD89   | Fcc00086   | Laminin A                    | GAGTTGCCAGAGCAGCCTGTAAT       | GAAATCCTACCTAATCAGGCAGG        | 1.9        | 0.47**             | 0.67***         | 0.001           |

$Linear regression analysis was performed to investigate dose-dependent transcriptional regulation at significance level *p < 0.05, ** p < 0.01, or *** p < 0.001.
$$One way ANOVA was performed to investigate significant differential regulation between extract- or soil exposure.

Collembase, EST numbers from Collembase.org. Efficiency, PCR efficiency resulting from a standard curve analysis.
are also elevated in the soil sample, including lead, copper, cadmium, and zinc. The exact chemical composition of the extract is not known, however due to the nature of the organic extraction procedure, metals are expected to be removed from the extract (Hubert et al., 2000). Moreover, microarray analysis generated important mechanistic information that can explain this discrepancy in toxicological effects. In combination with the QPCR data, we can conclude that developmental processes, fatty acid metabolism, and defense processes are adversely affected depending on which sample type was analyzed. Indeed, most of the genes (1929) were regulated in response to sample type. This is reflected in the PCA graph (Figure 5) where the first principle component explaining 42% of the variance divides the samples into either Extract or Soil. As much as 395 genes showed treatment-specific regulation. Among them are genes (CYP6N4v1) that show a highly significant dose-dependent transcriptional activation, which was confirmed by QPCR analysis. Such genes may proof invaluable as genetic markers for soil pollution assessment.

**ECOLOGICAL RELEVANCE**

The ISO standardized *Folsomia* test resulted in a consistent difference in toxicity between soil samples and organic extracts, with

![Figure 4](image-url)
the former being much more toxic than the latter. This confirms the concern raised in earlier reports that toxicity tests based on extracts may generate uncertain levels of protection due to the fact that they do not address bioavailability of the original sample (Fent, 2004). The reason for this result could lie in differences in the availability of the toxic compounds between the original landfill soil and the natural soil spiked with the organic extracts, or in the loss of some of these substances during the extraction procedure. The extraction method using acetone/dichloromethane does not remove all toxic compounds; therefore it is likely that part of the soil toxicity is due to polar substances such as hydrophilic xenobiotic compounds and/or heavy metals.

**MOLECULAR MECHANISMS**

A considerable difference was found between the two kinds of samples, concerning the number of genes affected by the exposure. This is in accordance with a recently published survey of ecotoxigenomics studies by Van Straalen and Feder (2012), who showed that in most studies expression profiles at sublethal toxic effects of 10% DiR (EC10) and EC50 cluster together, while the main differences can be observed between sample type. Soil samples evoked gene regulatory changes in more than one order of magnitude fewer genes than the extract samples although we aimed to assess the two sample types at similar toxicity levels (NOEC and DiR of around 50% reduced reproduction). Due to the steepness of the logistic model through the soil toxicity data we may have chosen a soil sample that exerts more toxic effects than the DiR in Extract, although DiR in Soil did not significantly deviate from 50% DiR deduced from the logistic model. It is worth to mention that Nota et al. (2009) studied the effects of phenanthrene on *F. candida* and also found a smaller number of differentially expressed genes in response to high toxic concentration. Very recently, we obtained a similar result in a toxicogenomic study assessing stress–responses in *F. candida* exposed to the anti inflammatory drug Diclofenac (Roelofs et al. unpublished data). The explanation for this drop in transcriptional regulation needs further investigation. We speculate that higher toxic levels induce intense detoxification responses in the organisms, thus leaving less energy for other less essential processes, although Timmermans et al. (2009) showed that increased desiccation stress strongly increased the number of up- and down-regulated genes.

