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## Differential representation of sunflower ESTs in enriched organ-specific cDNA libraries in a small scale sequencing project

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### Abstract

**Background:** Subtractive hybridization methods are valuable tools for identifying differentially regulated genes in a given tissue avoiding redundant sequencing of clones representing the same expressed genes, maximizing detection of low abundant transcripts and thus, affecting the efficiency and cost effectiveness of small scale cDNA sequencing projects aimed to the specific identification of useful genes for breeding purposes. The objective of this work is to evaluate alternative strategies to high-throughput sequencing projects for the identification of novel genes differentially expressed in sunflower as a source of organ-specific genetic markers that can be functionally associated to important traits.

**Results:** Differential organ-specific ESTs were generated from leaf, stem, root and flower bud at two developmental stages (R1 and R4). The use of different sources of RNA as tester and driver cDNA for the construction of differential libraries was evaluated as a tool for detection of rare or low abundant transcripts. Organ-specificity ranged from 75 to 100% of non-redundant sequences in the different cDNA libraries. Sequence redundancy varied according to the target and driver cDNA used in each case. The R4 flower cDNA library was the less redundant library with 62% of unique sequences. Out of a total of 919 sequences that were edited and annotated, 318 were non-redundant sequences. Comparison against sequences in public databases showed that 60% of non-redundant sequences showed significant similarity to known sequences. The number of predicted novel genes varied among the different cDNA libraries, ranging from 56% in the R4 flower to 16% in the R1 flower bud library. Comparison with sunflower ESTs on public databases showed that 197 of non-redundant sequences (60%) did not exhibit significant similarity to previously reported sunflower ESTs. This approach helped to successfully isolate a significant number of new reported sequences putatively related to responses to important agronomic traits and key regulatory and physiological genes.

**Conclusions:** The application of suppressed subtracted hybridization technology not only enabled the cost effective isolation of differentially expressed sequences but it also allowed the

identification of novel sequences in sunflower from a relative small number of analyzed sequences when compared to major sequencing projects.

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## Background

Cultivated sunflower (*Helianthus annuus L.*) is one of the most important sources of vegetable oil worldwide. During the last decade, rapid advances in applied genetics and genomic technologies have led to the development of saturated sunflower genetic maps based on different molecular markers including RFLP, AFLP and SSR [1–10]. More recently, large-scale cDNA sequencing projects have identified expressed sequence tags (ESTs) in different plant species. Today, more than 100 plant species are represented in the EST division (dbEST) of GenBank [http://www.ncbi.nlm.nih.gov/dbEST/dbEST\\_summary.html](http://www.ncbi.nlm.nih.gov/dbEST/dbEST_summary.html) with a total of 2,063,406 entries. However, *Arabidopsis*, rice, maize, tomato and soybean ESTs projects gather more than 50% of the total entries. The *Compositae* is represented by 113,149 entries, of which 44,961 correspond to sunflower ESTs. These projects allow the characterization of full sets of transcribed genes in the target organisms and provide, at the same time, a source of genetic markers that can be functionally associated to important agronomical traits reinforcing and complementing the use of anonymous markers. Thus, the use of EST-based markers could lead to genetic mapping of a gene that directly affects the trait or a specific sequence could be targeted due to its predicted function based on sequence comparison [11]. ESTs generated from cDNA libraries should represent, ideally, all expressed genes in a target organ/tissue, at a specific developmental stage and/or in a specific environment. However, differences in expression level among genes in a given tissue yield mRNAs that differ in abundance, making it difficult to capture rare mRNA in cDNA libraries. This problem also leads to redundant sequencing of clones representing the same expressed genes, affecting the efficiency and cost effectiveness of the EST approach [12] which hinders research laboratories with small budgets to perform EST characterization studies. To avoid this problem, different strategies based on normalized cDNA libraries have been reported in many different organisms [12–14] including plants [15–17]. In this study, we report for the first time in sunflower, the isolation and characterization of ESTs from organ-specific cDNA libraries constructed by suppressed subtractive hybridization [18] as an alternative to identify differentially expressed sunflower transcripts. We analyzed the efficiency of the subtraction and enrichment methods for each cDNA library generated and present the differential level of representation for functional EST groups based on Gene Ontology annotation [19], as well as a comprehensive description of individual non-redundant sequences generated.

