# Gene Comparison Between Arabidopsis thaliana, Prunus mume and Prunus persica 

Joanna M. Cross<br>Department of Horticultural Science, Faculty of Agriculture, Inonu University (Battalgazi Campus), 44000 Battalgazi, Malatya, Turkey


#### Abstract

Apricot (Prunus armeniaca) is a stone fruit consumed fresh or dried. Turkey is the top world producer with the Malatya area supplying over half of the crop. Market demands are stringent as apricot trees need to be resistant to both heat and cold while producing fruits satisfying the customer. Therefore, crop improvement involves many parameters. The genomes of peach (Prunus persica) and Japanese apricot (Prunus mume) have been sequenced. Both belong to the same family as apricots. Consequently, around 250 genes were collected for both species along with the reference Arabidopsis. Both targeted and non targeted approaches were applied to diversify the range of protein functions covered. Thus, a set of genes involved in amino acid metabolism was studied along with a second group selected based on the phenotype conferred. Comparison of the three plants shows that gene allocation to a given function is conserved assignment of a clear gene to gene correspondence between organisms is a delicate task and clear gene counterparts do not necessarily share the same cellular compartment.


Key words: Prunus, Arabidopsis, gene comparison, cold response, amino acid synthesis, protein localization

## INTRODUCTION

Apricot (Prunus armeniaca) is a stone fruit from the same family (Rosaceae) and genus (Prunus) as plum, peach, cherry and almond. It is a diploid with a relatively small genome size of 590 Mb (Hagen et al., 2002). Trees are cultivated in the Mediterranean Basin, the former Soviet Union countries, Iran, China, South Africa and the United States (Asthma, 2007). Turkey is the top world producer. The Malatya region ( $38^{\circ} 21^{\prime} \mathrm{N} / 38^{\circ} 17^{\prime} \mathrm{E}$ ) contributes over half of the country's apricots with its 10 million trees.

Trees bloom in the middle of March while leaves develop a month after flowering. As a result, the first stages of fruit development rely on reserves accumulated the previous season. Apricot is a climacteric fruit which means that ripening is accompanied by an ethylene burst. Fruits mature at different periods depending on the variety grown. Early ripening fruits are harvested in June while mid and late ripening cultivars are picked in July and end of August, respectively. Hence, harvest season spreads over several months.

Temperatures drop below freezing during the Winter while reaching $40^{\circ} \mathrm{C}$ in the Summer. Consequently, trees have to be hardy to both cold and heat. In addition, the species is sensitive to diseases with Sharka (Plum Pox Virus) being the major threat (Sochor et al., 2012). So far,
the disease is absent from the Malatya region though present in Turkey. However, night frosts occur until late April. They are particularly damaging in March at the time of flowering. For instance, in 2014, 3 days of cold, end of March, destroyed $90 \%$ of the crop. Since, Spring frosts occur roughly every 3 years, apricot production is quite irregular.

Apricots are appreciated for their taste. Fruit quality depends on sugar and organic acid content (Drogoudi et al., 2008). In addition, over 200 volatile compounds have been identified (Gonzalez-Aguero et al., 2009). Panel studies have shown that a combination of 18 compounds can mimic apricot taste (Greger and Schieberle, 2007). Apricots harbor high levels of antioxidants including vitamin A. The molecules are suggested to have therapeutic effects on a variety of pathological conditions such as cancer, diabetes and neurodegenerative or cardiovascular diseases. Consequently, there is an interest in identifying apricot varieties with enhanced antioxidant content (Ruiz et al., 2005a, b).

