Data in Brief

Microarray analysis of *Arabidopsis* under gold exposure to identify putative genes involved in the synthesis of gold nanoparticles (AuNPs)

Devesh Shukla, Sneha Krishnamurthy, Shivendra V. Sahi *

Department of Biology, Western Kentucky University, 1906 College Heights, Bowling Green, KY 42101-1080, USA

**Article info**

**Abstract**

Very little is known about the genes responsible for Au uptake, reduction and detoxification in plants, which indeed essential to understand the complex trait of AuNP biosynthesis. We designed a targeted experiment to elucidate the response of plant at transcriptional level under Au exposure, and a microarray was performed on root tissue treated with AuCl₄⁻ in the absence of nutrient media to record specific gene expression signature. Here, we describe the experimental procedures and data analysis in detail to reproduce the results (available at GEO database under GSE55436) published by Shukla et al. (2014) [1] in the Frontiers in Plant Sciences. The data produced from this study provide significant information of genes which may be used to enhance the AuNP biosynthesis.

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**Keywords:** Au, Arabidopsis, Gene expression, Microarray, AuNPs

**Data format**

Raw data: CEL files, analyzed data: RMALog2_ANOVA Results.csv, normalized data: rma-gene-full.chp

**Sequencer or array type**

Affymetrix Arabidopsis Gene 1.0 ST Array

**Sex**

N/A

**Organism/cell line/tissue**

*Arabidopsis thaliana* Col-0

**Sample source location**

Bowling Green, USA (36°59′W 86°26′N 86°26′W)

**Sample size**

Approximately 80 seeds were germinated hydroponically [2] on polypropylene mesh (pore sice 250 μm, Amazon, USA) settled on square plastic wedges in the magenta light boxes (GA-7) containing 0.5× MS medium for 5 days under a long day photoperiod, 16 h cool white fluorescent light (120 μmol m⁻² s⁻¹)/8 h dark, at 23 °C. Thereafter, seedlings were transferred to 1× MS medium and allowed to grow for 7 days to get sufficient amount of root tissue (Fig. 1a). After 12 days, the seedlings were transferred into magenta boxes containing 10 ppm AuCl₄⁻ (HAuCl₄, Sigma-Aldrich, USA) solution at pH 4.2 and incubated for 12 h. Similarly, parallel experiments were performed in distilled water at pH 4.2 to serve as control. After 12 h, root and shoot tissues were separated using sharp blade and snap frozen in liquid nitrogen and stored in −80 °C until use. To avoid the circadian related effect if any, we performed all operations in the middle of the day. The whole set of experiment repeated three times independently and approximately 240 seedlings were used in each biological replicate.

**Total RNA isolation and quality control**

Frozen samples were transferred into liquid nitrogen filled polystyrene container to prevent thawing of the tissue during handling. Total...
RNA was isolated from root and shoot tissue using RNeasy plant Mini Kit equipped with on column DNase I digestion (Qiagen, USA), following the manufacturer’s instructions. We ensured the grinding of the root tissues as fine as possible in liquid nitrogen using mortar and pestle as it is the critical factor in isolating high quality RNA from plant tissue. Further, the quality of the total RNA was determined by visualizing the nearly 2:1 ratio of 28S:18S ribosomal RNA by running the samples in the 0.8% TBE (Tris-Borate-EDTA) agarose gel (Supplementary Fig. 1). The concentration and purity of the samples were measured at 260 nm and the ratio of 260 nm/280 nm (generally a ratio $\approx 2$, is considered as pure RNA), respectively, using Nanodrop (Wilmington, DE, USA). In addition, quality of the total RNA was also assessed by using 2100 Bioanalyzer (Agilent Technologies) before performing microarray.