## Results and Discussion

### Construction of organ-specific cDNA libraries

Different cDNA libraries were constructed after subtractive hybridization. Firstly, a reciprocal experiment was designed to determine the efficiency of the subtraction procedure to clone differentially expressed genes in two different plant organs. Poly (A<sup>+</sup>) RNAs from R4 flower and from leaf were used to generate tester and driver cDNAs, respectively, for the flower library and vice versa for the leaf library. Clearly distinctive patterns of differential transcript abundance could be observed when these two cDNA libraries were compared. Sequence comparison among the two generated cDNA showed that as much as 92% (209 out of 227 sequences) and 62.5% (80 out of 128 sequences) of the analyzed sequences were unique to R4 flower and leaf libraries, respectively (Table 1). These results indicate the high efficiency of this technique to isolate organ specific transcripts compared to other reports on organ derived cDNA libraries [20]. Endo et al. [20] reported that 64.8% of ESTs sequences isolated from *Lotus japonicum* flower bud had not been found in EST sequences of the whole plant. Other reports indicated that only 12% of ESTs of an equalized cDNA library constructed from different developmental stages of inflorescence in *Arabidopsis thaliana* were unique to inflorescence tissue [21]. This high percent of organ-specific sequences for the flower and leaf libraries encouraged us to construct additional cDNA libraries. Stem and root cDNA libraries were subtracted with leaf cDNA in order to avoid high redundancy of photosynthesis related sequences. The R1 flower bud cDNA library was subtracted with R4 flower cDNA with the aim to identify specific gene induction during early stages of development.

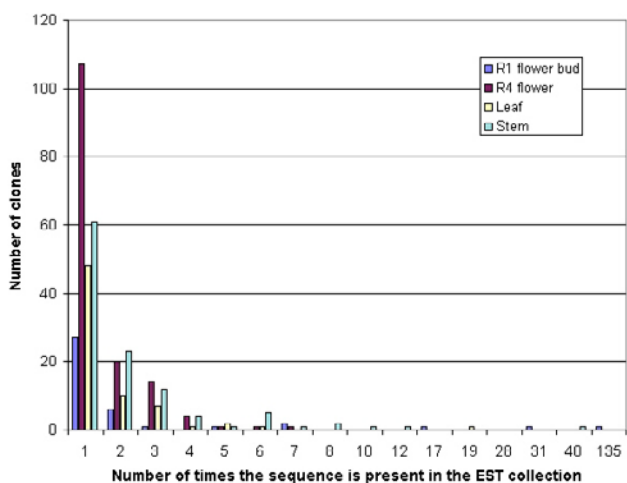
### Sequence analysis

Table 1 summarizes the total number of isolated, sequenced and analyzed clones, differential and non-redundant sequences and the average insert size and the average ORF per cDNA library. A total of 1073 randomly selected non-directionally clones from the different cDNA libraries were sequenced from which, after removing low quality and contaminant ribosomal RNA sequences, 919 readable sequences were generated, edited and annotated as described in experimental procedures. 5' and 3' sequences were equally represented in the generated EST database. The analysis of sequence redundancy was performed by sequence comparison using local BLASTN through a clustering system running under an alpha version of Biopipeline<sup>®</sup> and by using the Cap3 contig assembly program [22]. The Biopipeline<sup>®</sup> clustering revealed a

**Table 1: Number of isolated, analyzed, differential and non redundant sequences by organ-specific cDNA library.**

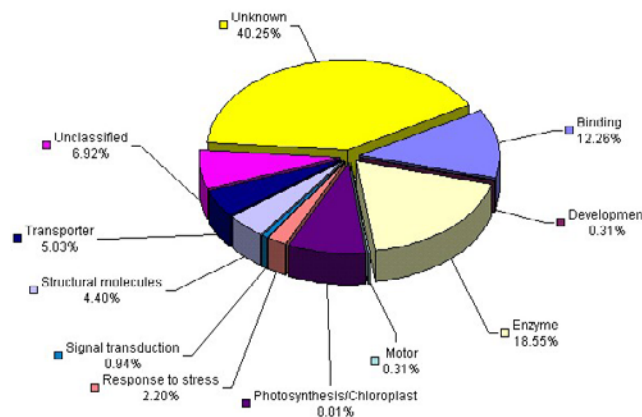
Total ESTs	R1 flower (1)	R4 Flower (2)	Leaf (3)	Stem (4)	Root (5)	Total
Isolated	504	384	268	400	115	1671
Sequenced	261	269	159	312	72	1073
Analyzed	245	227	128	282	55	919
Differential sequences between (2) and (3) <sup>a,b</sup>		209	80			
Non-redundant	32	140	63	79	4	318
Non-redundant percentage <sup>b</sup>	13	62	49	28	7	36
Differential sequences <sup>c</sup>	31	131	42	81	4	289
Average ORF <sup>c</sup>	269	220	257	265	225	247
Average insert size (bp) <sup>b</sup>	495	365	443	463	370	441

