

## KAGIANA: An Excel-Based Tool for Retrieving Summary Information on *Arabidopsis* Genes

Yoshiyuki Ogata<sup>1</sup>, Nozomu Sakurai<sup>1</sup>, Koh Aoki<sup>1</sup>, Hideyuki Suzuki<sup>1</sup>, Koei Okazaki<sup>1</sup>, Kazuki Saito<sup>2</sup> and Daisuke Shibata<sup>1,\*</sup>

<sup>1</sup>Kazusa DNA Research Institute, Kazusa-Kamatari 2-6-7, Kisarazu, Chiba, 292-0818 Japan

<sup>2</sup>Graduate School of Pharmaceutical Science, Chiba University, Yayoi-cho 1-33, Inage-ku, Chiba, 263-8522 Japan

Various public databases provide *Arabidopsis* gene information via the internet. It is useful to abstract information obtained from such databases. We have developed the KAGIANA tool, which allows a user to retrieve summary information obtained from selective databases and to access pages for a gene of interest in those databases. The tool is based on Microsoft Excel and provides several macro programs for gene expression analyses. It can assist plant biologists in accessing omics information for plant biology. The KAGIANA tool is freely available at <http://pmnedo.kazusa.or.jp/kagiana/>.

**Keywords:** Annotation • *Arabidopsis* • Database • Gene expression • Omics.

**Abbreviations:** AGI, *Arabidopsis* Genome Initiative; GO, Gene Ontology; NCBI, The National Center for Biotechnology Information; TAIR, The *Arabidopsis* Information Resource.

Since the completion of the genome sequence of the model plant *Arabidopsis thaliana* (*Arabidopsis Genome Initiative* 2000), advances in genome and gene expression analysis have resulted in a vast number of data sets generated for *Arabidopsis*. Data sets of the *Arabidopsis* genome sequence are available at GenBank (Benson et al. 2008; <http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=Nucleotide>) and DDBJ (Sugawara et al. 2008; <http://www.ddbj.nig.ac.jp/>). In *Arabidopsis*, the sequence is separated into five chromosomes, which have 33,282 gene loci according to TAIR8 (Swarbreck et al. 2008; <http://www.arabidopsis.org/>). Amino acid sequences of proteins encoded by genes have been analyzed for various purposes. On the basis of localization signals included in such sequences, subcellular localization of proteins has been predicted using tools such as WoLF PSORT (Horton et al. 2007; <http://wolffpsort.org/>) and TargetP (Emanuelsson et al. 2007;

<http://www.cbs.dtu.dk/services/TargetP/>). Domain structures, which show functional properties in proteins, have been predicted on the basis of amino acid sequences using tools such as SCOP (Andreeva et al. 2008; <http://scop.berkeley.edu/>), and can be found in databases such as InterPro (Mulder et al. 2007; <http://www.ebi.ac.uk/interpro/>). Analysis of transmembrane domains has been performed using tools such as TMHMM (Moller et al. 2001; <http://www.cbs.dtu.dk/services/TMHMM/>) and that of hydrophathy can be found in databases such as SUBA (Heazlewood et al. 2007; <http://www.plantenergy.uwa.edu.au/suba2/>). Data sets of gene expression are available from databases such as the Gene Expression Omnibus (GEO) database (Barrett et al. 2006; <http://www.ncbi.nlm.nih.gov/geo/>). Several databases such as ATTED-II (Obayashi et al. 2007; <http://atted.jp/>) and Genevestigator (Zimmermann et al. 2004; <https://www.genevestigator.ethz.ch/>) provide a function to perform gene expression analysis. Using a vast number of gene expression data sets, approaches for detecting co-expressed genes, such as the ARACNE tool (Margolin et al. 2006; <http://amdec-bioinfo.cu-genome.org/html/ARACNE.htm>), the average clustering coefficient index (Gupta et al. 2006) and the Confeito algorithm (<http://pmnedo.kazusa.or.jp/kagiana/coexprocess/>) have been developed. On the basis of these analyses, molecular function, subcellular localization and biological processes of genes have finally been consistently assigned to 'molecular function', 'cellular component' and 'biological process', respectively, the three aspects of the Gene Ontology (GO) terminology (Gene Ontology Consortium 2008; <http://geneontology.org/>).