Some interesting biological processes were influenced by the exposure to contaminated samples. The GO term “lipid metabolic process” includes the chemical reaction and pathways involving all kinds of lipids. It is the biological process most significantly affected in this study. Within living systems, polar lipids have a fundamental structural role, being the main constituents of biological membranes. Furthermore, apolar lipids act as a reserve of energy. Lipid metabolism is regulated in order to ensure the correct balance between degradation and synthesis of lipids, according to the needs of the cells and of the whole organism. When facing environmental stress, an organism will activate a series of stress–response mechanisms in order to face the new conditions. These processes require energy and this might explain the changes in lipid metabolism suggested by the gene expression pattern. In fact, within this GO term category, we found a series of significant genes linked to lipid metabolism and transport. For example, Enoyl-CoA hydratase was found to be up-regulated in response to toxicant exposure: this enzyme is very efficient in metabolizing fatty acids to produce acetyl CoA and energy (Agnihotri and Liu, 2003), so...
it might be up-regulated in order to sustain the energy-requiring processes. On the other side, fatty acid desaturase, that causes increase of unsaturated bonds in fatty acids of membranes (Los and Murata, 1998), was found to be down-regulated, probably because the double bonds in fatty acids are a target of oxidative stress. Another interesting biological process is “response to wounding,” which includes any process resulting from a stimulus indicating damage of the organism. Within this group we found, for example, hedgehog, a developmentally active transcription factor that plays a vital role during early embryonic stages (Tabata and Kornberg, 1994). The hedgehog signaling pathway is intimately linked to cell growth and differentiation, so this protein could also be involved in the healing response. Vascular development is another significantly affected biological process. Springtails do not have a vascular system; thus it is difficult to translate this biological process to an invertebrate response. In fact, in this category we find hypoxia-inducible factor, a transcription factor that responds to changes in the level of available oxygen and mediates responses to hypoxia (Jiang et al., 1996). Furthermore, matrix metalloproteinase are represented in this category. This family of enzymes hydrolyze components of the extracellular matrix and play a central role in many biological processes, such as embryogenesis, normal tissue remodeling, wounding, etc (Nagase and Woessner, 1999). When investigating the single genes that fall into a GO term, it is interesting to notice that many of them are actually annotated to more than one term, and this in fact can be seen as a reflection of the number of interconnections that exist between different kinds of stress–responses. For example, within the context of lipid metabolism, a series of genes are annotated that are responding to different kinds of stress: glucosyl glucuronosyl transferases (involved in phase II metabolism of xenobiotics), glutathione S transferases, CYP450, catalases, dismutases, and other gene products found, by other authors, to be associated to oxidative stress or exposure to toxicants.

Some significant genes were found to both high and low levels of exposure in either kind of sample, soil and extract. Heat-shock proteins, that are part of the general stress–response and have a chaperone function (Feder and Hofmann, 1999), were always found to be up-regulated. A similar result was observed for ubiquitins, regulatory proteins that are involved in the degradation of damaged or unneeded proteins (Glickman and Ciechanover, 2002). An increase in expression of enzymes responsible for biotransformation and detoxification reactions was also observed in all cases of exposure. Mono-oxygenases such as cytochrome P450 for phase I, conjugation enzymes for phase II and ABC transporters for phase III were all found to be significantly up-regulated in contaminated samples compared to the control ones. These enzymes are responsible for the detoxification of organic
compounds, so their induction is expected in case of exposure to organic xenobiotic substances (Xu et al., 2005). Interestingly, two components of the antibiotic biosynthetic pathway were found to be up-regulated in soil samples: aminoadipyl–cysteinyl–valine synthetase and isopenicillin–N–synthetase. Cathepsins, a family of proteases that break apart other proteins and might play a role in apoptosis, also resulted overexpressed, in the extract samples. Hedgehog was down-regulated following exposure to the environmental samples, so the stress caused by toxic exposure is likely to influence important signaling pathways and therefore the correct development of the organism. Some antimicrobial genes were also found to be down-regulated after exposure, suggesting that polluted might adversely affect the insect immune system and increase its susceptibility to invading pathogens. Finally, down-regulation of genes coding for proteins with a folding function is likely to affect directly or indirectly important cellular structures and functions.

In conclusion the microarray experiment shows that transcriptomics data can add relevant information on the nature of the compounds that cannot be recovered by traditional bioassays. We showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012). In this case, the patterns showed this in a recent study on aged copper contamination in an agricultural field (de boer et al., 2012).

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