<sup>a</sup> Sequences detected exclusively either in the flower or leaf cDNA libraries. <sup>b</sup> Out of analyzed sequences. <sup>c</sup> Sequences detected exclusively in the indicated cDNA library out of non-redundant sequences.



**Figure 1**  
**Frequency of redundant clones among ESTs from different organ-specific cDNA libraries.**

total of 318 non-redundant sequences, meanwhile Cap3 running with an overlap cut-off identity of 95% and a minimum overlap of 25 bases detected a total of 29 contigs composed of two to four sequences and 249 singletons. The observed discrepancy between the unigene set outcomes from both methods is based on the different algorithms used by each program. Manual check of the outcome results confirmed that comparison using Biopipeline<sup>®</sup> was more efficient in detecting redundancy without losing sensitivity in the detection of gene variants. Thus, further sequence comparisons were performed using the 318 non-redundant sequences as unigen set. Sequence redundancy varied among the different cDNA libraries (Figure 1). The least redundant library was the R4 flower library with a total of 140 unique sequences from



**Figure 2**  
**Expression analysis of ESTs from organ-specific cDNA libraries.** cDNA clones with significant similarity to protein sequences in SWALL were classified according to Gene Ontology annotation. Sequences with no hits to known protein sequences from BLASTX comparison were classified as unknown. ESTs with significant similarity according to BLASTX comparison but with no GO term definition associated to them were referred as unclassified. Functional analysis includes all non-redundant generated ESTs.

227 analyzed ESTs (62%). In contrast, the most redundant cDNA libraries were: the R1 flower bud (13% of unique sequences) and the root cDNA library (7% of unique sequences). The leaf and stem cDNA libraries exhibited intermediate redundancy levels, with 49 and 28 % of non redundant sequences, respectively. The high level of redundancy in the early flower bud library compared to the late flower bud is likely to be related to differences in this specific subtraction protocol. While the R4 flower library was subtracted with a non-related driver

cDNA (leaf cDNA), allowing the detection of transcripts not represented (or represented at a lower level) in the leaf tissue, the R1 flower bud cDNA was arrested with an mRNA population from the same organ but at a different developmental stage. Transcripts from the same organ/tissue share a high number of identical mRNAs and, consequently, a relatively reduced pool of differentially expressed transcripts remains unsubtracted at a specific developmental stage. In the case of the root library, the analysis of redundancy should be treated with caution due to the small number of cDNA molecules that remained unsubtracted after the hybridization step. Thus, studies on predicted functionality were not conducted for this latter cDNA library. The leaf and stem cDNA libraries exhibited higher levels of redundancy compared to the R4 flower cDNA library. The higher redundancies in these two libraries are due to a high representation of photosynthesis related sequences.

Analysis of organ-specificity among non-redundant sequences confirmed that a high proportion of the non-redundant sequences in each library corresponded to sequences only detected in that tissue. In the R4 flower and stem cDNA libraries, 93.5% and 98.7% of the analyzed sequences were unique to those libraries, respectively (Table 1). A global analysis including all constructed libraries revealed that 87.8% of the generated non-redundant ESTs were indeed differentially expressed sequences.

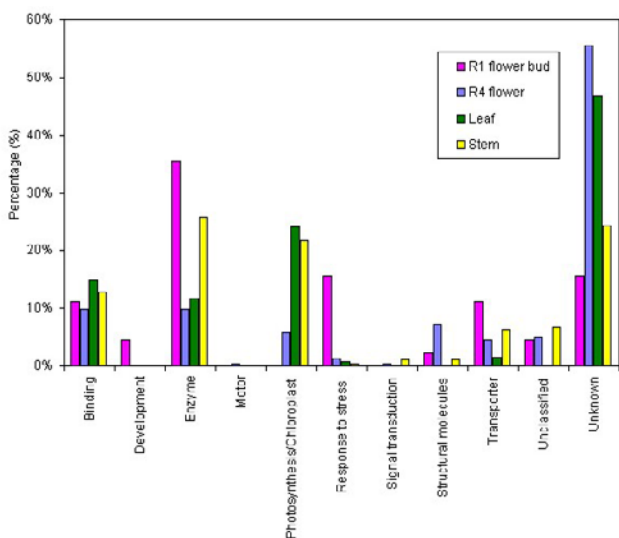
#### **EST analysis based on predicted gene function**