To obtain genomic and transcriptomic information on genes of interest, a user can visit these databases and access these tools via the internet or download them for personal

\*Corresponding author: E-mail, [shibata@kazusa.or.jp](mailto:shibata@kazusa.or.jp); Fax, +81-438-52-3948.

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use. However, to retrieve the information, users generally require knowledge of the omics information published in the databases; for example, how to select an adequate website and how to set an adequate threshold value such as the gene-to-gene correlation coefficient for acquiring data of interest in the website. For biological users, unfamiliar with omics analyses such as genomics and transcriptomics, it is useful to have access to abstracted gene information from such databases and analyses and to use quick links to these databases.

We have developed the KAGIANA (Kazusa *Arabidopsis* Gene Information And Network Analysis) tool to summarize various *Arabidopsis* omics analyses from the above-mentioned databases and tools, and to provide links to pages for genes of interest in the databases. The tool is based on Microsoft Excel (version 2003 or higher) and thus requires only enough skill for basic Excel operation. The implementation of this tool is verified using Windows XP or higher for PC, and OS X or higher for Macintosh. The macro programs of the tools are available only for Windows users as of November 2008. Our goal is to assist plant biologists in accessing information from omics analyses so that they can incorporate it into their plant biology research.

The KAGIANA tool is downloadable as a ZIP-format file at <http://pmnedo.kazusa.or.jp/kagiana/>. The KAGIANA tool is formatted as a Microsoft Excel workbook file, composed of five worksheets [one database sheet ('Data20080524'), two readme sheets ('ReadMe\_1st' and 'ReadMe\_Tools') and two retrieval sheets ('Selected\_Link' and 'Selected\_GO')] and one macro program ('Tools') comprising four analysis tools ('Confeito', 'GX bar chart', 'GO pie chart' and 'ATTED chart'). In KAGIANA, AGI codes (e.g. At1g01010) are used for the retrieval and performance of the tool.

The database sheet is composed of the following information for 33,362 loci (Fig. 1A), which was obtained from the TAIR database. First, the A to D columns represent AGI codes, a short description, description, and identifiers for NCBI, respectively. Secondly, the E to J columns display representative GO terms, which certainly accompany the evidence codes, and their Evidence Code categories, which are abbreviated as 'X' (experimental) for EXP, IDA, IPI, IMP, IGI and IEP; 'S' (statement) for TAS and IC; 'C' (computational) for ISS, ISO, ISA, ISM, IGC and RCA; 'L' (electronic) for IEA; and 'N' (not available) for NAS and ND, in the three aspects of GO terminology, i.e. cellular component (the E and F columns), molecular function (the G and H columns) and biological

**A**

	A	B	C	D	E	F	G	H	I	J	K	L	M	N	O	P
1	AGI Code	Short Description	Description	NCBI ID	Comp. Term	CS	Func. Term	FS	Proc. Term	PS	WoLF	WI	TgtP	SCOP	SI	#TD
2	At1g01010	ANAC001 (Arabidopsis NAC domain similar to ANAC001)	similar to ANAC001	839580	cellular_component_unkn	N	transcription factor activi	C	multicellular organismal d	C	cyto	2	O9	NAC domain ID101941	39	1
3	At1g01020	ARV1	similar to ARV1	839569	membrane GO:0016020	C	molecular_function_unkno	N	biological_process_unknov	N	#N/A	-	M6	#N/A	-	3
4	At1g01030	NGA3 (NGATHA3)	similar to NGA3	839321	cellular_component_unkn	N	transcription factor activi	C	leaf development GO:0048	X	nucL	8	O8	alpha-catenin/vinculin	1	0
5	At1g01040	DCL1 (DICER-LIKE1)	Identical to E	839574	nuclear dicing body GO:00	X	protein binding GO:00055	X	virus induced gene silenc	X	nucL	3	O8	RNase III catalytic domain	38	0
6	At1g01046	MIR838a	#N/A	6240410	#N/A	-	#N/A	-	#N/A	-	#N/A	-	#N/A	#N/A	-	-
7	At1g01050	ATPPA1 (ARABIDOPSIS THALIANA similar to ATI	similar to ATI	839579	nucleus GO:0005634	X	inorganic diphosphatase	X	metabolic process GO:000	C	cyto	8	M4	Inorganic pyrophosphatase	70	0
8	At1g01060	LHY (LATE ELONGATED HYPOCOT similar to CC	similar to CC	839341	#N/A	-	transcription factor activi	C	response to ethylene stir	X	nucL	10	O8	Homeodomain-like ID461	16	0
9	At1g01070	nodulin MtN21 family	similar to noc	839550	membrane GO:0016020	L	#N/A	-	#N/A	-	plas	2	O4	Multidrug resistance efflu	7	8
10	At1g01080	33 kDa ribonucleoprotein, chloroplast similar to CP	similar to CP	839463	chloroplast GO:0009507	L	RNA binding GO:0003723	C	biological_process_unknov	N	chlo	9	O9	RNA-binding domain, RBC	34	0
11	At1g01090	PDH-E1 ALPHA (PYRUVATE DEHYD similar to AT	similar to AT	839429	chloroplast GO:0009507	L	pyruvate dehydrogenase	C	metabolic process GO:000	C	chlo	9	O9	Thiamin diphosphate-bind	109	0
12	At1g01100	60S acidic ribosomal protein P1 (RP	Identical to 6	839410	ribosome GO:0005840	C	structural constituent of	C	translational elongation	G	cyto	5	O5	Ribosomal protein L7/12	2	0
13	At1g01110	IOD18 (IQ-domain 18)	similar to IQC	839394	mitochondrion GO:00057	L	molecular_function_unkno	N	#N/A	-	nucL	6	M9	P-loop containing nucleoi	4	0

