Extreme sensitivity of gene expression in human SH-SY5Y neurocytes to ultra-low doses of Gelsemium sempervirens

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Abstract

Background: Gelsemium sempervirens L. (Gelsemium s.) is a traditional medicinal plant, employed as an anxiolytic at ultra-low doses and animal models recently confirmed this activity. However the mechanisms by which it might operate on the nervous system are largely unknown. This work investigates the gene expression of a human neurocyte cell line treated with increasing dilutions of Gelsemium s. extract.

Methods: Starting from the crude extract, six 100 × (centesimal, c) dilutions of Gelsemium s. (2c, 3c, 4c, 5c, 9c and 30c) were prepared according to the French homeopathic pharmacopoeia. Human SH-SY5Y neuroblastoma cells were exposed for 24 h to test dilutions, and their transcriptome compared by microarray to that of cells treated with control vehicle solutions.

Results: Exposure to the Gelsemium s. 2c dilution (the highest dose employed, corresponding to a gelsemine concentration of $6.5 \times 10^{-9}$ M) significantly changed the expression of 56 genes, of which 49 were down-regulated and 7 were overexpressed. Several of the down-regulated genes belonged to G-protein coupled receptor signaling pathways, calcium homeostasis, inflammatory response and neuropeptide receptors. Fisher exact test, applied to the group of 49 genes down-regulated by Gelsemium s. 2c, showed that the direction of effects was significantly maintained across the treatment with high homeopathic dilutions, even though the size of the differences was distributed in a small range.

Conclusions: The study shows that Gelsemium s., a medicinal plant used in traditional remedies and homeopathy, modulates a series of genes involved in neuronal function. A small, but statistically significant, response was detected even to very low doses/high dilutions (up to 30c), indicating that the human neurocyte genome is extremely sensitive to this regulation.

Background

Gelsemium sempervirens (Gelsemium s.), also called yellow jasmine, is a plant belonging to the Loganiaceae family. All parts of the plant contain the major active principle gelsemine as well as other toxic strychnine-related alkaloids, such as gelseminine and sempervirine [1-3]. In the phytotherapy literature, Gelsemium s. has been reported to show sedative, analgesic and anti-seizure properties [4,5] while in the homeopathic Materia Medica and literature, Gelsemium s. is described as a remedy for a variety of anxiety-like psychological and behavioral symptoms [6-9]. The anxiolytic, antidepressant and/or analgesic action of Gelsemium s. extracts and its purified components has been recently demonstrated in animal models [10-16]. Other reports in the literature suggest this plant species may exhibit anticancer and immune-modulating activity [17-20].

The question of dosage is obviously central to pharmacology and of particular interest in homeopathic pharmacopoeia, where the procedure of serial dilutions followed by shaking has sparked much debate. The original extract (Mother Tincture, MT) is generally obtained by grinding the medicinal plant matter with a mortar and pestle and dissolving it in ethanolic solution. According to the most widely-used French pharmacopoeia, the first centesimal (1c) dilution is obtained by dissolving
one volume of MT in 99 volumes of 30% ethanol in water and then subjecting it to vigorous shaking (succussion or "dynamization"). Subsequent c dilutions are prepared by repeating the same procedure. Although the lower dilutions (i.e., 2c to 5c) contain substantial amount of the original active phytochemical substances, their concentration progressively decreases as the number of dilutions increases. Thus, in order to address possible mechanisms of action of high dilutions, physical or chemical mechanisms involving changes imparted to the solvent itself have been hypothesized [21-23]. This is a fairly controversial question in the literature on *Gelsemium s.*, since most authors have investigated only a narrow range of doses or dilutions. It is also important, when dealing with elusive phenomena such as biological responses to diluted and dynamized substances, to take special care with the controls: the recent consensus recommendation among researchers in this field is for protocols that use the diluted and succussed vehicle solution as a control, however this is still a debated theme and has been done only in few cases [24,25].

Previous investigations in our laboratory [26,27] have shown a significant anxiolytic-like activity of *Gelsemium s.* high dilutions (namely 5c, 7c, 9c and 30c according to different test paradigms) in mice, using emotional response models. Other laboratories have also reported in vivo [16,19,28,29] or in vitro [30] effects of *Gelsemium s.* in extremely low doses or high dilution/dynamization, but its action at the cellular level has not been fully clarified. To follow up the above evidence of an anxiolytic effect in animal models, we decided to investigate the *Gelsemium s.* mechanism of action in neuronal models by assessing the drug effects on whole genome expression changes. The SH-SY5Y and IMR-32 human neuroblastoma cells were used since are widely employed in neuropharmacology [31-33]. Finally, this approach allowed us to test several replications of multiple doses and dilutions of the remedy, taking advantage of high-throughput and easily reproducible microarray technology.

Cells were treated with a wide variety of doses: in total, we tested 6 increasing dilutions - which was the maximum sample size permitted by technical constraints - from the low dilution 2c (dilution factor 104) to the extremely high dilution 30c (dilution factor 1060). The 5c, 9c, 30c dilutions are among the most frequently used drug formulations in complementary therapies on humans [34]. The drug effects were compared with those of the same solvent used for the dilutions of *Gelsemium s.*, just without the plant extract (control solutions). After testing for possible toxic effects of any dilution on cell viability, their effectiveness in changing gene expression was evaluated using a microarray designed for the whole human transcriptome. *Gelsemium s.* 2c was checked in SH-SY5Y and IMR-32 cells and the most responsive cell line was chosen for testing also higher dilutions/dynamizations.

**Methods**

**Preparation of Gelsemium s. and control solutions**

The homeopathic dilutions/dynamizations were prepared in a manner comparable to methods used by commercial manufacturers, i.e. using 30% ethanol for all dilution/succussion steps. Since ethanol at higher concentration may be toxic for cells the 100x, last dilution/succussion was made in pure water. The detailed procedure was carefully repeated in all experiments and precisely reported below, since it is relevant as basic science research on homeopathic medicine progresses. Whole hydroalcoholic extract (MT) of *Gelsemium s.* was produced by Boiron Laboratoires, Lyon (F) according to the French Homeopathic Pharmacopoeia [35]. The gelsemine content in the MT was 6.5 × 10−4 M. MT was diluted 100 times in 30% ethanol/distilled water to obtain the 1c dilution. Subsequent serial 100× dilutions up to 29c, each followed by vigorous succussion (shaking) were then prepared in the same solvent using glass bottles.

30-ml bottles containing 1c, 2c, 3c, 4c, 8c and 29c dilutions were supplied by the manufacturer wrapped in aluminum foil and stored in the dark at room temperature in a metal cupboard. The control solutions (solvent) were prepared as the drug dilutions just without the plant extract. The 1c, 2c, 3c, 4c, 8c and 29c solvent samples contain only 30% ethanol/distilled water, but differ for the number of succussions performed. To prepare the final dilutions used in the tests, immediately before the experiments, 0.05 ml of the solutions (*Gelsemium s.* and controls) were added to 4.95 ml of distilled sterile-filtered water (Sigma-Aldrich) in a sterile 15 ml Falcon polystyrene plastic tube and shaken in a DinaA mechanical shaker for 7.5 sec (150 strokes). This yielded the 2c, 3c, 4c, 5c, 9c and 30c succussed dilutions, with ethanol concentration lowered to 0.3% (v/v) (final 0.03% in the assay system).

UV-visible absorption spectra of *Gelsemium s.* samples were performed with a Jasco V550 double-beam spectrophotometer using quartz cuvettes with 1-cm optical path and control solutions as the reference samples.

**Exposure of cells to Gelsemium s. and control dilutions**

Human neuroblastoma cell line SH-SY5Y [36,37], kindly provided by prof. Ubaldo Armato (Department of Life and Reproduction Sciences, University of Verona), was grown in DMEM-F12 (1:1) medium (Lonza, Walkersville, MD, USA), supplemented with 10% foetal bovine serum (FBS; Lonza), penicillin (100 units ml−1) and streptomycin (100 mg ml−1) (Lonza). The culture medium was replaced every three days. The cells were grown in Greiner plastic culture flasks at 37°C in a 5% CO2 atmosphere, until 80% confluence was reached. Cells were propagated after
reactivation of cryogenates until the fourth culture passage and then used for the gene expression assay. Cells were counted in duplicate in a Thoma counting chamber after staining with Turk blue reagent. For the analysis of differential gene expression, SH-SY5Y cells were plated onto 96-well plates in the DMEM-F12 medium with 10% FBS and 5% CO2 atmosphere (90% humidity) for a further 24 h. Four replicate experiments were carried out under identical conditions. In three experiments, Gelsemium s. c and the respective control were tested on IMR-32 neuroblastoma cell line (CCL-127 purchased from ATCC, Manassas, VA, USA), grown and treated under the same conditions, except that EMEM medium (Lanza) was used instead of DMEM-F12.