Sunflower ESTs were grouped into different functional categories according to their predicted gene products based on sequence comparison with the current SWISS-PROT/ TrEMBL (SWALL) data bases. Annotation was performed based on Gene Ontology (GO) [24] terms and functional categories were defined accordingly (Figure 2). This annotation allows the classification of generated ESTs by function [23] with the aim to create universal vocabulary for consensus annotation [24]. A complete list of non-redundant sequences generated here, including BLASTX top hit sequence in SWALL, GO term definition and GO identification number for each sequence is provided on Additional file 1.

A total of 190 sequences (60 %), out of 318 non-redundant ESTs, showed significant similarity to known gene sequences in the database with a stringency level ( $E$  value) of  $10^{-3}$  and a score value higher than 80. No significant differences in average insert length in both were detected between the sequences that match previous entries on GenBank and those that did not show similarities. These results indicate that the lengths of the sequences reported in this study are good enough to retrieve significant hits in GenBank database. Out of the remaining 128 sequences

(40 %) that exhibited no significant similarity to known genes, 90 sequences (70%) exhibited significant homology to ESTs with unknown function on public databases while the remaining sequences representing 32 % are new reported sequences. The *Compositae* is represented in the GenBank by 113,149 entries of which 44,961 correspond to recently deposited sunflower ESTs and the rest to lettuce (*Lactuca sativa*; Composite Genome Initiative, CGI, <http://cgpdb.ucdavis.edu/database/cgpdb.php>). Out of these 44,961 sunflower ESTs, 15,248 are unique sequences, and only circa 2,061 are functionally annotated sequences (TIGR Gene Indices, <http://www.tigr.org/tdb/tgi/hagi>). In spite of this extensive amount of available information, sequence comparison of the 318 non-redundant sequences generated in this study against 37,208 unique *H. annuus* and *L. sativa* sequences (HaGI and LsGI <http://www.tigr.org/tdb/tgi/lsgi>, TIGR) showed that 197 (59.9%) did not exhibit significant similarity to previously reported sunflower ESTs whilst 228 (69.3%) did not match *L. sativa* ESTs. The important level of homology found with other plant ESTs that do not belong to the *Compositae* family indicate that this fact was not due to highly variable or non-coding sequences present at the 3' end of the mRNA. Since the ESTs in this study are derived from polyA RNA and thus enriched in 3' end sequences of the mRNAs, while the ESTs recently deposited at the CGI are enriched in the 5' end of the mRNA, a comparative study of outcome BLASTX was performed in order to determine if the 197 newly detected sunflower genes were indeed represented at GenBank by previously deposited sunflower ESTs from different gene regions. This analysis revealed that most of these sequences share annotation but do not share identities at a nucleotide level, thus some of them are likely to be variants of gene families.

The "unclassified" class correspond to sequences that showed significant similarity to SWALL sequences using BLASTX search but they do not have an associated GO term. Most of these sequences correspond to hypothetical proteins with unknown function. The relative abundance of EST categories varied according to the analyzed library (Figure 3). ESTs showing no significant similarity (unknown) represented 56% of the analyzed sequences in the R4 flower cDNA library, while this category was considerably lower in the other libraries, ranging from 16 to 47%. Previous studies reported similar values of predicted novel genes isolated from different normalized cDNA libraries. Asamizu *et al.* [15] reported that 45% of non-redundant ESTs generated from different plant tissues including aboveground organs, flower buds, roots and liquid-culture seedlings were predicted to be novel genes. In a drought-stressed normalized cDNA library from rice seedlings, up to 28.2% of the non-redundant sequences were novel [17].



**Figure 3**  
**Comparison of ESTs classified by predicted function among four organ-specific cDNA libraries.** Functional classification of all generated ESTs was done as described in Figure 2. Percentage of ESTs included in each functional class is compared among four differential cDNA libraries.