Production of new apricot varieties demands a careful balance between farmer satisfaction and customer approval. A regular and optimal ratio between yield and size must be ensured while taste and nutritional content are preserved. Consequently, parameters of importance span a large number of reactions as well as various
signaling pathways. So far, efforts have mainly focused on assessing genetic variation with Random Amplified Polymorphic DNA (Takeda et al., 1998), Amplified Fragment Length Polymorphism (Hagen et al., 2002; Yuan et al., 2007; Krichen et al., 2008), Inter-Simple Sequence Repeat markers (Yilmaz et al., 2009) and Sequence Characterized Amplified Regions. The results are used for conservation efforts and variety classification (Hagen et al., 2002; Bourguiba et al., 2012), quantitative trait locus identification (Salazar et al., 2014) and as an information basis to breed new varieties (Asthma, 2012). Efforts are under way to develop cultivars with cold and/or Sharka resistance.

At the molecular level, genes differentially expressed during fruit maturation were identified (Geuna et al., 2005). Moreover, correlations were made between volatile composition and expression patterns of genes involved in aroma synthesis (Gonzalez-Aguero et al., 2009). Recently, the species entered the omics field. Indeed, transcriptional analysis of fruit maturation was performed by collecting expressed sequenced tags (Grimplet et al., 2005), designing expressed sequenced tag based microarrays (Li et al., 2012) and using chips constructed for peach (Manganaris et al., 2011). The data were complemented by a proteomic time-course of fruit development. Moreover, the BGI lists a sequencing project in its early phase. Meanwhile, the genomes of peach (Prunus persica) and of Japanese apricot (Prunus mume) have been sequenced (Verde et al., 2013) and the pathways of peach collected into PeachCyc (Jung et al., 2014).

Unfortunately, resources are still incomplete for apricot mainly due to the lack of a sequenced genome. Therefore, it would be beneficial to use data from other species. A logical first choice consists of the two closely related species Prunus persica and Prunus mume. Those are related to the extent that microarrays designed for Prumus persica or Prunus mume also function for apricots (Manganaris et al., 2011; Li et al., 2012). As mentioned, all three species are classified in the genus Prunus. However, both Prunus mume and Prunus armeniaca belong to the same subgenus Prunus and section Armeniaca while exhibiting similar fruit and tree morphology. Nonetheless, markers clearly separate each plant into different species (Hagen et al., 2002). On the other hand, Prunus persica is either classified in a separate subgenus Amygdalus or listed in the subgenus Prunus albeit in a separate section from apricot (Shi et al., 2013).

The model plant Arabidopsis represents an appealing second resource with its wealth of experimental data. Thus, a subset of genes was compared between Prunus persica, Prunus mume and Arabidopsis to assess
cautions required when translating data from one organism to another. Genes were first assembled for a specific function, i.e., amino acid metabolism. Second, databases of Arabidopsis mutants were screened to identify genes based on phenotype. This ensured that a diversity of gene functions was collected.

## MATERIALS AND METHODS

Selection of the amino acid metabolism pathways: The curated pathways Aracyc12.0 (http://pmn.plantcyc.org/) (Mueller et al., 2003) and Peachcycl.0 (http://www. rosaceae.org/) (Jung et al., 2014) were used to collect genes for Prunus persica and Arabidopsis. They were checked for consistency and redundancies. On the whole, the two databases are consistent except in the following cases. Regarding methionine recycling, Peachcyc harbors two versions of the S-adenosyl-Lmethionine cycle while Aracyc contains one version in addition to the Yang cycle. The latter salvages the sulfur of 5 '-Methylthioadenosine (MTA) synthesized during the production of ethylene (Sauter et al., 2013) and therefore seems quite relevant to developing apricot fruits. Effectively, combination of the early steps of ethylene biosynthesis with the methionine salvage route of Peachcyc produced the Arabidopsis Yang cycle. Moreover, Peachcyc lists two conversions of adenosyl-homocysteine to homocysteine, one direct, one through ribosylhomocysteine. However, the Arabidopsis counterparts of the annotated peach genes potentially converting ribosylhomocysteine were shown to catalyze a Yang cycle reaction with poor or no affinity for their hypothesized substrate (Siu et al., 2008). Hence, only the direct conversion of adenosyl-homocysteine to homocysteine was considered.