Cell viability assay
The cytotoxic action of the Gelsemium s. or ethanol dilutions on SH-SY5Y cells was assessed by the WST-1 assay [38]. In this test, cell viability is reflected by mitochondrial dehydrogenase activity in cleaving tetrazolium salts (WST-1 reagent, Roche Molecular Biochemicals -Mannheim, Germany) to soluble formazan. A total of 20,000 cells per well were seeded in a 96-well microplate in the DMEM-F12 medium with 10% FBS and left to adhere for 16 h. Then the culture medium was replaced with 200 μl of the same medium supplemented with 2% FBS. Drug and control solutions (22 μl) were then added (6 replicates for each condition for each plate) and the plate was incubated at 37°C in a 5% CO2 atmosphere. After 24 h, 1:10 (v/v) pre-warmed WST-1 solution was added to the cells and the plate incubated for 3 h. The absorbance (OD) of the samples was measured using a Victor3 multilabel reader (PerkinElmer) for the measurements. Each dose dilutions on SH-SY5Y cells was assessed by the WST-1 assay [38]. In this test, cell viability is reflected by mitochondrial dehydrogenase activity in cleaving tetrazolium salts (WST-1 reagent, Roche Molecular Biochemicals -Mannheim, Germany) to soluble formazan. A total of 20,000 cells per well were seeded in a 96-well microplate in the DMEM-F12 medium with 10% FBS and left to adhere for 16 h. Then the culture medium was replaced with 200 μl of the same medium supplemented with 2% FBS. Drug and control solutions (22 μl) were then added (6 replicates for each condition for each plate) and the plate was incubated at 37°C in a 5% CO2 atmosphere. After 24 h, 1:10 (v/v) pre-warmed WST-1 solution was added to the cells and the plate incubated for 3 h. The absorbance (OD) of the samples was measured using a Victor3 multilabel reader (PerkinElmer, Shelton, CT, USA) at 450 nm, and cell metabolic activity was evaluated as the difference between OD at 3 h and OD at T0.

Measurement of intracellular Ca2+ concentration
Increase in intracellular Ca2+ was monitored as described [39] with minor modifications. SH-SY5Y cells were inoculated in 96-well black microplates (flat transparent bottom) with a density of 80,000 cells/well and left to adhere for 16 h. The culture medium was removed from the wells, and the cells were washed with warm Hank’s basal saline solution (HBSS, Sigma-Aldrich) with 20 mM Hepes (Sigma-Aldrich) and incubated with loading medium (100 μl/well) at 37°C for 40 min in the dark, with 5% CO2 in a humidified atmosphere. The loading medium was made up of the Ca2+-sensitive dye Fluo-4 AM (4.5 μM) (Invitrogen, Paisley, UK) and probenecid (2.5 mM) (Invitrogen) in HBSS. After incubation, the cells were washed and incubated with warm HBSS containing 2.5 mM probenecid at 37°C for 30 min in the dark. At the indicated time, carbachol (Sigma-Aldrich) was added at the final concentrations of 1, 5, 10, 20 μM and the plate was transferred to a Victor3 multilabel reader (PerkinElmer) for the measurements. Each dose was measured in triplicate and compared with a blank for about 15 min using the kinetic mode.

RNA isolation and quality controls
Cells exposed to 24 h Gelsemium s. or control solutions were harvested with trypsin-EDTA-PBS treatment (5 mg L−1, Lonza) and counted. Then, total RNA was promptly extracted (from 3.5 × 106 cells) using the Qiagen RNAeasy Mini Kit (Qiagen GmbH, Hilden, Germany) following the manufacturer’s instructions (Animal cells Spin protocol), including genomic DNA elimination step in column. RNA extraction was performed within 20 min from cell detachment. The RNA samples were centrifuged by precipitation with 2.5 volumes of ice-cold absolute ethanol in presence of 0.3 M Na acetate. RNA yield was determined by a NanoDrop 1000 spectrophotometer (Thermo Scientific, Wilmington, DE, USA) and RNA integrity was then evaluated using the 2100 Bioanalyzer (Agilent, Santa Clara, CA, USA).

cDNA synthesis, labelling and microarray hybridisation
Microarray analysis was performed on a 12 × 135 K (i.e. made with 12 sub-arrays and 135,000 probes per sub-array) human NimbleGen microarray chip (Roche NimbleGen, Madison, WI, USA, catalogue no. 05 543 789 001, design 100718_HG18_opt_expr_HX12) containing 45033 genes with 3 probes per target gene. The microarray is based on HG18, Build 36; cDNA synthesis, labelling and hybridization were performed according to manufacturer’s protocols (http://www.nimblegen.com/support/dna-microarray-support.html; see file 05434505001_NG_Expression_U-Guide_v6p0.pdf). Briefly, 10 μg total RNA for each sample was used to synthesize cDNA using a SuperScript double-stranded cDNA synthesis kit (Invitrogen) with oligo(dT) primers for amplification. After further evaluation of integrity and yield with the Bioanalyzer, the cDNA samples were labelled with Cy3 using a NimbleGen One-Color DNA labelling kit (Roche). 4 μg of Cy3-cDNA were hybridized on each subarray for 16 h at 42°C. All 12 samples (6 Gelsemium s. and 6 controls) for each experiment were hybridized in the same chip and processed simultaneously. Sample tracking controls were used to ensure against cross contaminations or erroneous loading in the array. The procedure was repeated for four and three biological replicates with SH-SY5Y and IMR-32 cells, respectively.

The arrays were scanned with a GenePix 4400A scanner (Molecular Devices Corp., Sunnyvale, CA, USA) and...
scanned images (TIFF format) were then imported into the NimbleScan software for grid alignment and expression data analysis. Quality control of the array images was performed on the basis of the parameters reported in the Experimental Metrics Report as indicated by the NimbleGen Software Guide v3.0. The parameters assessed the absence of spatial biases of the fluorescence within each subarray, the homogeneity of the mean signal among the subarrays and the acceptable level of background (empty and random spots) before background correction and intra-array normalization. Gene calls were generated using the Robust Multichip Average (RMA) algorithm as described by Irizarry et al. [39]. Normalization was performed using quantile normalization as described by Bolstad et al. [40]. The data have been deposited in NCBI’s Gene Expression Omnibus [41] and are accessible through GEO Series accession number GSE42236.

Real time quantitative RT-PCR

A qRT-PCR analysis was performed on SH-SY5Y neuroblastoma treated with *Gelsemium s.* 2c or the control 2c, to verify the gene expression profile of APL1, ALPK3, BIRC8, C1ORF167, DD1, EN2, GALR2, GPR25, LST1, OR4X1, OR5C1, KKL4 and TAC4 genes, that were identified by microarray analysis. UPL hydrolysis probes and primers (RealTime ready Assays, Roche) were specifically designed and experimentally validated to match the differentially expressed transcript Id identified by Nimblegen microarray. One µg of RNA previously extracted (Qiagen), quantified spectrophotometrically (Nanodrop) and further DNase treated (Turbo DNA-free kit, Ambion), was reverse transcribed using Transcriptor First Strand Synthesis kit with oligo dT (Roche) and subsequently 250 ng of cdNA were pre-amplified with a pool of primers following the instruction of RealTime Ready cDNA Pre-Amp Master kit (Roche). The pool consisted of the RealTime Ready Assays primers specific for genes listed above diluted 1:10 each in water PCR-grade. One to 20 diluted pre-amplified cdNA was put in qPCR with the gene specific RealTime Ready Assays and with FastStart Universal Probe Master-Rox (Roche). Briefly, the reaction mixture consisted of 10 µl of 2X FastStart Universal Probe Master-Rox, 1 µl of 20X RealTime ready Assay, 1 µl of template cdNA diluted 1:10 and nuclease free water up to 20 µl. Three different technical replicates were analyzed for each cdNA sample in the same assay and β–actin (ACTB gene ID: 60) and β-2 microglobulin (B2M gene ID: 567) were used as housekeeping genes for the normalization. The StepOne Plus Real-time PCR System (Applied Biosystem, USA) was used to monitor the hydrolysis probe signal generated with a standard thermal profile specific for this kind of probe, i.e.10 min of 95°C, followed by 40 cycles of 95°C 15 sec, 1 min of 60°C. The quantification cycle (Cq) was determined by using the log view of the ΔRn amplification plots, normalized by the internal ROX reference dye, whereas the relative fold change (FC) in the expression levels was determined with the ΔΔCq method, taking the mean of the three PCR replicates. Data are presented as Log2 transformation of FC.

Statistics

The experimental model had dose–response setup, including 6 dilutions of *Gelsemium s.* and 6 corresponding controls. The main working variable was the Log2-transformed fluorescence value of microarray analysis of gene expression. Data from 4 independent experiments were considered. Expecting effects to diminish with increasing dilution, we focused to a pair-wise comparison between *Gelsemium s.* dilutions and the vehicle controls instead of an overall comparison analysis. Two consecutive statistical approaches were followed. The first approach analyzed the complete transcriptome dataset and was aimed to select the DEGs that were most significantly affected by treatment at the highest dose; linear model (Limma) was applied to compare the expression values from *Gelsemium s.* 2c treated and the mean of controls (see details below). The second approach analyzed only the expression values of the selected DEGs when treated with highly diluted drugs or the corresponding controls. The main focus was to verify the null hypothesis that the higher *Gelsemium s.* dilutions did not affect the expression of the genes compared to control. For this analysis we used Friedman test as nonparametric ANOVA and Fisher’s exact test (see details below). The tests analyzed the distributions of the fold changes in the down- or up-regulated DEGs and determined whether the direction of effect for the DEGs detected in the 2c concentration was maintained across all other dilutions (3c-30c).