"Structural proteins" and "motor" as well as sequences related to cell growth and metabolisms, here included in the "enzyme" class, showed a low level of representation compared to the corresponding values obtained by non-normalized cDNA libraries [25,26]. A similar under representation of ESTs from the cell metabolisms category was reported for other normalized cDNA libraries [16]. This result shows that the normalization step that took place in the construction of the cDNA libraries was efficient in diminishing the level of highly abundant transcripts equally represented in the different analyzed tissues.

As expected, ESTs related to the "photosynthesis/chloroplast" class were highly abundant in the leaf (24%) and stem (22%) libraries while these sequences were absent in the R1 flower bud and very low represented (6%) in the R4 flower cDNA library. Conversely, the leaf and stem libraries showed a low representation of ESTs homologous to stress related sequences, which barely reached 1%. The proportion of "response to stress" sequences showed a higher representation in the R1 flower bud library (16%). Besides the sequences classified as "response to stress" class according to GO terms, there are some other ESTs included in other categories such as "enzyme", "transporter" and "binding" that have been associated to biotic and/or abiotic stress in previous studies [27–32]. Table 2 includes a complete list of 33 non-redundant dif-

ferential ESTs related to defence and stress response according to GO terms and literature references. Interestingly, agronomical important sequences related to response and/or defence to pathogens such as glucanases (BU671807), germin-like proteins (BU671889) and polygalacturonase inhibitor proteins (BU671906) are new reported ESTs sequences for sunflower as they are not represented in the current EST database. Most of these defence related transcripts were differentially detected in the R1 flower bud library without exposure to any external stimuli, thus reinforcing the importance of designing highly specific organ/developing stage cDNA libraries for detection of low abundant transcripts. This observation confirmed previous reports of higher level of defence related transcripts in developing flowers [33–36]. The "enzyme" class is highly represented in the stem (30%) and R1 flower bud (36%) libraries compared to the leaf and R4 flower library. This class includes a significant number of defence related enzymes differentially detected in the R1 flower bud and stem cDNA libraries. Within this group, those ESTs with significant similarity to pathogen defence-related genes like those coding for germin-like proteins, lipid transfer proteins, polygalacturonase inhibitor factors, protease inhibitors, as well as those genes related to abiotic stress responses like fructosyl transferase, salt-stress induced tonoplast, aquaporin protein, dehydrin protein were mostly detected as unique or low copy number sequences. On the other hand, the more abundant stress related protein genes like glucanases, catalases, peroxidases, jasmonate-induced proteins, thaumatin-like proteins, heat shock proteins were detected more frequently in most of the constructed cDNA libraries. These results are consistent with a previous report on the identification of defence-related genes by suppression subtractive hybridization in rice [27]. In that study the authors compared this strategy with a differential screening performed on a non-differential cDNA library. They found that the suppressed subtracted hybridization allowed the detection of medium-low abundant genes such as protein kinases and transcription factors whilst the differential screening technique detected mostly abundant transcripts such as *PR* genes.

In the present study, the "binding" class is equally represented in all the analyzed libraries, although this class includes sequence with putative involvement in diverse processes such as transcription and translation factors, ATPase and cation binding proteins. Within this group low abundant transcripts like those coding for transcription factors and homeotic factors were specially detected in the R4 flower cDNA library. ESTs with homology to genes coding for signalling enzymes as MAP protein kinase and serine/threonine phosphatase were only detected in the stem cDNA library (Figure 2b). The functional category of "transporter" is represented by