Regarding cysteine biosynthesis, Peachcyc lists an interconversion route between homocysteine and cysteine which is absent from Aracyc. However, the conversion of homocysteine to cysteine with cystathionine beta synthase seems specific to mammals while plants synthesize homocysteine and then methionine from cysteine (Kushwaha et al., 2009). Hence, the interconversion route was not considered.

Finally, Peachcyc lists two routes to convert prephenate to phenylalanine, one through arogenate, another via phenylpyruvate. However, the arogenate pathway appears to be the major route with the phenylpyruvate bypass potentially having a minor role (Tzin and Galili, 2010). Therefore, both pathways were tentatively kept.

Selection of the genes involved in each reaction: Genes were collected as listed in Aracyc and Peachcyc. The final list was determined based on literature, sequence alignments and annotations. Indeed, Arabidopsis genes were checked in TAIR (http://www.arabidopsis.org/) Lamesch et al., 2012) for experimental evidence regarding their function and localization. They were also blasted (Altschul et al., 1990) against a Prunus + Arabidopsis thaliana database to verify that the Prunus persica counterparts were selected correctly and to identify Prunus mume homologues. Peach genes with no Arabidopsis homologues were blasted against the non-redundant database to validate the annotation. All blastings were performed with nBLAST using default settings (Match/Mismatch Scores, 2/-3; Gap costs, existence 5 , extension 2 ). Reactions were associated with a loci rather than a mRNA. Hence, alternative splicing variants were not considered.

Selection of genes based on phenotype: Phenotypes were screened in the Chloroplast 2010 (Ajjawi et al., 2010; Lu et al., 2008; Lu et al., 2011), the RIKEN Phenome (Kuromori et al., 2004, 2006) and the Arabidopsis Stress Responsive (Borkotoky et al., 2013) databases. The Chloroplast 2010 mutants were screened for altered C/N ratio in seeds or modified starch levels in leaves using default $z$ scores and the option "two siblings with altered parameter". Moreover, hits were checked for consistency of results within two different knock-out lines. Morphological parameters were examined to eliminate plants terminally diseased. The RIKEN Phenome Project was screened for smaller siliques and decreased yields. Plants noted as sterile or small were eliminated to ensure that defects were due to silique development and not to growth problems. The Arabidopsis stress database was screened for cold responses. The literature supplied with each hit was used to identify potential signaling partners. Both metabolic and phenotype base genes were run through the Stanford Interactome database (Jones et al., 2014) selecting partners identified in two primary and two confirmation screens.

Gene to gene correspondence assignment: Arabidopsis genes identified in the phenotype based search were blasted as described above against a Prunus + Arabidopsis thaliana database. Next, gene to gene correspondence was determined relatively. All genes which aligned over more than a conservative $16 \%$ of the target were selected. The DNA sequences of Prunus mume, Prunus persica and Arabidopsis were then aligned using Clustal Omega (Goujon et al., 2010;

Sievers et al., 2011) and default parameters. The identity matrixes and alignments were used to assign the correct Prunus homologues to their Arabidopsis counterparts. Those were considered to be genes aligning preferentially and solely with the Arabidopsis target. Next, Prunus persica genes were blasted against Arabidopsis to check that they aligned preferentially with their selected homologue.

Localization: Reactions were assigned to a compartment based on previous literature, TAIR annotations, protein sequence alignment and targeting prediction programs. The prediction programs WoLF PSORT and TargetP (Emanuelsson et al., 2007) were used to sort out reactions with multiple cell locations. WoLF PSORT aligns a target with proteins of known compartmentalization while TargetP specializes in detecting mitochondrial, chloroplasticand secretory targeting sequences. Therefore, both approaches are complementary. Default parameters were used with the setting on "plant sequences". NucPred (Brameier et al., 2007) and PredPlantPTS1 (Reumann et al., 2012) were used to discriminate nuclear and peroxisomal targeted proteins, respectively.

