Set 3B -	
coleoptilar not	
cotyledonary -	Set 4 -

			cotyledonary -	Set 4 -
GO Term	Onto	Description	arab. IDs	cotyledonary
GO:0005975	Р	carbohydrate metabolic process	0.0016	0.0000076
GO:0009719	Р	response to endogenous stimulus	0.002	6.6E-11
GO:0009056	Р	catabolic process	0.002	0.00000014
GO:0006810	Р	transport	0.01	1.6E-17
GO:0051234	Р	establishment of localization	0.01	9.9E-18
GO:0051179	Р	localization	0.011	2.6E-18
GO:0009856	Р	pollination	0.05	
GO:0003824	F	catalytic activity	0.000055	2E-55
GO:0005215	F -	transporter activity	0.0011	0.0000004
GO:0016788	F	hydrolase activity, acting on ester bonds	0.012	0.0000018
GO:0005886	С	plasma membrane	0.013	2E-22
GO:0005576	С	extracellular region	0.013	
GO:0030312	C	external encapsulating structure	0.032	0.007
GO:0005618	C	cell wall	0.032	0.007
GO:0016020	С	membrane	0.037	4.6E-44
GO:0009987	P	cellular process		3.1E-77
GO:0044237 GO:0008152	P P	cellular metabolic process		6.2E-50
GO:0008132 GO:0044238	P P	metabolic process		2E-46 2.9E-37
GO:0044238 GO:0016043	P	primary metabolic process		1.3E-30
GO:0016043 GO:0050896	P P	cellular component organization response to stimulus		5.1E-28
GO:0009628	P	response to abiotic stimulus		9.7E-28
GO:0009028	P	nucleobase-containing compound metabolic process		1.2E-27
GO:0050789	r P	regulation of biological process		8.4E-27
GO:0030703	P	developmental process		1E-26
GO:0032302 GO:0048856	P	anatomical structure development		2.9E-25
GO:0050794	Р	regulation of cellular process		3.5E-23
GO:0007275	P	multicellular organism development		4.5E-23
GO:0032501	P	multicellular organismal process		5.9E-23
GO:0009791	Р	post-embryonic development		1.6E-21
GO:0043412	Р	macromolecule modification		2.9E-19
GO:0044260	Р	cellular macromolecule metabolic process		5E-19
GO:0043170	Р	macromolecule metabolic process		5.5E-19
GO:0048608	Р	reproductive structure development		1.2E-17
GO:0065007	Р	biological regulation		4.3E-17
GO:0006464	Р	cellular protein modification process		5.4E-17
GO:0000003	Р	reproduction		1.7E-16
GO:0006807	Р	nitrogen compound metabolic process		2E-16
GO:0022414	Р	reproductive process		2.9E-16
GO:0003006	Р	developmental process involved in reproduction		4.2E-15
GO:0019222	Р	regulation of metabolic process		5.3E-12
GO:0006950	Р	response to stress		2.6E-11
GO:0044249	Р	cellular biosynthetic process		6.6E-11
GO:0009058	Р	biosynthetic process		1.7E-10
GO:0006259	P	DNA metabolic process		2.1E-10
GO:0009790	P	embryo development		8.3E-10
GO:0006091	P	generation of precursor metabolites and energy		8.3E-10
GO:0015979	P	photosynthesis		8.5E-10
GO:0007049	P	cell cycle		8.5E-10
GO:0060255	P	regulation of macromolecule metabolic process		4.3E-09
GO:0009908	P	flower development		5.8E-09 0.000000007
GO:0009653	P D	anatomical structure morphogenesis		0.00000007
GO:0007165 GO:0007154	P D	signal transduction cell communication		0.0000003
GO:0007154 GO:0010468	P P	regulation of gene expression		0.00000023
GO:0010468 GO:0006629	P P	lipid metabolic process		0.0000072
GO:0000029	P	protein metabolic process		0.000088
GO:0019338	P	growth		0.000098