**B**

	A	B	C	D	E	F	G	H	I	J	K	L	M	N	O	P	Q	R	S	T	U	V	W
1	Title:																						081022
2																							
3	AGI Code	? TA	EG	MP	IP	TD	SB	KG	GR	DN	SH	GA	GC	RV	eB	AV	AT	AC	KV	AP	Short Description		Description
4	At1g01010	TA	EG	MP	IP	TD	SB	KG	GR	DN	SH	GA	GC	RV	eB	AV	AT	AC	KV	AP	ANAC001 (Arabidopsis NAC domain containing protein 1)		similar to ANAC069 (Arabidopsis NAC domain containing protein 69), transcription factor [Arabidopsis thaliana] (TAIR:AT4G01550.1), similar to unnamed protein product [Vitis vinifera] (GB:CA062872.1), contains InterPro domain No apical meristem (NAM) protein; (InterPro:IPR003441)
5	At1g01020	TA	EG	MP	IP	TD	SB	KG	GR	DN	SH	GA	GC	RV	eB	AV	AT	AC	KV	AP	ARV1		similar to ARV2 [Arabidopsis thaliana] (TAIR:AT4G01510.1), similar to Os03g0784300 [Oryza sativa (japonica cultivar-group)] (GB:NP_001051472.1), similar to hypothetical protein [Oryza sativa] (GB:AAK72898.1), similar to hypothetical protein Os013313 [Oryza sativa (indica cultivar-group)] (GB:EAY92080.1), contains InterPro domain Arv1-like protein (InterPro:IPR007290)
6	At1g01030	TA	EG	MP	IP	TD	SB	KG	GR	DN	SH	GA	GC	RV	eB	AV	AT	AC	KV	AP	NGA3 (NGATHA3)		similar to NGA1 (NGATHA1), transcription factor [Arabidopsis thaliana] (TAIR:AT2G46870.1), similar to AT1G01030 [Arabidopsis lyrata subsp. petraea] (GB:AA72472.1), contains InterPro domain Transcription factor B3 (InterPro:IPR003340)

**C**

	A	B	C	D	E	F	G	H	I	J	K	L	M	N	
1	Title:														081022
2															
3	AGI code	Short Description	Cellular Component Term	CS	Molecular Function Term	FS	Biological Process Term	PS	WoLF	WI	TgtP	SCOP	SI	#TD	
4	At1g01010	ANAC001 (Arabidopsis NAC domain containing protein 1)	cellular_component_unknow	N	transcription factor activity	C	multicellular organismal development	C	cyto	2	O9	NAC domain ID101941	39	1	
5	At1g01020	ARV1	membrane GO:0016020	C	molecular_function_unknow	N	biological_process_unknow	N	#N/A	-	M6	#N/A	-	3	
6	At1g01030	NGA3 (NGATHA3)	cellular_component_unknow	N	transcription factor activity	C	leaf development GO:0048366	X	nucL	8	O8	alpha-catenin/vinculin ID:47220	1	0	