Statistics and probabilities: All calculations were performed in R Version 3.1.2 for Windows 64 bit with the packages gmp, lpSolveAPI and Xnomial. The number of genes per reaction was viewed as a distribution problem of a determined number of isozymes between different functions. The number of possible ways of splitting $n$ objects into $k$ groups of $n_{1}, n_{2}, n_{k}$ elements is given by $n!/ n_{1}!n_{2}!\ldots n_{k}!$. This was calculated for observed gene distributions in all three species with package gmp. The latter handles very large integers and decimals. Moreover, it expresses decimals as quotients thus enabling high precision calculations. Therefore, all calculations were done on quotients with a number obtained at the last step for rounding.

Next, it was sought to express results on a relative basis for comparison purposes. A first choice was the Stirling number of second kind which measures the number of ways of splitting n objects into k groups with at least one element. Unfortunately, this could not be calculated reliably due to the size of n ranging from 193-233. Therefore all results were expressed relative to the most frequent distribution in each species. The most frequent combination is obtained by minimizing the denominator in $n!/ n_{1}!n_{2}!\ldots n_{k}!$. Moreover, the latter is smaller with reduced $n$ values. Therefore, the assumption was made that a solution could be found with all groups
harboring 1-6 genes. Thus, the denominator could be expressed as $(1!)^{81} \times(2!)^{22} \times(3!)^{x^{3}} \times(4!)^{24} \times(5!)^{25} \times(6!)^{26}$. As $\log _{10}(\mathrm{x})$ varies in the same direction as x the $\log _{10}$ form was used, i.e., $x_{1} \times \log _{10}(1!)+x_{2} \times \log _{10}(2!)+x_{3} \times \log _{10}(3!)+x_{4} \times$ $\log _{10}(4!)+x_{5} \times \log _{10}(5!)+\mathrm{X}_{6} \times \log _{10}(6!)$.

The function was minimized with package lpSolveAPI using constraints $\sum \mathrm{x}_{\mathrm{i}}=103$ and $\sum \mathrm{ixx}_{\mathrm{i}}=193$ (Prunus mume), 201 (Prunus persica) and 233 (Arabidopsis) with all $\mathrm{x}_{\mathrm{i}}$ being integers between 0 and 103 . Identical results were obtained by replacing $\log _{10}(1)$ by a small value. In addition, a few simple solutions were confirmed by manual checking. Additional constraints were added to determine the maximum distributions for a given number of reactions with 2 genes or when the observed number of reactions with $>3$ genes is kept.

A given random gene distribution was calculated for two species using the observed values. Next, categories were reduced to reactions with 0 or $1,2,3$ and $>3$ genes. The small sample size precluded the use of a $\chi^{2}$-test to determine whether the gene distribution between two species was random or not. Hence, a Multinomial Goodness of Fit test was performed with package XNomial. Unfortunately, the Full Enumeration Method (xmulti) required examining $10^{18}$ possible combinations. Therefore, Monte Carlo simulations were used (xmonte) on 10000 trials. Simulations with 100000 or even 1000000 trials produced more precise results albeit at a reduced speed. Since, the outcome of the test was identical all calculations were performed based on 10000 trials. Moreover, results were similar for a log-likelyhood ratio, probability or even a $\chi^{2}$-test.

## RESULTS

Genes were collected for Arabidopsis thaliana, Prunus mume and Prunus persica using both a targeted and non targeted approach: Gene information was gathered for Arabidopsis thaliana, Prunus mume and Prunus persica because Prunu smume is the closest species to apricot with a sequenced genome but Prunus persica proteins have been curated into pathways and Arabidopsis thaliana harbors the most experimental evidence regarding protein function, localization and mutation effects. Moreover, both targeted and non-targeted approaches were used. First, genes encoding proteins involved in amino acid metabolism were collected. The latter are precursors for the synthesis of ethylene, volatile and phenolic compounds. Transcriptomic studies showed significant variations in different biosynthetic pathways (Li et al.,

2012; Manganaris et al., 2011). Both Aracyc12.0 (http://pmn.plantcyc.org/) (Mueller et al., 2003) and Peachcyc1.0 (http://www.rosaceae.org/) (Jung et al., 2014) were screened and reactions assigned on a metabolic rather than on a gene basis. Hence, catalytic steps with enzymatic evidence are listed even in the absence of corresponding genes. Moreover, several reactions correspond to the same gene list for bifunctional proteins. Enzymes are also placed in their compartments which produces further duplications for multiple locations.