In the first part of analysis, a linear modelling approach and the empirical Bayes statistics as implemented in the Limma package [42] were employed for differential expression analysis. The Limma test was applied to compare *Gelsemium s.* dilutions with controls, or controls between each other. The p-values were adjusted for the False Discovery Rate (FDR) on the 45033 cases using the Benjamini and Hochberg method [43]. No pre-filtering to the dataset (variant-based or minimal expression-based) was applied to avoid a-priori loss of results when studying minimal doses of drug. Log2 fold change was calculated as Log2-transformed fluorescence value of *Gelsemium s.* dilutions minus Log2-transformed fluorescence value of mean of controls. DEGs were selected as significant and interesting for further analysis if their absolute value of Log2 fold change (|log2 fold change|) was higher than 0.5 and the adjusted p-value was <0.05.

In the second part of the analysis, the significant DEGs in 2c treatment were divided in two groups (considered
as gene-sets) according to their direction of change, including down- and up-regulated genes; data referred to the same dilution (from 2c to 30c of both treatments and respective controls) were joined, treating the single gene as a statistical unit and the mean of four replications as the corresponding datum. Statistical significance of the overall differences between expression profiles of gene-sets (down- and up-regulated) in various treatment conditions was calculated by the Friedman multi-sample test using SPSS software, version 17 (SPSS Inc., Chicago, IL, USA). The Friedman test is a nonparametric test for multiple related samples (in our case, the expression level of multiple matched samples from cells treated with 6 *Gelsemium s.* dilutions or 6 control solutions) that checks the null hypothesis that multiple ordinal responses come from the same population. Following a significant result of Friedman test, frequency of down-regulated vs upregulated genes were calculated; |log2 fold change| lower than or equal to 0.05 were considered to be null. The significance of distributions for each dilution was analyzed by the Fisher’s exact test, which calculates the exact probability of getting, only by chance, the observed values or more extreme ones. A randomly selected set of 49 genes, for comparison of frequency of down-regulated vs up-regulated genes with the *Gelsemium s.* specific gene-set, was generated from the whole microarray dataset, using the specific function of the SPSS 17 software.

Gene expression profiles were clustered by the k-means clustering method and Pearson correlation metrics using the MeV 4.8.1 software (http://www.tm4.org/mev.html). The application “Figure of merit” (FOM) was used to set the number of clusters that best fit the dataset variability [44]. The FOM measures the average intra-cluster variance of the observations, estimating the mean error using predictions based on the cluster averages [45]. Gene functional classification and enrichment analysis were performed by DAVID Bioinformatics Resources 6.7 (http://david.abcc.ncifcrf.gov) [46]. Results of viability assay were analyzed by ANOVA and t-test comparing data from each *Gelsemium s.* dilution (G2c, G3c, G4c, G5c, G9c and G30c) with the corresponding controls (n = 12 replicates for each group).

Results

Cell morphology and function

The SH-SY5Y neuroblastoma cells used in the assay exhibited a neuron-like shape with visible axons and junctions when grown in Petri dishes (Figure 1A). To assess the basal neuronal reactivity of these cells, the culture was stimulated with different doses of the acetylcholine-analogue carbachol and the change in concentration of intracellular calcium was measured with the Ca\(^{2+}\)-sensitive probe Fluo4-AM. As shown in Figure 1B, the cells were sensitive to the varying amounts of the neurotransmitter, and intracellular calcium concentration increased in a dose-dependent way.

UV–VIS Spectra of *Gelsemium s.*

Figure 2 shows the absorption spectra of some of the preparations used in this study. The spectrum of the lowest dilution (1c) was considered as marker for the actual presence of plant extract. Spectra of the subsequent dilutions checked the effectiveness of the 100× dilution steps, i.e. verified that a) the lower dilutions (from pharmaceutical factory and prepared in the laboratory) were comparable and b) the provided higher potencies were effectively diluted. The original 1c dilution supplied by the factory (panel A) was characterized by high absorption in UV region near 210 nm and by two absorption shoulders at 280 nm and 330 nm; no absorption in the visible spectrum region above 450 nm was detected. The original 2c dilution (panel B) showed a qualitatively similar spectrum, but with an absorption intensity about 100 times lower than that of 1c, indicating that the dilution was done correctly during the preparation process. The 3c dilution (panel C) has no significant absorption over the background noise level, which is as expected since it was produced by a 100× dilution of 2c. The 2c solution prepared in the laboratory by a 100× dilution of the original 1c in water (panel D) shows a spectrum with absorption features corresponding roughly to 1/100 of the spectrum of (panel A), indicating that the final dilution of the samples for use in cell assays was done correctly. The spectra of higher dilutions were below the detection limit for this technique (data not shown).

Effect of *Gelsemium s.* dilutions on SH-SY5Y viability

To evaluate whether the *Gelsemium s.* dilutions had any toxic effects, the viability of SH-SY5Y cells after exposure to drug dilutions for 24 h was checked by WST-1 spectrophotometric assay. As can be seen in Figure 3, *Gelsemium s.* dilutions did not impair cell viability as compared to controls. No significant differences in cell viability were observed between cells treated with the ethanol control solution 0.03% (v/v) and untreated cells (data not shown).

Gene expression changes induced in SH-SY5Y cells

Global changes in gene expression produced by exposure to low doses or high dilutions of *Gelsemium s.* extracts in human SH-SY5Y neuroblastoma cells were investigated by microarray analysis, and the results selectively compared with the gene expression of cultures exposed to the vehicle solutions. Cells were incubated for 24 h with the 6 dilutions of *Gelsemium s.* or the corresponding controls, after which the 12 samples were rapidly processed and...
tested simultaneously on a 12 × 135 K NimbleGen chip. After running a total of 4 experiments, differential gene expression was analyzed. The general correlation values among the 48 normalized sub-arrays compared in the analysis (12 conditions and 4 replicates) was very high (Pearson correlation coefficient >97%, mean = 0.988), demonstrating the reproducibility of the experiments.

Preliminary Limma analysis, performed only with the control samples, excluded the presence of significant differences among the different diluted/succussed vehicles (adjusted p > 0.05), and authorized merging them in a unique control group. The difference in expression (Log2 fold change) between Gelsemium s. and the average of the controls was calculated for each dosage, and the results were compared to detect any trend in the response to increasing drug dilutions. In general, the range of changes in gene expression was quite narrow: out of a total of 45033 transcripts, in Gelsemium s. 2c and 3c,
the lowest dilutions, only a small subset of genes (577 and 165, respectively) showed |log2 fold change| > 0.5. Among these DEGs, exposure to Gelsemium s. 2c promoted a statistically significant down-expression of 49 genes, while 7 genes were overexpressed (Limma analysis with adjusted \( p < 0.05 \)) (Table 1). In general, mean fold changes in the mRNAs levels of cells treated with Gelsemium s. were small and only 4 genes showed |log2 fold change| > 0.8. No significant changes of housekeeping genes were recorded, as expected.

**Gene expression changes induced in IMR-32 cells**

To verify whether the effect of Gelsemium s. could be reproducible in different types of neurocytes, the Gelsemium s. 2c treatment and the corresponding ethanol controls were applied to the IMR-32 human neuroblastoma cell line. After three replicate experiments, the analysis did not detect significant changes between Gelsemium s. and controls if the cut-off values of |log2 fold change| > 0.5 and adjusted \( p < 0.05 \) (Limma analysis with Benjamini and Hochberg correction) were applied. As observed in the SH-SY5Y cells, the global gene expression change in the IMR-32 cells was slight, since only 116 genes (0.25% of the transcripts) registered |log2 fold change| >0.5 (compared to the 577 genes in SH-SY5Y cells, corresponding to 1.3% of total). In any case, as shown in Figure 4, the changes in the 56 selected genes of SH-SY5Y cells were in the same direction in the IMR32 cells. In fact, 44 of the 49 genes that were down-regulated in SH-SY5Y also had a negative fold change in IMR-32, and 6 of the 7 genes that were up-regulated in SH-SY5Y also had a positive fold change in IMR-32. These data show that the expression of the same gene-set was also modified in a second type of neurocyte, although the most sensitive model for detecting the effect of Gelsemium s. is the SH-SY5Y cell line.

**Real time quantitative PCR: Validation of the microarray data**

To validate the microarray results, RT-qPCR analysis was performed on SH-SY5Y cells exposed to the Gelsemium s. 2c and the corresponding control. RT-PCR was carried out on the cDNA obtained from the RNA samples of 3 replicate experiments tested by microarray assay. Among the list of DEGs (see Table 1) we selected 13 genes according to the extent of expression changes and their potential relevant functions (e.g. transcription factors, G-protein coupled receptors or neuropeptides) (Table 2). The genes investigated by quantitative PCR generally confirmed the changes obtained by microarray assay. DDI1, EN2, GALR2, GPR25, OR5C1, Klkbl4 and TAC4 genes were down-regulated in Gelsemium s. 2c samples compared to Control 2c in the three replicated experiments. Negative fold changes were observed also with BIRC8 genes, although with variable values. The applied RT-qPCR assays could not detect AIPL1, C1orf167, LST1 and OR4X1, because their expression was under the sensitivity of the assay, and did not confirm the up-regulation of ALK3 gene.