**Fig. 1** Composition of the KAGIANA worksheets. (A) The database sheet, including summary information of *Arabidopsis* genes obtained from the selected databases. (B) The sheet for hyperlinks to the selected public databases, as shown in Table 1. (C) The sheet for summary information of the selected omics analyses, i.e. GO terms and results from analyses of WoLF PSORT, TargetP, SCOP and TMHMM.

process (the I and J columns), respectively. A GO term was selected as the representative term for each aspect for a gene, according to the order of Evidence Code categories, i.e. X, S, C, L and N. The following columns represent information from the analytical tools. The K and L columns represent data from WoLF PSORT, which predicts the subcellular localization of proteins, and the reliability index, whose best score is 14, respectively. The M column represents information from TargetP, which also predicts subcellular localization, and the reliability index, ranging from 0 to 9 at the maximum. The N and O columns represent that from SCOP, which predicts domains of proteins, and the reliability index, which is the negative logarithm of the actual value, respectively. The P column represents TMHMM, which predicts the number of transmembrane domains of proteins.

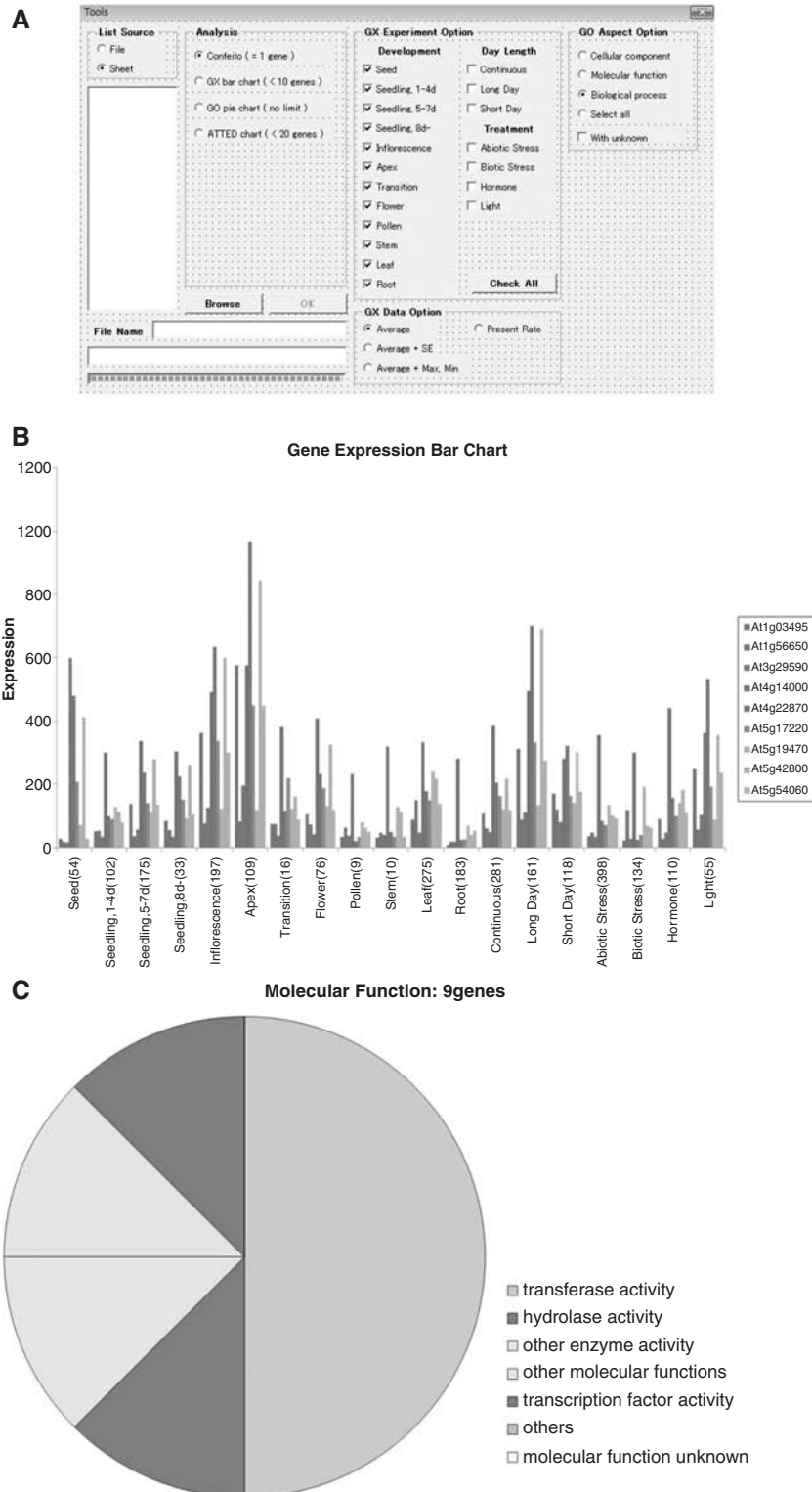
The 'Selected\_Link' sheet provides hyperlinks to 19 selected public databases for information retrieval of genes of interest and their Short Description and Description (Fig. 1B). These hyperlinks lead a user to the pages for individual genes in the individual databases by the following steps: (i) input AGI code(s) in the A column from the A4 cell to the lower cells (e.g. input 'At1g01010' in the A4 cell and 'At1g01020' in the A5 cell); (ii) select the range of the B4 to the W4 cells; and (iii) double-click the right lower corner (a black square) to copy the equations in the fourth row into the lower rows in the same columns (e.g. copy the B4-W4 into the B5-W5). Then, a user can access a database of interest from among the C to U columns (e.g. click the T4 cell for access to the page for the query gene in the KaPPA-View tool). The tool provides access to the databases shown in Table 1. The way to use this sheet is also described in the 'ReadMe\_1st' sheet.