The targeted approach has the drawback of limiting the selected genes to those encoding enzymes. Therefore, a phenotype based search was also completed to identify transcripts important for the early stages of fruit development. The period stands out in several ways. First, as mentioned cold frost damages flowers. In addition, fruits rely on reserves accumulated by the tree the previous season until leaves develop. Hence, three types of phenotypes were searched: those related to cold responses, those suggesting a deficiency in reserve accumulation or metabolic balance and those implicating poor fruit growth.

Several databases were screened to that effect. The Chloroplast 2010 Project (Ajjawi et al., 2010; Lu et al., 2008, 2011) measures a set of metabolic parametersin Arabidopsis knock-out mutants. It has the advantage of providing data for two different knock-outs for many genes but as the name implies focuses on chloroplastic proteins. Mutants were identified with consistent alterations in morning starch levels of leaves or in seed C/N ratio. The RIKEN Phenome Project morphologically characterizes Arabidopsis mutants obtained by transposon insertion (Kuromori et al., 2004, 2006). Mutant with smaller siliques or reduced yield were identified. The Arabidopsis Stress Responsive Database curates genes involved in abiotic stress (Borkotoky et al., 2013). Consequently, the database was screened for cold response genes. Some of the references provided listed proteins interacting with the curated product. They were included. Finally, potential interaction partners were searched for all genes in the Stanford Interactome Database (Jones et al., 2014). The project uses the split ubiquitin system to identify interactions between all Arabidopsis membrane proteins.

The Prunus species show a high conservation of isozyme numbers per reaction: Table 1 and 2 give a summary of the genes collected with literature used. In total, 133 reactions and 48 phenotypic genes were studied. This corresponded to 290 Arabidopsis genes or roughly 250

Table 1: Metabolic reactions curated for Arabidopsis thaliona (At), Prunus persica ( Pp ) and Prunus mume ( Pm )

| Pathway | Rx | Nb of genes |  |  | Genes perreaction |  |  | Local | References |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | At | Pp | Pm | At | Pp | Pm |  |  |
| Leu/Val/Ile | 19 (10) | 23 | 16 | 19 | 2.3 | 1.6 | 1.9 | Mostly Cl | Binder |
| Lys/Thr/Met | 14 (13) | 26 | 22 | 22 | 2.0 | 1.7 | 1.7 | Mostly Cl | Ravanel et al. (1998) |
|  |  |  |  |  |  |  |  |  | Ravanel et al. (2004) <br> Jander and Joshi (2009) |
| Met salvage | 13 (10) | 28 | 25 | 24 | 2.8 | 2.5 | 2.4 | Cy | Ravanel et al. (1998) |
|  |  |  |  |  |  |  |  |  | Sauter et al. (2013) |
| Phe/Tyr/Trp | 21 (16) | 55 | 41 | 38 | 3.4 | 2.6 | 2.4 | Cl | Tzin and Galili (2010) |
| Ser/Gly/Ala | 20 (15) | 38 | 33 | 31 | 2.5 | 2.2 | 2.1 | $\mathrm{Cl}, \mathrm{Mt}, \mathrm{Pe}, \mathrm{Cy}, \mathrm{Nu}$ | Couturier et al. (2013) |
|  |  |  |  |  |  |  |  |  | Liepman and Olsen (2003) |
|  |  |  |  |  |  |  |  |  | Ros et al. (2013) |
| Arg | 10 (9) | 13 | 17 | 16 | 1.4 | 1.9 | 1.8 | $\mathrm{Cl}, \mathrm{Cy}$ | Slocum (2005) |
| Asn/Asp/Glu/Gln | 10 (9) | 24 | 19 | 20 | 2.7 | 2.1 | 2.2 | $\mathrm{Cl}, \mathrm{Mt}, \mathrm{Cy},(\mathrm{Pe})$ | Coruzzi |
|  |  |  |  |  |  |  |  |  | Liepman and Olsen (2004) |
| Cys | 6 (6) | 14 | 11 | 11 | 2.3 | 1.8 | 1.8 | $\mathrm{Cl}, \mathrm{Mt}, \mathrm{Cy}$ | Novero (2009) |
| His | 10 (8) | 11 | 12 | 10 | 1.4 | 1.5 | 1.3 | Cl | Rajani |
| Pro | 10 (6) | 9 | 12 | 9 | 1.5 | 2.0 | 1.5 | $\mathrm{Cl}, \mathrm{Mt}, \mathrm{Cy}$ | Szabados and Savoure (2010) |
| Total | 133 (102) | 241 | 208 | 200 | 2.4 | 2.0 | 2.0 |  |  |