**Statistical analysis of data from Gelsemium s. dilutions and controls**

SH-SY5Y cells treated with higher Gelsemium s. dilutions (3c, 4c, 5c, 9c, 30c) showed changes in gene expression due to treatment, which were rated by Limma
| Gene ID | Transcript ID | Symbol | Log2 fold change | p     | Description                                      |
|--------|---------------|--------|-----------------|-------|--------------------------------------------------|
| 7940   | AF0500424     | LST1   | −0.84 ± 0.14    | 0.04  | Leukocyte specific transcript 1                  |
| 390113 | NM_001004726  | OR4X1  | −0.83 ± 0.06    | 0.01  | Olfactory receptor, family 4, subfamily X, member 1 |
| 23746  | AJ830742      | AIPL1  | −0.82 ± 0.16    | 0.04  | Aryl hydrocarbon receptor interacting protein-like 1 |
| 284498 | AL833920      | C1orf167| −0.80 ± 0.17    | 0.05  | Chromosome 1 open reading frame 167              |
| 221191 | AK058068      | Klk4   | −0.79 ± 0.12    | 0.04  | Plasma kallkrein-like protein 4                  |
| 26658  | NM_012377     | OR7C2  | −0.77 ± 0.07    | 0.01  | Olfactory receptor, family 7, subfamily C, member 2 |
| 112401 | BC039318      | BIRC8  | −0.76 ± 0.11    | 0.00  | Baculoviral IAP repeat-containing 8               |
| 2848   | NM_005298     | GPR25  | −0.75 ± 0.15    | 0.02  | G protein-coupled receptor 25                    |
| 55803  | NM_018404     | ADAP2  | −0.74 ± 0.11    | 0.02  | ArfGAP with dual PH domains 2                    |
| 386676 | NM_198690     | KRTAP10-9| −0.73 ± 0.12    | 0.04  | Keratin associated protein 10-9                  |
| 4353   | X04876        | MPO    | −0.72 ± 0.15    | 0.04  | Myeloperoxidase                                  |
| N/A    | AY358413      | N/A    | −0.71 ± 0.18    | 0.02  | Homo sapiens clone DNA59853 trypsin inhibitor    |
| 392391 | NM_001001923  | OR5C1  | −0.71 ± 0.05    | 0.04  | Olfactory receptor, family S, subfamily C, member 1 |
| N/A    | AK094115      | N/A    | −0.70 ± 0.11    | 0.04  | Homo sapiens cDNA FLJ36796 fis, clone ADRLG2006817 |
| 55287  | BC020658      | TMEM40 | −0.70 ± 0.15    | 0.02  | Transmembrane protein 40                        |
| 54209  | NM_018965     | TREM2  | −0.69 ± 0.10    | 0.02  | Triggering receptor expressed on myeloid cells 2 |
| 150365 | AK097834      | RPS5-821D11.2| −0.68 ± 0.17    | 0.02  | Similar to mouse meiosis defective 1 gene        |
| 400934 | NM_207478     | FLJ44385| −0.68 ± 0.09    | 0.04  | FLJ44385 protein                                 |
| 255061 | NM_170685     | TAC4   | −0.67 ± 0.14    | 0.02  | Tachykinin 4 (hemokinin)                         |
| 644065 | XM_931993     | LOC644065| −0.65 ± 0.23    | 0.04  | Hypothetical protein LOC644065                   |
| 1339   | NM_005205     | COX6A2 | −0.64 ± 0.17    | 0.01  | Cytochrome c oxidase subunit Via polypeptide 2   |
| N/A    | AK128093      | N/A    | −0.63 ± 0.09    | 0.04  | Homo sapiens cDNA FLJ46214 fis, clone TESTI4012623 |
| 53841  | AY358368      | CDHR5  | −0.63 ± 0.11    | 0.04  | Mucin-like proteocadherin                         |
| 9332   | NM_004244     | CD163  | −0.63 ± 0.18    | 0.03  | CD163 molecule                                   |
| 441239 | XM_499305     | LOC441239| −0.63 ± 0.22    | 0.05  | Hypothetical gene supported by BC063653          |
| 7164   | NM_001003397  | TPDS2L1| −0.62 ± 0.09    | 0.02  | Tumor protein DS2-like 1                         |
| 11136  | NM_014270     | SLC7A9 | −0.62 ± 0.09    | 0.04  | Solute carrier family 7 member 9                 |
| 389084 | NM_206895     | UNQ830 | −0.62 ± 0.11    | 0.04  | ASCL830                                          |
| 400224 | XM_375090     | FLJ44817| −0.62 ± 0.20    | 0.04  | Similar to pleckstrin homology domain protein (S V327) |
| 647240 | XM_934559     | LOC647240| −0.60 ± 0.06    | 0.00  | Hypothetical protein LOC647240                   |
| 846    | BC104999      | CASR   | −0.59 ± 0.06    | 0.00  | Calcium-sensing receptor                         |
| 116123 | NM_138784     | RP11-45| −0.58 ± 0.09    | 0.04  | Flavin-containing monoxygenase pseudogene        |
| 644280 | XM_497769     | LOC644280| −0.58 ± 0.06    | 0.05  | Hypothetical protein LOC644280                   |
| 57452  | AB032956      | GALNT1 | −0.57 ± 0.17    | 0.05  | Alpha-D-galactosamine N-acetylgalactosaminyltransferase |
| 414301 | NM_001001711  | DD1   | −0.56 ± 0.11    | 0.04  | DD1, DNA-damage inducible 1, homolog 1 (S. cerevisiae) |
| 116535 | BC016964      | MRGPRF| −0.55 ± 0.17    | 0.01  | MAS-related GPR, member F                        |
| 8811   | NM_003857     | GALR2  | −0.55 ± 0.07    | 0.04  | Galanin receptor 2                               |
| 10880  | NM_006686     | ACTL7B | −0.55 ± 0.12    | 0.04  | Actin-like 7B                                    |
| 6368   | NM_145898     | CCL23  | −0.55 ± 0.11    | 0.05  | Chemokine (C-C motif) ligand 23                   |
| 64581  | BC071746      | CLEC7A | −0.54 ± 0.08    | 0.04  | C-type lectin domain family 7, member A          |
| 644003 | XM_927256     | LOC644003| −0.54 ± 0.11    | 0.04  | Similar to Mucin-2 precursor (Intestinal mucin 2) |
| 643514 | XM_931594     | LOC643514| −0.54 ± 0.10    | 0.03  | Hypothetical protein LOC643514                   |
| 374569 | XM_935431     | LOC374569| −0.54 ± 0.07    | 0.04  | Similar to Lysoosphopholipase                    |
| 84504  | BC101635      | NKX6-2 | −0.53 ± 0.13    | 0.03  | NK6 transcription factor related, locus 2 (Drosophila) |
statistics above the 5% of FDR. Inspection of data reported in Table 3, concerning the expression profiles of the 56 DEGs (49 down-regulated and 7 up-regulated by Gelsemium s. 2c), highlights small expression changes (i.e. |log2 fold change| from 0.05 to 0.6) in 52, 48, 39, 36 and 48 genes of cells treated with Gelsemium s. 3c, 4c, 5c, 9c and 30c respectively. In order to analyze the statistical significance of these effects, a further approach was applied to these 56 DEGs. The hypothesis tested was to determine whether treated samples were different from controls or not and, in particular, if the direction of DEGs’ changes detected in the 2c was maintained across all other dilutions rather than randomly distributed.

Table 1 Differentially expressed genes after treatment with Gelsemium s. 2c in SH-SY5Y cells (Continued)

| Gene ID   | Symbol  | Transcript ID | Log2 fold change ± SEM | p value | Description |
|-----------|---------|---------------|------------------------|---------|-------------|
| 732       | NM_000066 | C8B           | −0.53 ± 0.06           | 0.05    | Complement component 8, beta polypeptide |
| 146336    | NM_182510 | FLJ32252      | −0.52 ± 0.03           | 0.01    | Hypothetical protein FLJ32252 |
| 150763    | BC042847  | LOC150763     | −0.51 ± 0.10           | 0.04    | Hypothetical protein LOC150763 |
| 2020      | NM_001427 | EN2           | −0.51 ± 0.08           | 0.04    | Engrailed homolog 2 |
| 646258    | XM_029203 | LOC646258     | −0.51 ± 0.11           | 0.04    | Hypothetical protein LOC646258 |
| 154872    | NM_001024603 | LOC154872 | 0.51 ± 0.10         | 0.03    | Hypothetical LOC154872 |
| 400866    | NM_001001789 | C2orf24   | 0.52 ± 0.12         | 0.05    | Chromosome 21 open reading frame 24 |
| 9457      | NM_020482 | FHL5          | 0.55 ± 0.19           | 0.04    | Four and a half LIM domains 5 |
| 55816     | NM_018431 | DOK5          | 0.56 ± 0.04           | 0.03    | Docking protein 5 |
| 1446      | NM_001890 | CSN151        | 0.57 ± 0.09           | 0.04    | Casein alpha 1 |
| 285600    | AK130941  | KIAA0825      | 0.63 ± 0.06           | 0.01    | KIAA0825 protein |
| 57538     | NM_020778 | ALPK3         | 0.76 ± 0.10           | 0.01    | Alpha-kinase 3 |

The table includes the genes with absolute log2 fold change higher than 0.5. Each gene is described via GeneBank accession number (Gene ID), Gene symbol (Symbol), NimbleGen array transcript designation (Transcript ID). The log2 fold change of expression compared to mean control vehicle-treated cells is displayed as mean ± SEM (n = 4 replicate experiments). Adjusted p value with Benjamini-Hochberg correction, obtained by Limma statistical test.