In the 'Selected\_GO' sheet, a user can retrieve information on genes of interest from various omics analyses (Fig. 1C), i.e. the three GO term aspects, WoLF PSORT, TargetP, SCOP and TMHMM as mentioned above. Steps for retrieval are similar to those in the 'Selected\_Link' sheet. The terms in the third row are the same as those in the database sheet mentioned above, and the 'ReadMe\_1st' sheet has the explanation for such retrieval. By selecting the 'Selected\_Link' and the 'Selected\_GO' sheets, a user can manage to operate them simultaneously, e.g. when inputting AGI codes.

KAGIANA provides 'Tools' macro programs including the four analyses (Fig. 2A), i.e. including 'Confeito', 'GX bar chart', 'GO pie chart' and 'ATTED chart'. The 'Confeito' tool allows a user to extract co-expressed genes using the Confeito algorithm on the basis of a co-expression network approach (<http://pmnedo.kazusa.or.jp/kagiana/coexpress/>). The way to use the tools is described in the 'ReadMe\_Tools' sheet. The 'GX bar chart' tool allows a user to depict bar charts of gene expression profiles for multiple genes of interest (Fig. 2B). Bar charts are depicted using 1,245 DNA microarray data from the AtGenExpress project, which are available at <http://www.weigelworld.org/resources/microarray/AtGenExpress/>. The 'GO pie chart' tool allows a user to depict a pie chart of the distribution of GO-SLIM terms associated with multiple genes of interest (Fig. 2C). GO-SLIM terms are available at the TAIR database. This tool counts all multiple GO-SLIM terms assigned to a gene. For this version of KAGIANA, such terms were obtained at May 2008. The 'ATTED chart' tool helps users download the charts of AtGenExpress gene expression profiles for individual genes from the ATTED database onto a worksheet in KAGIANA per gene.