The second column (Rx) provides the total number of reactions for the pathways listed in the first column and in parenthesis the number of stepss with unique genes. Total number of genes involved, number of genes per reaction, pathway localization (Local) and references are provided for each metabolic group. Localization abbreviations are Cl: Chloroplast; Cy: Cytoplasm; Mt: Mitochondria; Nu: Nucleus; Pe: Peroxisome

Table 2: Genes collected based on phenotype conferred. Information was gathered from the Chloroplast 2010 project (Cl2010, Ajjawi et al., 2010; Lu et al., 2008, 2011), the RIKEN Phenome collection (RIKEN, Kuromori et al., 2004, 2006), the Stress Responsive Database (Stress, Borkotoky et al., 2013) and the Stanford Interactomeproject (Interactome, Jones et al., 2014). The table lists the number of genes associated with a given phenotype, the type of proteins encoded with their numbers in parenthesis and in the last column the number of homologues found in Prunus. Unambiguous Prunus counterparts are specified in parenthesis

| Database | Phenotype | Genes in At | Proteins encoded | Hits in Pp and Pm |  |
| :--- | :--- | :---: | :--- | :--- | :--- |
| Cl2010 | Excess starch am | 3 | Catalysis (2), kinase/phosphatase (1) | $3(3)$ |  |
| Cl2010 | Altered C/N | 1 | Interactions (1) | $11(1)$ |  |
| RIKEN | Low yield | 3 | Catalysis (1), interactions (1), transport (1) | $2(2)$ |  |
| RIKEN | Short siliques | 7 | Catalysis (2), interactions (1), transcription factor (1), other (3) | $4(3)$ | $25(7)$ |
| Stress | Involved in response to cold | 25 | Catalysis (3), kinase/phosphatase (8), |  |  |
|  |  |  | transcription factor (6), other (8) |  |  |
| Interactome | Interact with genes identified above | 9 | Catalysis (2), kinase/phosphatase (1), transport (5), other (1) | $9(2)$ |  |
| Total |  | 48 | Catalysis (10), kinase/phosphatase (10), interactions (3), |  | transcription factors (7), transport (6), other (12) |

counterparts in Prunus. Of note, 102 reactions were encoded by unique genes with the other steps provided as an additional gene function. The database screen enlarged the category of functions studied as 38 genes encoded non catalytic proteins (Table 2). Four Arabidopsis genes did not produce hits in either Prunus sp . It is important to note that the amino acid metabolism study evaluated genes by family while the phenotypic search concentrated on single units. Therefore, the two lists are studied separately although as will be seen, the conclusions reached are similar.