Figure 4 Differential effect of Gelsemium s. on two cell lines. Fold changes of the 56 selected genes in SH-SYSY (red bars) and IMR32 (blue bars) cells after 24 h treatment with Gelsemium s. 2c.
Table 2 Validation of microarray data of selected genes by RT-qPCR in *Gelsemium* s. 2c versus Control 2c treated samples

| Symbol | Gene ID | Fold change microarray | Fold change RT-PCR |
|--------|---------|------------------------|--------------------|
|        | R1  | R2  | R3  | Mean | SEM  | R1  | R2  | R3  | Mean | SEM  |
| AIPL1  | 23746 | −0.60 | −0.59 | −1.06 | −0.75 | 0.13 |     |     |     |     |     |
| ALPK9  | 57538 | 1.16  | 0.04  | 0.96  | 0.72  | 0.28 | −0.87 | 0.24 | −0.25 | 0.29 | 0.26 |
| BIRC8  | 112401 | −0.79 | −0.77 | −0.91 | −0.82 | 0.04 | 1.64 | −1.12 | −1.43 | −0.30 | 0.80 |
| C10orf167 | 284498 | −0.69 | −0.51 | −1.00 | −0.73 | 0.12 | n.d. | n.d. | n.d. |     |     |
| DDI1   | 414301 | −0.93 | 0.02  | −1.30 | −0.74 | 0.32 | −0.62 | 0.16 | −1.23 | −0.56 | 0.33 |
| EN2    | 2020  | −0.62 | −0.13 | −0.77 | −0.51 | 0.16 | −1.53 | −0.12 | −0.41 | −0.69 | 0.35 |
| GALR2  | 8811  | −0.57 | −0.36 | −0.72 | −0.55 | 0.08 | −0.94 | −0.19 | −0.61 | −0.58 | 0.18 |
| GPR25  | 2848  | −1.15 | 0.20  | −1.08 | −0.68 | 0.36 | −0.74 | 0.60  | −0.11 | −0.08 | 0.32 |
| LST1   | 221191 | −0.71 | −0.76 | −1.18 | −0.88 | 0.12 | n.d. | n.d. | n.d. |     |     |
| OR4X1  | 7940  | −0.96 | 0.05  | −0.55 | −0.74 | 0.10 | n.d. | n.d. | n.d. |     |     |
| OR5C1  | 390113 | −0.84 | −0.33 | −0.79 | −0.66 | 0.13 | −0.95 | −0.51 | −1.00 | −0.82 | 0.13 |
| Kika4  | 392391 | −0.05 | −0.70 | −1.04 | 0.24  | 1.10 | −0.17 | −0.41 | −0.56 | 0.23 |     |
| Tac4   | 255061 | −0.23 | −0.28 | −1.34 | −0.62 | 0.30 | −1.14 | −0.10 | −0.46 | −0.57 | 0.25 |

1Fold change was calculated as Log2 ratio between G2c and Ct2c expression values within the three replicated experiments (R1-R3); 2fold change was calculated as Log2 transformation of 2−ΔΔCq between G2c and Ct2c. N.d., not detectable.

Statistical inference

Expression values (mean of 4 experiments) of the 49 down- and 7 up-regulated genes referred to the same dilution (2c + 30c) of treatments and respective controls were compared. Additional file 1 reports Log2 data of all samples tested in this analysis. Friedman test estimated the overall variance among the samples and showed that the value distributions of the 12 different treatment groups (6 *Gelsemium* s. and 6 controls, n = 49 or n = 7 data for down-regulated and up-regulated genes, respectively) are significantly different (p < 0.0001). For a direct evaluation of the differences between *Gelsemium* s. treatments and the corresponding controls, Figure 5 shows the distribution of the fold changes in the 49 down-regulated genes for all the dilutions tested. Even though the size of the differences was distributed in a small range, the number of genes with negative fold change (Log2 *Gelsemium* s. < Log2 control, blue in Figure 5) was systematically higher than the number of genes with positive fold change (Log2 *Gelsemium* s. > Log2 control, pink bars in Figure 5). In particular, the frequency of down-regulated vs up-regulated genes was 49 vs. 0 (100% vs. 0%) in 2c, as expected, 47 vs. 2 (96% vs. 4%) in 3c, 42 vs. 3 (86% vs. 6%) in 4c, 38 vs. 3 (78% vs. 6%) in 5c, 30 vs. 9 (61% vs. 18%) in 9c, 27 vs. 7 (55% vs. 14%) in 30c. By applying Fisher exact test, the exact probability of the distributions, under the null hypothesis of indifference, was calculated and significant p values resulted for all dilutions (p < 0.001 for 3c, 4c and 5c treatments, p = 0.0035 for 9c and p = 0.004 for 30c). The absence of an equal scattering between the two signs (positive and negative fold changes) suggests that *Gelsemium* s. at high dilutions affects the expression of a significant portion of these genes. This conclusion is reinforced by a separate Fisher exact test carried out on a list of 49 genes randomly selected by the SPSS software from the 45033 transcripts (excluding the 56 DEGs); as reported in Additional file 2, no significantly different distribution of down-regulated or up-regulated genes in this random gene-set was observed with any *Gelsemium* s. dilution. Figure 6 reports the results for the panel of 7 up-regulated genes. Due to the small number of these genes, a distribution of fold changes could not be drawn and the statistical power of analysis was low. By Fisher exact test, a statistically significant prevalence of positive fold changes was observed only in 2c, as expected, while the prevalence of positive fold changes in the other dilutions was not significant.

Cluster analysis

With the aim to describe the trends of gene expression when exposed to higher *Gelsemium* s. dilutions, k-means cluster analysis was applied on the Log2-fold change profiles of the 56 selected genes. The effect of all the tested *Gelsemium* s. dilutions was visualized as a heat map (Figure 7A) and as mean fold changes in each cluster of genes (Figure 7B). This allowed to identify gene subsets with similar expression profiles, and to detect some trends in the changes induced by increasing *Gelsemium* s. dilutions. Most of the genes down-regulated in the 2c-treated samples were also under-expressed in 3c and, to a varying extent, even in higher dilutions. The frequency of genes with negative fold changes was above 65% in all conditions, and in the sample treated with *Gelsemium* s. 30c the number of common genes that were down-regulated in all dilutions was 20 out of 49 (41%). Cluster 1 contains 20 genes whose expression was down-regulated by the 2c
Table 3 Fold changes of the 56 differentially expressed genes and the 4 housekeeping transcripts in cells treated with the 6 *Gelsemium s. dilutions* compared to means of controls

| Transcript ID | Symbol | G 2c | G 3c | G 4c | G 5c | G 9c | G 30c |
|---------------|--------|------|------|------|------|------|-------|
| AB032956      | GALNTL1| −0.57| −0.20| 0.16 | −0.13| 0.02 | 0.09  |
| AF000424      | LST1   | −0.84| −0.18| −0.20| −0.05| −0.14| −0.23 |
| AJ830742      | AIP1   | −0.82| −0.43| −0.25| −0.24| −0.07| −0.08 |
| AK058068      | Klkb4  | −0.79| −0.41| −0.17| 0.11 | −0.15| −0.07 |
| AK094115      | N/A    | −0.70| −0.35| −0.04| −0.01| 0.01 | −0.41 |
| AK097834      | RPS-821D112| −0.68| −0.31| −0.03| −0.21| −0.01| 0.01  |
| AK128093      | N/A    | −0.63| −0.28| −0.15| 0.05 | −0.19| −0.30 |
| ALB33920      | C1orf167| −0.80| −0.60| 0.17 | −0.04| 0.18 | −0.22 |
| AY358368      | CDHR5  | −0.63| −0.23| −0.09| −0.10| −0.19| −0.25 |
| AY358413      | N/A    | −0.71| −0.15| 0.10 | 0.04 | 0.10 | −0.28 |
| BC010964      | MRGPRF | −0.55| −0.08| −0.12| −0.02| 0.06 | 0.07  |
| BC026558      | TMEM40 | −0.70| −0.56| −0.15| −0.15| 0.04 | −0.12 |
| BC039318      | BIRC8  | −0.76| −0.42| −0.10| −0.09| 0.09 | −0.02 |
| BC042847      | LOC150763| −0.51| −0.29| −0.12| 0.01 | −0.03| −0.03 |
| BC071746      | CLEC7A | −0.54| −0.32| −0.12| −0.17| 0.14 | 0.17  |
| BC101635      | NIK6X-2| −0.53| −0.56| −0.12| −0.12| −0.14| −0.12 |
| BC104999      | CASR   | −0.59| −0.17| −0.25| 0.08 | 0.06 | −0.30 |
| NM_000066     | C8B    | −0.53| −0.07| 0.00 | −0.07| −0.03| −0.23 |
| NM_001001711 | DD11   | −0.56| −0.21| −0.27| −0.31| −0.17| −0.26 |
| NM_001001923 | OR5C1  | −0.71| −0.28| −0.22| 0.08 | −0.13| −0.24 |
| NM_001003397 | TPDS2L1| −0.62| −0.31| −0.33| −0.18| 0.04 | −0.31 |
| NM_001004726 | OR4X1  | −0.83| −0.34| 0.07 | 0.03 | −0.11| −0.18 |
| NM_001427     | EN2    | −0.51| −0.33| −0.22| −0.06| 0.02 | −0.13 |
| NM_003857     | GALR2  | −0.55| −0.31| −0.13| 0.02 | −0.01| −0.13 |
| NM_004244     | CD163  | −0.63| −0.30| −0.20| −0.13| 0.04 | −0.25 |
| NM_005205     | CDX6A2 | −0.64| −0.39| −0.38| −0.07| −0.30| −0.17 |
| NM_005298     | GPR25  | −0.75| −0.41| 0.02 | −0.05| −0.02| 0.02  |
| NM_006686     | ACTL7B | −0.55| −0.44| −0.13| −0.02| −0.01| −0.15 |
| NM_012377     | OR7C2  | −0.77| −0.22| −0.03| −0.14| −0.14| −0.03 |
| NM_014270     | SLC7A9 | −0.62| −0.16| −0.20| 0.01 | −0.27| −0.19 |
| NM_018404     | ADA5   | −0.75| −0.40| −0.30| −0.19| 0.04 | 0.03  |
| NM_018965     | TREM2  | −0.69| −0.34| −0.08| −0.14| 0.09 | −0.20 |
| NM_138784     | RP11-4S J16.2| −0.58| −0.29| 0.06 | −0.05| 0.20 | −0.17 |
| NM_145808     | CCL23  | −0.55| 0.03 | −0.09| −0.20| 0.02 | −0.20 |
| NM_170685     | TAC4   | −0.67| −0.30| −0.19| −0.06| −0.13| −0.24 |
| NM_182510     | FLJ32252| −0.52| −0.33| −0.26| −0.10| −0.10| 0.02  |
| NM_198690     | KRTAP10-9| −0.73| −0.29| −0.10| −0.03| −0.28| −0.13 |
| NM_206895     | UNQ830 | −0.62| −0.45| −0.22| −0.25| −0.10| −0.18 |
| NM_207478     | FLJ44385| −0.68| −0.12| −0.10| −0.08| −0.01| −0.26 |
| XM_04876      | MPO    | −0.72| −0.36| −0.20| 0.19 | −0.01| −0.16 |
| XM_375090     | FLJ44817| −0.62| −0.58| −0.16| 0.00 | −0.27| −0.11 |
| XM_497769     | LOC644280| −0.58| −0.19| −0.20| −0.02| −0.07| −0.05 |
| XM_499305     | LOC441239| −0.63| −0.31| −0.21| −0.08| −0.15| −0.24 |
Table 3 Fold changes of the 56 differentially expressed genes and the 4 housekeeping transcripts in cells treated with the 6 Gelsemium s. dilutions compared to means of controls (Continued)