**Table 2: Differential ESTs related to response to biotic and/or abiotic stress**

AN <sup>a</sup>	GO ID <sup>b</sup>	GO functional definition <sup>b</sup>	AN <sup>c</sup>	BLASTX hit <sup>d</sup>	E value
BU671794	GO:000238	Phosphoethanolamine N methyltransferase	Q944HO	Putative phosphoethanolamine N methyltrans	2.00E-26
BU671801	GO:0004332	Fructose biphosphate aldolase	Q9SXX5	Plastidic aldolase	2.00E-91
BU671803	GO:0042027	Cyclophilin-type peptidyl-prolyl cis-trans isomerase	Q9M530	Cyclophilin (EC 5.2.1.8) (Peptidyl-prolyl cis-trans isomerase) (PPlase)	3.00E-18
BU671805	GO:0008246	Electron transfer flavoprotein	Q39640	Glycolate oxidase	2E-94
BU671807	GO:0004553	Hydrolase	Q9M473	Putative-beta 1,3-glucanase	5.00E-38
BU671832	GO:0003773	Heat shock protein	Q02028	Stromal 70 kDa heat shock-related protein	3.00E-27
BU671840	GO:0004096	Catalase	P45739	Catalase	1.00E-95
BU671841	GO:0004332	Fructose biphosphata aldolase	P93565	homologous to plastidic aldolases	3.00E-21
BU671845	GO:0000287	Magnesium binding	Q93WE2	Magnesium chelatase subunit	4.00E-36
BU671864	GO:0004332	Fructose-bisphosphate aldolase	Q9SXX5	Plastidic aldolase	1.00E-54
BU671866	GO:0009058	Biosynthesis	Q39049	Magnesium chelatase subunit.	4E-16
BU671867	GO:0030145	Magnesium binding	Q943W1	Oxygen-evolving enhancer protein I	2.00E-92
BU671886	GO:0004299	Proteasome endopeptidase	O23708	Proteasome subunit alpha type 2	2.00E-17
BU671887	GO:0005215	Transporter	Q9ZR68	Aquaporin I	2.00E-84
BU671888	GO:0004553	Hydrolase, hydrolyzing O-glycosyl compounds	Q9M453	Putative beta-1,3-glucanase.	3.00E-58
BU671889	GO:0030145	Manganese binding	O48999	Germin-like protein 3	2.00E-35
BU671904	GO:0004564	Beta-fructofuranosidase	O81985	1,2-beta-fructan 1F-fructosyltransferase	2.00E-51
BU671906	GO:0005489	Electron transport	Q94L58	Polygalacturonase inhibitor protein	3.00E-39
BU671909	GO:0016068	Immediate hypersensitivity response	Q93YX9	Lipid transfer protein	1.00E-37
BU671910	GO:0016068	Immediate hypersensitivity response	Q9M6B8	Lipid transfer protein	2.00E-15
BU671924	GO:0004197	Cysteine-type endopeptidase	Q8VVS1	Putative cysteine proteinase	4.00E-42
BU671928	GO:0003755	Peptidyl-prolyl cis-trans isomerase	Q8L5T1	Peptidylprolyl isomerase (Cyclophilin)	3.00E-18
BU671929	GO:0004222	Metalloendopeptidase	O22941	Putative zinc protease	1.00E-61
BU671944	GO:0005509	Calcium ion binding	O49301	T26J12.7 protein.	7.00E-13
BU671955	GO:0009607	Response to biotic stimulus	P13046	Pathogenesis-related protein R major form	3.00E-34
BU671960	GO:0004766	Spermidine synthase	48658	Spermidine synthase I (EC 2.5.1.16)	1.00E-21
BU671972	GO:0008168	Methyltransferase	Q9LW67	Ankyrin-like protein	7.00E-33
BU671989	GO:0004601	Peroxidase	O64970	Cationic peroxidase 2	1.00E-40
BU671977	GO:0005489	Electron transporter	O04002	Chloroplast drought-induced stress protein of 32 kDa	6.00E-41
BU672016	GO:0004867	Serine protease inhibitor	Q8LNY0	Protease inhibitor 2	2.00E-13
BU672055	GO:0004869	Cysteine protease inhibitor	Q9MB08	Multicystatin.	2.00E-12
BU672102	GO:0005489	Electron transporter	O04002	Chloroplast drought-induced stress protein of 32 kDa	2.00E-56
BU672106	GO:0004124	Cysteine synthase	Q8Y0X6	Probable cysteine synthase B (CSASE B) protein (EC 4.2.99.8).	9.00E-21

<sup>a</sup> GenBank accession number of ESTs from the present study. <sup>b</sup> Identification number and functional definition according to GO annotation <sup>c</sup> The GenBank accession number of most similar sequence to the sunflower EST. <sup>d</sup> Similarity search was conducted using BLASTX program. EST putative function was assigned according to the highest similar sequence on GenBank.

sequences with similarity to carrier protein genes as ATP-binding cassette (ABC) and electron transporters that were mainly detected in the R4 flower library. ESTs with similarity to homeobox genes here included in "development" were only detected in the early flower bud cDNA library. The homeobox sequences isolated in this work did not show similarity to previously reported sunflower homeobox genes [37–39]. Preliminary results showed that some of the agronomical interesting sequences, including those putatively related to response to biotic and abiotic stress, revealed polymorphisms when used as genetic markers in the analysis of genetically segregant populations derived from the crossing of parental lines with contrasting biotic and abiotic stress resistance behaviour (not shown).