GO:0040007	P	cellular protein metabolic process		0.00027
GO:0044869	P	cellular developmental process		0.00027
GO:0042592	P	homeostatic process		0.00032
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GO:0019725	P	cellular homeostasis	 0.0013
GO:0030154	P	cell differentiation	 0.019
GO:0008219	Р	cell death	 0.02
GO:0016049	Р	cell growth	 0.021
GO:0010467	Р	gene expression	 0.035
GO:0005515	F	protein binding	 4.3E-57
GO:0005488	F	binding	 7E-56
GO:0000166	F	nucleotide binding	 3.8E-29
GO:0016787	F	hydrolase activity	 1.3E-17
GO:0016740	F	transferase activity	 3.5E-17
GO:0016817	F	hydrolase activity, acting on acid anhydrides	 1.8E-11
		hydrolase activity, acting on acid anhydrides, in phosphorus-	
GO:0016818	F	containing anhydrides	 1.9E-11
GO:0016462	F	pyrophosphatase activity	 2.4E-11
GO:0017111	F	nucleoside-triphosphatase activity	 3.3E-10
GO:0016772	F	transferase activity, transferring phosphorus-containing groups	 0.000000072
GO:0005198	F	structural molecule activity	 0.00000053
GO:0003677	F	DNA binding	 0.000001
GO:0016301	F	kinase activity	 0.0000032
GO:0004518	F	nuclease activity	 0.00034
GO:0003676	F	nucleic acid binding	 0.0021
GO:0003774	F	motor activity	 0.013
GO:0008135	F	translation factor activity, RNA binding	 0.017
GO:0060089	F	molecular transducer activity	 0.019
GO:0004872	F	receptor activity	 0.019
GO:0003682	F	chromatin binding	 0.021
GO:0004871	F	signal transducer activity	 0.036
GO:0005623	С	cell	 4.3E-187
GO:0044464	С	cell part	 5.1E-187
GO:0005622	С	intracellular	 1.7E-144
GO:0044424	С	intracellular part	 1.8E-143
GO:0044422	С	organelle part	 1.4E-128
GO:0044446	С	intracellular organelle part	 1.8E-128
GO:0005737	С	cytoplasm	 5.5E-111
GO:0043226	С	organelle	 6.7E-111
GO:0043229	С	intracellular organelle	 7.4E-111
GO:0043227	С	membrane-bounded organelle	 8.9E-101
GO:0043231	С	intracellular membrane-bounded organelle	 1.4E-100
GO:0044444	С	cytoplasmic part	 4.8E-96
GO:0009536	С	plastid	 3.1E-59
GO:0032991	С	macromolecular complex	 3E-48
GO:0005829	С	cytosol	 4.8E-47
GO:0031975	С	envelope	 2.8E-32
GO:0031967	С	organelle envelope	 2.8E-32
GO:0012505	С	endomembrane system	 3.4E-28
GO:0043232	C	intracellular non-membrane-bounded organelle	 7.6E-28
GO:0043228	C	non-membrane-bounded organelle	 7.6E-28
GO:0005773	C	vacuole	 6.1E-27
GO:0031974	C	membrane-enclosed lumen	 5.2E-25
GO:0043233	С	organelle lumen	 5.2E-25
GO:0070013	С	intracellular organelle lumen	 5.2E-25
GO:0044428	C	nuclear part	 6.7E-25
GO:0009579	C	thylakoid	 2.5E-22
GO:0031981	C	nuclear lumen	 1.1E-20
GO:0005634	C	nucleus	 5.8E-19
GO:0005794	C	Golgi apparatus	 1.7E-17
GO:0005783	C	endoplasmic reticulum	 2.3E-13
GO:0005768	C	endosome	 2.7E-11
GO:0005730	C	nucleolus	 1.2E-10
GO:0003730	C	intracellular ribonucleoprotein complex	 2.6E-10
GO:0030329 GO:0005654	C	nucleoplasm	 3.3E-09
GO:0005840	C	ribosome	 0.000000049
GO:0005856	C	cytoskeleton	 0.000000049
GO:0005830	C	peroxisome	 0.0000025
GO:0003777 GO:0042579	C	microbody	 0.00015
GO:0042379 GO:0005635	C	nuclear envelope	 0.00013
30.0003033	C	nacion chivolope	3.00077