| ID         | Gene        | 2c  | 3c  | 5c  | 7c  | 9c  |
|------------|-------------|-----|-----|-----|-----|-----|
| XM_927256  | LOC644003   | -0.54 | -0.57 | -0.31 | -0.19 | -0.27 | -0.10 |
| XM_929203  | LOC646258   | -0.51 | -0.20 | -0.38 | -0.13 | -0.11 | -0.08 |
| XM_931594  | LOC643514   | -0.54 | -0.23 | -0.11 | -0.01 | -0.21 | -0.16 |
| XM_931993  | LOC644065   | -0.65 | -0.29 | -0.15 | 0.05  | -0.06 | -0.21 |
| XM_934559  | LOC647240   | -0.60 | -0.40 | -0.23 | -0.16 | -0.08 | -0.08 |
| XM_935431  | LOC374569   | -0.54 | -0.31 | -0.14 | -0.12 | -0.13 | -0.10 |
| AK130941   | KIAA0825    | 0.63  | 0.30  | -0.07 | -0.07 | -0.06 | 0.15  |
| NM_001890  | CSN1S1      | 0.57  | -0.04 | 0.03  | 0.30  | -0.07 | 0.11  |
| NM_001024603| LOC154872   | 0.51  | 0.13  | 0.15  | 0.05  | 0.06  | 0.21  |
| NM_001890  | CSN1S1      | 0.57  | -0.04 | 0.03  | 0.30  | -0.07 | 0.11  |
| NM_018431  | DOK5        | 0.56  | 0.04  | 0.03  | 0.01  | -0.02 | 0.31  |
| NM_020482  | FH5         | 0.55  | 0.38  | 0.01  | 0.10  | -0.02 | -0.08 |
| NM_020778  | ALPK3       | 0.76  | 0.45  | 0.23  | 0.16  | 0.03  | 0.19  |
| BC001601   | GAPDH¹      | 0.01  | 0.09  | 0.02  | 0.10  | 0.03  | 0.04  |
| NM_002046  | GAPDH¹      | 0.09  | -0.14 | -0.02 | -0.10 | -0.01 | 0.00  |
| BC009081   | GAPDH¹      | 0.01  | -0.05 | 0.01  | -0.04 | -0.04 | -0.03 |
| NM_001101  | ACTB¹       | -0.04 | -0.05 | -0.05 | 0.05  | 0.02  | 0.00  |

N = 4 experiments. ¹Housekeeping transcripts.

dilution but which were less sensitive to higher dilutions, thus drawing a curve with asymptotic direction. Clusters 2 and 3 group together the genes also down-regulated by the Gelsemium s. high dilutions (but on which 5c or 9c, respectively, had no effect), while cluster 4 includes the genes that were clearly responsive to Gelsemium s. 2c and 3c only. Cluster 5 contains the 7 up-regulated genes. Though significant up-regulation occurred only with 2c, most of those genes showed a similar effect trend in all dilutions.

Functions of the modulated genes

To obtain a functional classification, the 56 genes whose expression changed following exposure to Gelsemium in SH-SY5Y cells were subjected to analysis of the enriched annotation terms associated with the list. Table 4 reports the top enriched biological themes, particularly the GO terms discovered in the gene list by the DAVID software. A total of 28 genes (all down-regulated) from the list were classified into functional-related gene groups, while 17 IDs were unmapped in the DAVID database (see Table 1) because they have unknown functions. The remaining genes (3 up-regulated and 8 down-regulated) have known functions but were not rated as enriched in the list compared to the whole human transcriptome. The main group of functional features includes genes coding for membrane receptors, and in particular involved in G-protein coupled receptor (GPCR) transduction systems (OR4X1, CASR, OR5C1, CCL23, GPR25, GALR2, OR7C2, MRGPRF). Among these receptors, three have specific functions in olfactory transduction, attuned to detecting different types of stimuli including molecular vibrations [47]. The other clusters of genes may have a role in calcium signaling, inflammatory pathways, neuropeptide/receptor systems or as transcription factors. Of particular relevance for neuronal functions is the small but significant down-regulation of the gene TAC4 and GALR2. The first gene codes for the neuropeptide hemokinin-1 an analog of substance-P [48], and the second for the receptor 2 of the neuropeptide galanin. Both are involved in the complex system of psycho-neuro-immune-endocrine axis which correlates the emotional responses with the hormone release and the immune functions [49,50].

Discussion

Natural remedies are increasingly viewed as potentially valuable complements to conventional drugs in integrated treatment strategies for a number of disorders, and many consumers use natural health products alongside prescription medications [51]. Anxiety and depression are among the ailments most frequently reported by patients seeking complementary and alternative medical remedies and/or naturopathic care [9,52,53]. Gelsemium s. is a traditional remedy used in complementary and alternative therapies for treating patients who exhibit neurological complaints such as headache and anxiety-like symptoms [9,52,53], but evidence-based clinical studies are few and with contrasting results [53,54].
Homeopathy is a 200-year-old therapeutic system that uses extremely small doses of various substances to stimulate auto-regulation and self-healing processes [55]. Although some conventional physicians find such notions implausible [56], use of highly diluted drugs from homeopathic pharmacopoeia has recently seen a worldwide revival [57,58] and laboratory investigations are increasing in this field [26,27], but scientific evidence of underlying molecular mechanisms is still lacking. Moreover, the experimental approaches adopted to study

Figure 5 Frequency of fold change values in the down-regulated gene-set after Gelsemium s. treatments. In this analysis the 49 genes whose expression was down-regulated by Gelsemium s. 2c were considered. Mean Log2 fluorescence values from Gelsemium s.-treated samples (Gnc) and those from controls (Ctnc) were obtained from 4 microarray experiments and their difference was considered as fold change attributable to Gelsemium s. effect (see Methods). Absolute fold changes less than or equal to 0.05 were considered null. Blue bars: frequencies of genes with negative fold change (< −0.05); grey bars: frequency of unaffected genes (from −0.05 to 0.05); pink bars: frequencies of genes with positive fold change (> 0.05). Fisher exact p values are reported in each panel except the G2c-Ct2c that are significant by definition.
Figure 6 Number of genes modulated by Gelsemium s. dilutions in the panel of up-regulated genes. In this analysis the 7 genes whose expression was up-regulated by Gelsemium s. 2c were considered. Differences less than or equal to 0.05 were considered null. Blue bars: number of genes with negative fold change (< -0.05); grey bars: number of unaffected genes (from -0.05 to 0.05); pink bars: number of genes with positive fold change (> 0.05). Fisher exact test is not significant in any dilution except in the 2c dilution that is significant by definition.