The application of suppressed subtracted hybridization technology for the detection of differential ESTs allowed

the identification of novel sequences in sunflower from a relative small number of analyzed sequences in spite of the large number of ESTs that have been recently release. Particularly interesting was the detection of a significant number of ESTs related to response to both abiotic and biotic stresses, as well as low abundance transcripts with high similarity to homeobox genes, transcription factors and signalling component genes that were not represented in the sunflower EST division at the GenBank. The R4 flower cDNA library was the library that provided the largest number of novel genes in sunflower, whilst the R1 flower bud library was particularly enriched in defence related genes. The detection of these novel sequences could contribute to the development of EST-based markers for important agronomic traits such as resistance to pathogens and tolerance to different environmental stresses such as extreme temperatures and drought, which

are aspects crucial for sunflower crop improvement in many of the cultivated areas in the world.

## Conclusions

The application of suppressed subtracted hybridization technology enabled the isolation of a significant number of organ-specific sunflower ESTs and allowed the identification of novel sequences from a relative small number of analyzed sequences. Redundancy level and percent of novel sequence detection varied among differential libraries reinforcing the importance of a careful selection of both target and driver transcript population according to project aims. In this work the R4 flower cDNA library provided the largest number of novel genes in sunflower, whilst the R1 flower bud library was particularly enriched in defence related genes. Some of the novel sequences reputed here share annotation but do not share identities at a nucleotide level with sunflower ESTs on public databases and thus, they are likely to be variants of gene families. We report for the first time in sunflower a significant number of novel sequences related to responses to abiotic and biotic stresses as well as low abundant transcripts with high similarity to homeobox genes, transcription factors and signalling components.

## Methods

### Plant material

Sunflower seedlings (public inbred line RHA89) were grown under controlled green house conditions (20–24°C and 16 h light/ 8 h dark cycle), and then transplanted to the field during the crop season to develop mature plants. Leaves, stems, and capitulum buds from 1 to 2 cm of diameter (early flower buds) and 3 to 4 cm of diameter (late flower buds) were harvested from two months old plants and immediately frozen in liquid nitrogen. Roots were harvested from 15 day old plants grown in sand under green house conditions. All samples were stored frozen at -80°C until processed.

### Total and poly (A+) RNA isolation

Total RNA was extracted from approximately 2 g of tissue using TRIzol® reagent following manufacturer recommendations (Invitrogen, USA). Poly (A+) RNA was isolated from 200–500 µg of total RNA using NucleoTrap® System (Promega, USA). RNA integrity was analyzed by checking its electrophoretic mobility on 1.5 % agarose gels in ME buffer (400 mM MOPS, 100 mM Na acetate, 10 mM EDTA pH 8.0, in diethyl-pyrocabonate treated water). mRNA quantification was performed by UV absorbance at 260 nm (GenQuant pro, Amersham-Pharmacia, UK).

### Construction of cDNA libraries

Differential cDNA libraries were constructed from different tissues including leaves, stems, roots and flower buds and from different developmental stages (e.g. R1 and R4

according to the description of sunflower growth stages by Schneider and Miller [40]) using PCR-Select cDNA Subtraction Kit® (Clontech, USA). Firstly, cDNA was synthesized from 0.5–2.0 µg of poly (A+) RNA from the two types of tissues being compared. The tester (target tissue) and driver (reference tissue) cDNAs were then digested with RsaI, that yields blunt end fragments of approximately 400 bp length in average. We defined different driver populations for the different specific libraries, depending on specific interests. Leaf cDNA collection was arrested against a late flower bud cDNA population. Stem early flower bud and root cDNA collections were arrested against a leaf cDNA population.

Both, tester and driver, cDNA populations were processed following manufacturer instructions, with some modifications. The tester cDNA was subdivided into two halves, and each half was ligated to different cDNA adaptors. Two hybridization rounds were performed with an excess of driver cDNA. Hybridization conditions were performed as recommended by the manufacturer. The resulted products were subjected to two cycles of PCR with adaptor targeted primers to amplify the desired differentially expressed sequences. Amplifications were performed on a PT-100 DNA thermocycler (MJ Research, USA). First PCR master mix contained 10x PCR reaction buffer, 0.2 mM dNTPs, 0.4 µM PCR primer 1 and Advantage cDNA polymerase (Clontech, USA). PCR was performed under the following conditions: 94°C (30 sec) denaturing step followed by 27 cycles each consisting of a denaturation step at 94°C (30 sec), an annealing step at 66°C (30 sec) and an elongation step at 72°C (10 min). The second PCR master mix contained 10x PCR reaction buffer, 0.2 mM dNTPs, 0.4 µM nested PCR primer 1, 0.4 µM nested PCR primer 2 and Advantage cDNA polymerase, PCR was run through 12 cycles each consisting of a denaturing step at 94°C (30 sec), an annealing step at 66° (30 sec) and an elongation step at 72°C (1.5 min).