**Table 1** Abstract of databases linkable from the KAGIANA tool

Abbreviation	Databases	URL
TA	TAIR (Swarbreck et al. 2008)	<a href="http://www.arabidopsis.org/">http://www.arabidopsis.org/</a>
EG	Entrez Gene (Maglott et al. 2005)	<a href="http://www.ncbi.nlm.nih.gov/sites/entrez?db=gene">http://www.ncbi.nlm.nih.gov/sites/entrez?db=gene</a>
MP	MPSS (Brenner et al. 2000)	<a href="http://mpss.udel.edu/at/">http://mpss.udel.edu/at/</a>
IP	InParanoid (Berglund et al. 2008)	<a href="http://inparanoid.sbc.su.se/cgi-bin/index.cgi">http://inparanoid.sbc.su.se/cgi-bin/index.cgi</a>
TD	SIGnAL (Alonso et al. 2003)	<a href="http://signal.salk.edu/cgi-bin/tdnaexpress">http://signal.salk.edu/cgi-bin/tdnaexpress</a>
SB	SUBA (Heazlewood et al. 2007)	<a href="http://www.plantenergy.uwa.edu.au/applications/suba2/">http://www.plantenergy.uwa.edu.au/applications/suba2/</a>
KG	KEGG (Kanehisa et al. 2008)	<a href="http://www.genome.jp/kegg/">http://www.genome.jp/kegg/</a>
GR	GRAMENE (Liang et al. 2008)	<a href="http://www.gramene.org/">http://www.gramene.org/</a>
DN	NASCArrays Digital Northern (Craigon et al. 2004)	<a href="http://affymetrix.arabidopsis.info/narrays/digitalnorthern.pl">http://affymetrix.arabidopsis.info/narrays/digitalnorthern.pl</a>
SH	NASCArrays Spot History (Craigon et al. 2004)	<a href="http://affymetrix.arabidopsis.info/narrays/spothistory.pl">http://affymetrix.arabidopsis.info/narrays/spothistory.pl</a>
GA	Geneinvestigator Gene Atlas (Zimmermann et al. 2004)	<a href="https://www.geneinvestigator.ethz.ch/gv/index.jsp">https://www.geneinvestigator.ethz.ch/gv/index.jsp</a>
GC	Geneinvestigator Gene Chronologer (Zimmermann et al. 2004)	<a href="https://www.geneinvestigator.ethz.ch/gv/index.jsp">https://www.geneinvestigator.ethz.ch/gv/index.jsp</a>
RV	Geneinvestigator Response Viewer (Zimmermann et al. 2004)	<a href="https://www.geneinvestigator.ethz.ch/gv/index.jsp">https://www.geneinvestigator.ethz.ch/gv/index.jsp</a>
eB	eFP Browser (Winter et al. 2007)	<a href="http://bbc.botany.utoronto.ca/efp/cgi-bin/efpWeb.cgi">http://bbc.botany.utoronto.ca/efp/cgi-bin/efpWeb.cgi</a>
AV	AtGenExpress Visualization Tool	<a href="http://jsp.weigelworld.org/expviz/expviz.jsp">http://jsp.weigelworld.org/expviz/expviz.jsp</a>
AT	ATTED-II (Obayashi et al. 2007)	<a href="http://atted.jp/">http://atted.jp/</a>
AC	AraCyc (Zhang et al. 2005)	<a href="http://www.arabidopsis.org/biocyc/index.jsp">http://www.arabidopsis.org/biocyc/index.jsp</a>
KV	KaPPA-View 3 (Sakurai and Shibata 2006)	<a href="http://kpv.kazusa.or.jp/kappa-view3/">http://kpv.kazusa.or.jp/kappa-view3/</a>
AP	AtProteome (Baerenfaller et al. 2008)	<a href="http://fgcz-atproteome.unizh.ch/">http://fgcz-atproteome.unizh.ch/</a>



**Fig. 2** Composition of the KAGIANA tools. (A) The window of the 'Tools' macro program, including analyses of 'Confeito', 'GX bar chart', 'GO pie chart' and 'ATTED chart'. (B) The result from the 'GX bar chart' analysis. (C) The result from the 'GO pie chart' analysis, showing the aspect of molecular function.



Detailed steps for using these tools are described in the 'ReadMe\_Tools' sheet in the KAGIANA workbook. Briefly, the steps are (i) click 'Tools' in the menu bar; (ii) select 'Macro' and click 'Macros'; (iii) select 'Tools' in the macro box and click 'Execute' (open the 'Tools' window); (iv) select a tool in the 'Analysis' frame in the window; (v) input AGI codes into different lines in the textbox left of the frame; (vi) select the option frame when selecting 'GX bar chart' and 'GO pie chart' tools; and then (vii) click the 'OK' button if the character color on the button is black (otherwise, there is insufficient information for retrieval).

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