Figure 7 K-mean clustering of the genes modulated upon exposure to Gelsemium s. dilutions. The expression profile of 56 genes significantly modulated by Gelsemium s. 2c was evaluated also upon exposure to increasing Gelsemium s. (G) dilutions. Fold change was calculated as the difference between Log2 fluorescence values of each Gelsemium s. dilution and the mean Log2 fluorescence of the controls. Data are means of 4 replicate experiments. A. K-mean clusters (KMC) visualized as a colour-coded heat map. The down-regulated genes (green) with similar expression profiles were grouped in 4 clusters and the up-regulated genes (red) in one cluster. B. Centroid graphs of the mean fold change of genes in the 5 clusters obtained in KMC analysis.
these remedies, particularly for highly diluted solutions, have suffered from problems with replicability between different laboratories. It is therefore important for any results in this field to be confirmed and consolidated through further investigations by independent laboratories, using rigorous protocols and statistical evaluations. The expression microarray analysis on whole genome, as other high-throughput technologies assisted by bioinformatics, could provide a strong clue as to the mechanism of action and the biological relevance of ultra-low doses and high dilutions interactions.

This is the first comparative transcriptomics approach to investigate changes in the human neurocytes induced by a natural plant remedy, traditionally used for anxiolytic-like effects. The chief innovation of our experimental design is that it employs a wide range of doses/dilutions. This enabled us to explore changes in gene expression from low dilutions (2c or 3c), where the active substances can still be expected to exert their normal pharmaceutical action, to high dilutions (9c or 30c), where the most controversial principles of high dilution pharmacology come into play. Thus, both conventional and ‘alternative’ pharmacological theories were evaluated and compared in the same investigation. In previous recent trials, *Gelsemium s.* showed anxiolytic-like effects in mouse emotional response models and appeared to work even at the high dilutions 9c and 30c [26,27]. Two other studies have also found that high dilutions of *Gelsemium s.* exert a preventive action against experimental stress (electric shock) in mice [29] and against convulsions provoked by lithium and pilocarpine in rats [5]. Other researchers have reported an anti-anxiety activity of *Gelsemium s.* [12] and of the alkaloids gelsemine, koumine, and gelsevirine [14,16], but have not explored the effect of ultra-low doses and high dilutions/dynamizations.

To follow up on these in vivo studies, we decided to investigate the action of *Gelsemium s.* at the cellular and transcriptional level. We adopted a validated microarray protocol and applied it to a series of replicate experiments designed to test: a) the null hypothesis that the effect of any *Gelsemium s.* dilutions is similar to that of the control vehicle, b) whether any dose-dependence of the putative effects can be demonstrated. As our model system, we chose the SH-SY5Y and IMR-32 neurocyte cell lines because these have been previously employed for investigations of natural compounds [31], neurotrophic factors [32], mood stabilisers [59], and antipsychotics [33]. In our conditions, this line proved to be more responsive to *Gelsemium s.* than IMR-32 and was used to compare the effects of high dilutions. The cells

| Category | Annotation term | P value | Genes | Fold enrichment |
|----------|----------------|---------|-------|-----------------|
| SP_PIR_KEYW | receptor | 8.32E-11 | AIPL1, OR4X1, CASR, OR5C1, GPR25, GALR2, OR7C2, MRGPRF, CLEC7A, TREM2 | 3.68 |
| GOTERM_BP_FAT | GO:0007186 ~ G-protein coupled receptor protein signalling pathway | 6.60E-11 | OR4X1, CASR, OR5C1, CCL23, GPR25, GALR2, OR7C2, MRGPRF, TAC4 | 4.17 |
| GOTERM_BP_FAT | GO:0007166 ~ cell surface receptor linked signal transduction | 0.001 | OR4X1, CASR, OR5C1, CCL23, DOK5, GPR25, GALR2, OR7C2, MRGPRF, CLEC7A, TAC4 | 0.08 |
| GOTERM_BP_FAT | GO:0051606 ~ detection of stimulus | 0.02 | AIPL1, CASR, CLEC7A | 13.23 |
| INTERPRO | IPR017452:GPCR, rhodopsin-like superfamily | 0.009 | OR4X1, OR5C1, GPR25, GALR2, OR7C2, MRGPRF | 4.44 |
| KEGG_PATHWAY | hsa04740:Olfactory transduction | 0.09 | OR4X1, OR5C1, OR7C2 | 5.31 |
| GOTERM_BP_FAT | GO:0006954 ~ inflammatory response | 0.02 | CBB, CCL23, CLEC7A, CD163 | 6.40 |
| GOTERM_BP_FAT | GO:0006952 ~ defense response | 0.02 | CBB, CCL23, MPO, CLEC7A, CD163 | 4.23 |
| GOTERM_BP_FAT | GO:0006955 ~ immune response | 0.03 | CBB, CCL23, LST1, CLEC7A, TREM2 | 3.77 |
| GOTERM_BP_FAT | GO:0006874 ~ cellular calcium ion homeostasis | 0.04 | CASR, CCL23, GALR2 | 8.53 |
| GOTERM_BP_FAT | GO:0030182 ~ neuron differentiation | 0.04 | LST1, NKX6-2, GALR2, EN2 | 4.75 |
| GOTERM_CC_FAT | GO:0005886 ~ plasma membrane | 0.02 | CASR, OR5C1, SLC7A9, MRGPRF, CDHRS, CD163, CBB, OR4X1, ADAP2, GALR2, GPR25, OR7C2, CLEC7A, TREM2 | 1.65 |
| GOTERM_CC_FAT | GO:0005576 ~ extracellular region | 0.071 | KLKB4, CBB, CCL23, MPO, TAC4, TREM2, CSN1S1, CD163 | 2.03 |
| SP_PIR_KEYW | Disulfide bond | 1.06E-12 | KLKB4, OR3C1, ALPK3, SLC7A9, GALNTL1, CSN1S1, CD163, CBB, OR4X1, CCL23, GALR2, OR7C2, MPO, CLEC7A, TREM2 | 2.99 |
| SP_PIR_KEYW | Glycoprotein | 0.06 | KLKB4, CASR, OR5C1, MRGPRF, CDHRS, CD163, CBB, OR4X1, GALR2, OR7C2, MPO, CLEC7A, TREM2 | 1.75 |

The analysis was performed by DAVID Bioinformatics Resources 6.7 (http://david.abcc.ncifcrf.gov). The genes associated with annotation terms with enrichment p < 0.1 are reported here.
were functional, as demonstrated by intracellular calcium increase following treatment with the neurotransmitter carbachol, and none of the *Gelsemium s.* dilutions affected their growth rate or metabolic activity.

**Low-dilution effects**
The most evident and statistically significant modification of cellular gene expression was induced by the lowest dilution of the medicine that we tested, namely *Gelsemium s.* 2c, as is to be expected with a dose-dependent effect. Spectroscopic analysis of the tested samples confirmed that the starting 1c solution supplied by manufacturer contained a considerable amount of original extract compounds, and proportionate quantities were also detected in the 2c dilutions prepared both by the manufacturer and in our laboratory. However, dilution equal or beyond 3c brought the concentration of detectable compounds below the minimum sensitivity threshold of optical spectroscopy.

Although *Gelsemium s.* contains several different compounds [2,3], the major active alkaloid of this plant is gelsemine, which was present in a concentration of 6.5 × 10⁻⁹ M in the final incubation mixture of cells treated with *Gelsemium s.* 2c. This nanomolar dosage is much lower than the toxic doses that have been reported in poisoning cases [60] and in experimental evaluations of LD50 [14]. In fact, *Gelsemium s.* 2c at the lowest dilution (highest dose) tested in this model system did not cause cell toxicity or viability impairment. This evidence is in agreement with recent hypotheses explaining the homeopathic effects (in the range of very low doses) in the framework of hormesis, where substance which are toxic at high doses turn into therapeutic when diluted to low and ultra-low doses [61]. According to the hormetic theory, ultra-diluted drugs and nanoparticles will act as low-dose stress conditions that could possibly evoke an adaptive response process producing effects that might modulate gene expression [62-64].

The effects on gene expression observed here are specifically targeted to the regulation of certain functions, possibly linked with the plant’s pharmacological activity. Of a total of 45033 transcripts, 49 were down-regulated and 7 up-regulated by the 2c dilution. This effect was quantitatively small (absolute value of fold change between 0.5 and 1.0) but statistically significant (adjusted \( p < 0.05 \)). In general, the prevalence of down-regulation seems to indicate a tendency to reduce cell excitability, especially because several of the genes in question belong to surface receptors involved in GPCR signaling and calcium homeostasis. Moreover, this first microarray screening of the effects of *Gelsemium s.* on neurocytes revealed a significant down-regulation of genes for inflammatory response, olfactory transduction and neuron differentiation. Clearly, this plant species contains a variety of active chemical principles which are presumably involved in different pathways of cell regulation besides the pure neural function, as suggested by reports of possible anticancer and immunomodulating activities [17-19].

A hypothetical neurological target of *Gelsemium s.* has been suggested by studies showing that gelsemine stimulates the biosynthesis of allopregnanolone in the rat brain [30,65], but the genes of neurosteroid enzymatic pathways were not modulated in our cell system. This apparent discrepancy may depend either on the fact that we used a cell line, whereas Venard et al. [30,65] used slices of spinal cord and limbic system, or on the fact that they studied a post-translational level of regulation, linked to enzyme function and not to gene expression. In any case, since in our model the effects of *Gelsemium s.* were quantitatively small, as confirmed by RT-PCR results, no definite conclusions regarding the role of single genes in the action mechanism of this plant can be drawn at this stage.

These microarray findings can be regarded as a preliminary screening of the sensitivity of SH-SY5Y cellular system to *Gelsemium s.*, while more robust conclusions about the possible role of the implicated genes will require to determine whether proteins encoded by the affected genes are similarly changed, through proteomic and phosphoproteomic approaches, and/or further studies using plant purified active compounds.