**Gene expression data analysis**

A novel whole-transcript expression array namely “Arabidopsis Gene 1.0 ST Array”, was used to carry out the microarray analysis. Advantage with this system is that it uses the entire transcript to measure expression of a particular gene thereby produces a complete, accurate and unbiased gene expression data. Target preparation, hybridization, washing, staining, and scanning were carried out following the standard protocols provided in the manufacturer’s instructions (Affymetrix, USA). Three independent biological experiments were carried out for control and experiment (AuCl$_4^-$ treated), hence total six Affymetrix array chips were used and no technical replicates were performed. GeneChips were scanned using GeneChip Scanner 3000 7G (Affymetrix) and Affymetrix GeneChip Command Console Software (AGCC version 3.2.4), and six CEL files were generated. Microarray experiments were performed at the Microarray Core Facility, University of Kentucky, USA. All six CEL files were normalized in freely available Expression Console Software v. 1.3.1 (Affymetrix) using RMA algorithm and probe set summarization files (CHP) were generated for genes or exons. The normalized CHP files processed using the freely available secondary analysis tool, Transcriptome Analysis Console software v. 1.0 (Affymetrix), to generate the .csv files containing Differentially Expressed Genes (DEGs) or Exons (DEEs) using a combined criterion of greater than two-fold change with ANOVA $p$-value of $<0.05$ (Condition unpaired) and corrected $p$-value (FDR) of $\leq 0.16$ in the analysis.

**Fig. 1.** A representative picture of the 12 days old Arabidopsis seedling grown in magenta boxes hydroponically (a). These seedlings treated with 10 ppm of AuCl$_4^-$ solution or water for 12 h, thereafter total RNA was isolated from root tissues for microarray. Overview of effect of Au exposure on root transcriptome of Arabidopsis (b). Volcano plot shows the significance, and fold change of differentially regulated genes obtained in the present experiment. X axis is the linear fold change calculated for Experiment vs Control; Y axis is $-10 \log_{10}$ $p$-value of the ANOVA $p$-values. The gray TCs (transcript clusters) possess $<2$-fold change in expression and filtered out. The green TCs show downregulation ($>2$-fold) and red TCs show upregulation ($>2$-fold).
To filter out the known transcription factors from DEGs, we submitted TAIR IDs of DEGs to a public Arabidopsis transcription factor database (AtTFDB) using a web based tool available on this link, http://arabidopsis.med.ohio-state.edu/AtTFDB/ [1]. In order to obtain a view of significantly changed biological function, GO terms affiliated to DEGs were analyzed using singular enrichment analysis (SEA, http://bioinfo.cau.edu.cn/agriGO/index.php) [3] as described in Shukla et al. [1]. Validation of microarray data was carried out by studying the gene expression of 12 key genes using the quantitative RT-PCR method [1]. For performing the quantitative RT-PCR, most of the primers were designed from 5′ or 3′ UTR to produce gene specific results. Effect of Au exposure on biochemical pathways was predicted using the Plant MetGenMap (http://bioinfo.bti.cornell.edu/cgi-bin/MetGenMAP/home.cgi) and Plant Metabolic Network (http://pmn.plantcyc.org/overviewsWeb/celOv.shtml) as described in earlier study [1]. To get the idea of the signaling mechanism operating under Au exposure, we identified overrepresented regulatory element in the 1000 bp upstream sequences of the upregulated genes using the web based Motif finder tool: SCOPE (http://genie.dartmouth.edu/scope/) [4] and the TAIR Motif analysis tool (http://www.arabidopsis.org/tools/bulk/motiffinder/index.jsp) as described in earlier study [1]. We summarized the steps involved in this study and prepared a flow diagram as shown in Fig. 2.

**Discussion**

Herein, we described a unique dataset of Au transcriptomics in root tissue of Arabidopsis. This dataset is composed of global gene expression measured by using novel unbiased Affymetrix Array ST 1.0. The transcriptomic data is of high quality as evidenced by the RMA analysis of the CEL files. The present dataset has been recently used in a study published in the peer reviewed high impact journal [1], signifying the importance of the data in the context of AuNP synthesis.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.gdata.2014.12.001.

**Conflict of interest**

The authors declare no conflict of interest.

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