cDNA molecules were size-selected and fractions larger than 250 bp were cloned non-directionally into the pGem-T-Easy Vector® (Promega, USA). Ligation was performed at 4°C for 48 h and the resulting ligation product was used to transform *Escherichia coli* (XL1-blue strain) by electroporation (Pulse Controller, BioRad, USA).

### Template preparation

cDNA libraries were plated onto solid Luria Bertani (LB) medium containing ampicillin. Recombinant clones were selected by β-galactosidase activity in media containing X-GAL and IPTG. White colonies were randomly picked to 364 well plates containing Freezing Medium (36 mM K<sub>2</sub>HPO<sub>4</sub>, 13.2 mM KH<sub>2</sub>PO<sub>4</sub>, 1.7 mM sodium citrate, 0.4 mM MgSO<sub>4</sub>, 6.8 mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, in LB medium and 4.4 % glycerol), grown overnight and later stored at -70°C.



Recombinant plasmids were isolated using REAL 96 prep kit (Qiagen, Germany) as recommended by the supplier. Insert sizes of individual recombinant clones were examined by electrophoresis of EcoRI digestion products on 1.2 % agarose gels in TAE buffer [41].

#### Sequencing and sequence analysis

Recombinant plasmids were single-pass sequenced from the T7 universal primer site at sequencing facilities (Laboratorio de Alta Complejidad, IMyZA – CICVyA – INTA Castelar, Argentina; Centro de Biología Molecular e Engenharia Genética – CBMEG Universidade Estadual de Campinas, Sao Paulo, Brazil and/or Department of Plant Pathology, Kansas University). Reverse sequencing was performed from the SP6 primer site, only when the forward sequences failed or were uninformative due to a short length. The generated EST sequences were stored in a relational database in which both 5' and 3' sequences were equally represented. Vector and uninformative sequences were automatically removed using computer program routines. The processed sequence were output to FASTA formatted files and a pile up (Biopipeline®) step routine written by in-house staff (S.L., Bioaxioma S.A.) was applied to detect remaining vector artifacts by comparing against a full vector sequence database. Redundancy was also analyzed by means of a clustering systems running under an alpha version of Biopipeline®. This system displays a graphic matrix which aligns the top scoring hits sequences in a score matrix. Sequences that exhibited more than 80% identity over total large sequence were considered identical or closely related and were assigned to a specific group. Sequence alignment of those highly similar sequences was confirmed by sequence alignment programs (ClustalW [42]). Contig analysis of the grouped ESTs was done using the contig assembly program Cap3 [21].

Sequence similarities searches against different protein databases were conducted using Advanced BLAST program [43]. Default BLAST parameter values were used except for the *E* value ( $E = 10^{-3}$ ). The top scoring hits were automatically annotated according to the putative function returned by BLASTX. Gene Ontology (GO) annotation was performed using the GOBlet software package [44] and a GO term associated to each sequence showing a significant similarity hit by BLASTX against SWALL search was defined. Sequences comparison against plant division ESTs, HaGI and LsGI were performed locally using BLASTN. These datasets were downloaded from public databases and the "Standalone WWW BLAST Server" from the National Center for Biotechnology Information (NCBI; <ftp://ftp.ncbi.nih.gov/blast>).

#### Authors' contribution

PF carried out subtracted cDNA libraries, DNA sequencing, and participated in data analysis, EST annotation and manuscript preparation. SL developed the Biopipeline® software for sequence comparison analysis, NP directed bioinformatic EST analysis, participated in local and global sequence comparison, HEH conceived the study and coordinated its development, and RH designed the construction of differential EST database, coordinated its analysis and drafted the manuscript. All authors read and approved the final manuscript.

#### Additional material

##### Additional File 1

A complete list of non-redundant EST, including GenBank accession number (AN) of the generated sequence and its corresponding GO identification number (GO ID), GO function definition inferred from sequence similarity, BLASTX top hit sequence and AN outcome from searches against SWALL is provided.

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