**Ultra-low doses and high dilutions**
The second major goal of this investigation was to study the dose-effect relationship, which is of central importance in any kind of pharmacological approach. As noted above, the *Gelsemium s.* 2c dilution yielded statistically significant results for 56 genes. This raised the question of whether those same genes, which appeared to be most sensitive to *Gelsemium s.*, would also be modified by higher dilutions. Since the quantitative changes for the 3c and higher dilutions were quite low (Table 3 and Figure 7B), the 4 replicates were insufficient to yield statistical confidence for analysis of single transcripts. We therefore employed cluster analysis to separately describe the trends of 6 gene subsets with similar expression profiles. All 4 down-regulated clusters included genes with negative mean fold changes, though of varying magnitude. Most notably, we found two clusters (2 and 3 in Figure 7) that included a total of 24 genes clearly responsive to *Gelsemium s.* 30c, and characterized by a bell-shaped dilution-effect curve. Exploring results accurately (Table 3), some genes showed an interesting pattern of expression in function of *Gelsemium* dilution. For instance, the EN2 gene that was underexpressed in treated cells exhibited a bell-shaped curve. This tendency can be seen in other genes in the cluster analysis. Moreover, it seems that after 9c, another wave
of expressions or no-expressions is recovered. Maybe future testing even higher dilutions, such as 100c or 200c, the bell-shaped curve could be more evident and, thus, the hypothesis of ultra-sensitive genes could be checked.

For the high dilutions, due to the small changes of gene expression, the only hypothesis statistically evaluable is the global effect of *Gelsemium s.* dilutions on the 49 down-regulated and 7 up-regulated genes, considered as gene-sets. Using the Fisher exact test (*Gelsemium s.* dilutions vs. their respective control solutions), the null hypothesis was rejected for every dilution in the down-regulated gene-set. This outcome of our microarray analysis is astounding if we consider that the 9c and 30c dilutions were obtained from MT extract by dilution factors of $10^{18}$ and $10^{60}$ respectively. Starting from a crude MT containing the active principle gelsermine at a concentration of $6.5 \times 10^{-4}$ M, the 9c dilution would theoretically contain $6.5 \times 10^{-22}$ M gelsermine, corresponding to less than 1 molecule per ml in the final working solution; even in the case of the 5c dilution, where the theoretical gelsermine concentration is $6.5 \times 10^{-15}$ M, it can be calculated that this would correspond to $3.9 \times 10^{7}$ molecules per culture plate, i.e. about 13 molecules per seeded cell. These results suggest that neurocytes have a number of genes with extreme sensitivity to *Gelsemium s.* effects, even if those effects of high dilutions are quantitatively very small (decrease in expression by approximately 10% to 20% compared to the control). The physiological or pharmacological implications of this observation remain to be clarified, but the rejection of the null hypothesis furnishes a new input for the open debate on this kind of therapeutic approach.

**Technical issues and confounding factors**

The puzzling evidence of gene expression changes under the influence of homeopathic dilutions prompt an analysis of the possible confounding factors that might explain the effects observed. We adopted different measures to address the issue of possible experimental artifacts. To avoid dye-bias artifacts a single-channel microarray was employed. We adopted a microarray design with probes of the same probe-set located in not contiguous positions on the array, so that artifacts due to uneven hybridization would only affect a subset of probes for a probe-set. Anyhow, the absence of spatial biases in fluorescence signal was assessed by checking the coefficient of variation of the mean signal intensities of different portions of each array. The experimental set up could have introduced biases and "position effects" if handling of control and *Gelsemium s.* matched dilutions was not equivalent. Actually, we conducted four independent experiments in which *Gelsemium s.* dilutions and the corresponding vehicle controls were processed in tandem (from drug addition to RNA extraction and cDNA synthesis). In every subarray of the chip, each transcript was targeted with three separate probes, merging the fluorescence values and attributing a statistical score.

Regarding the statistical analysis, the large number of genes of the complete set causes some problems concerning the choice of “interesting” genes. The approach followed here was quite stringent and limited the number of genes considered, reducing the probability of “false positive” results, but forcing to discard some possibly interesting genes from the analysis. Moreover, the small entity of the expression changes observed with high dilutions unavoidably reduced statistical inference in the single genes, especially since multiplicity corrections were applied. The choice of analyzing the sign of the fold changes in a pool of genes, rather than the variance of a single gene, may lead to a loss of statistical information, to the advantage of greater precision in discarding the null hypothesis. Further research specifically oriented on the most responsive genes, with suitable sample sizes, could possibly overcome this limitation of the microarray approach.

**Physico-chemical and biological hypotheses**

Our results are in keeping with a number of experimental observations from a variety of research fields, confirming that highly diluted compounds exert statistically significant effects on biological systems [66-69]. Thus far there is no satisfactory or unifying theoretical explanation for these claims, though some have hypothesized that the dynamics of the solvent water (or water-ethanol) on a mesoscopic scale may play a part [70]. Three major models for how this happens are currently being investigated: the water clusters or clathrates, the coherent domains postulated by quantum electrodynamics, and the formation of nanoparticles from the original solute plus solvent components. It has been suggested that a major role in the formation of water clusters is played by silica released from the glass containers which are usually employed in the preparation of homeopathic drugs [71]. Silica nanostructures formed during succussion in glass and/or biosynthesized by specific plant extract tinctures may also acquire and convey epitaxial information from the remedy source materials into the higher potencies [21,72,73]. In our experimental model, since the *verum* were succussed samples, we used the succussed ethanol/water solutions as negative controls and evaluated preliminarily the variability of the negative system before assessing the biological effect of the succussed/diluted drug. Notably, in our experiments serial dilutions/succussions were performed in glass bottles, with the exception of the last step, which was developed in polystyrene tubes. Thus the hypothetical role of silicates in nanoparticle formation is pertinent, but also the contribution of polystyrene should not be excluded [74].
Recent evidence supports the plausibility that homeopathic Gelsemium s. in the potencies tested could contain crudely formed nanoparticles. Bel-Haaj et al. [75] demonstrated that just extended ultrasonication of plant starch can create starch nanoparticles in water. Moreover, electron microscopic evidence of nanoparticles has been obtained in several different plants prepared homeopathically [76]. Gelsemium mother tincture itself, like many other plant extracts, can biosynthesize nanoparticles of silver metal from precursor substrate [77]. Nanoparticles have unique biological and physicochemical properties, including increased catalytic reactivity, protein and DNA adsorption, bioavailability, dose-sparing, electromagnetic, and quantum effects that are different from those of bulk-form materials [23]. As an example, Prakash and colleagues [78] compared in model animals the anti-anxiety effects of hypericum prepared as gold nanoparticles versus a bulk form and observed more significant effects with the nano-hypericum, even at a 10-fold lower dose. Higher cellular uptake of nano-encapsulated (poly lactide-co-glycolide) Gelsemium s. than of its bulk form has been observed by Bhattacharyya et al. [79].

The hypotheses regarding the possible biological mechanisms of highly diluted/dynamized solutions (beyond Avogadro-Loschmidt limit) at the level of DNA expression varies on the suggestion of dynamism of regulatory networks poised near the phase transition between order and chaos and extremely sensitive to initial conditions and small perturbations is well known to occur. Chaotic regimes have been found in a number of physiological systems, including neural systems [86-88], and this result in enhanced susceptibility to low energy inputs and to small changes of regulatory factors. According to this argument, the highly diluted drug might be regarded as a solution endowed with water clusters and/or nanoparticle structures capable of communicating some pharmacological information, through a resonance process, to biological networks and to cell critical systems such as macromolecules, alpha-helices, filamentous structures, receptors and DNA networks. This effect could be mediated by the participation of a dynamic intracellular network which may be presumed to exist in living cells [89].

Conclusions
This study provides evidence that Gelsemium s. exerts a prevalently inhibitory effect on a series of neurocyte genes across a wide dose-range. The effect decreases with increasing dilutions, but whole genome expression analysis allowed to detect statistically significant changes even at the highest dilutions tested (9c and 30c). The results suggest the extreme sensitivity of human gene expression to regulation by ultra-low doses and high dilutions/dynamizations of a plant remedy and encourage further efforts in research on this field. Studies using “omic-based” approaches and systems biology should be particularly worthy at generating new hypotheses on mechanisms for the effects of highly diluted natural compounds.

Additional files

Additional file 1: Microarray expression values of 56 transcripts in SH-SYSY neurocytes treated with Gelsemium s. or control dilutions. Data are reported as Log2 transformed fluorescence values from four replicate microarray assays.

Additional file 2: Frequency of fold change values in a randomly chosen gene-set after Gelsemium s. treatments. A list of 49 genes was generated by randomized selection from the whole transcriptome using SPSS software (excluding the differentially expressed genes) and fold change was calculated from the difference of mean Log2 fluorescence values of Gelsemium s-treated samples (Gnc) vs those of controls (Ctnc). Absolute fold changes less than or equal to 0.05 were considered null. Blue bars: frequencies of genes with negative fold change (< −0.05); grey bars: frequency of unaffected genes (from −0.05 to 0.05); pink bars: frequencies of genes with positive fold change (> 0.05). For these randomly selected genes, Fisher exact test is not significant in any dilution.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
PB conceived the experiments, MM, DO and MC designed and performed the experiments, PT, MB, MM, and DO analyzed the data, MM and PB wrote the paper. All authors read and approved the final manuscript.

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