Perusal of the amino acid metabolism reactions reveals that both Prunus sp. harbor 2.0 genes per reaction (Table 1) while Arabidopsis stands at 2.4 isozymes per reaction. The largest variation is observed for the synthesis of aromatic compounds with 3.4 genes per reaction for Arabidopsis versus 2.4 or 2.6 for the Prunus sp . The number of genes per reaction was counted for all species and listed in Table 3 (columns "obs"). The reaction with zero genes in Prunus persica corresponds
to the chloroplastic methionine synthase enzyme and will be developed later. Effectively, the number of isozymes per reaction was viewed as a distribution problem of a given number of genes between a certain amount of reactions. Therefore, frequencies of given combinations were calculated relative to the most common distribution. The latter was found to be a large number of reactions encoded by two genes with the complement performed by 3 isozymes (Arabidopsis) or only one (Prunus). Hence, the ratio gene to reaction favors functional redundancy.

Observed combinations harbor a majority of reactions encoded by 1-3 genes with a few larger families. The latter explain the low frequencies of the observed gene distributions. Indeed, Arabidopsis harbors a higher number of large gene families than either Prunus sp. A reduction of that figure to values observed in Prunus increases the frequency of the combination from $1.3 \times 10^{-11}-2.2 \times 10^{-3}$ (Table 3). Finally, setting the number of genes with two reactions to that observed, results in

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Table 3: Number of reactions with a given count of genes

|  |  | Arabidopsis thaliana |  |  |  | Prumus persica |  |  |  | Prumus mume |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | Max distrib. |  |  |  | Max distrib. |  |  |  | Max distrib. |  |  |  |
| per Rx | Obs | NA | $\mathrm{x}_{2}=36$ | $x>3$ | $\mathrm{x}>3(\mathrm{Pp})$ | Obs | NA | $\mathrm{x}_{2}=25$ | $x>3$ | Obs | NA | $\mathrm{x}_{2}=25$ | $x>3$ |
| 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| 1 | 38 | 0 | 20 | 25 | 0 | 48 | 5 | 42 | 31 | 52 | 13 | 46 | 35 |
| 2 | 36 | 76 | 36 | 62 | 92 | 25 | 98 | 25 | 62 | 25 | 90 | 25 | 59 |
| 3 | 13 | 27 | 47 | 0 | 1 | 19 | 0 | 35 | 0 | 17 | 0 | 31 | 0 |
| 4 | 9 | 0 | 0 | 9 | 5 | 5 | 0 | 1 | 5 | 6 | 0 | 1 | 6 |
| 5 | 3 | 0 | 0 | 3 | 4 | 4 | 0 | 0 | 4 | 2 | 0 | 0 | 2 |
| 6 | 1 | 0 | 0 | 1 | 1 | 1 | 0 | 0 | 1 | 1 | 0 | 0 | 1 |
| >6 | 3 | 0 | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| Rel. Freq. | $4.9\left(10^{-16}\right)$ | 1 | $3.0\left(10^{-4}\right)$ | $9.6\left(10^{-14}\right)$ | $2.2\left(10^{-3}\right)$ | 1.3 (10-11) | 1 | $2.3\left(10^{-7}\right)$ | $5.8\left(10^{-8}\right)$ | $1.1\left(10^{-9}\right)$ | 1 | $1.2\left(10^{-6}\right)$ | $1.8\left(10^{-9}\right)$ |

Observed distributions are given in the columns "Obs" along with the most probable combination (NA) and the most frequent ones for the observed number of reactions encoded by 2 genes ( $x_{2}=\ldots$ ) and for the observed number of reactions encoded by $>3$ genes ( $x>3$ ). The distribution with the $P p x>3$ was calculated for Arabidopsis as a comparison. Frequencies of all combinations are given relative to the most common distribution. All calculations were performed in $R$ Version 3.1.2 for Windows 64 bit with the packages gmp and lpSolveAPI. The study was performed on the set of metabolic reactions with unique genes. A reaction occurring in two different compartments was considered as two separate entities. The aspartate aminotransferase reactions were discounted due to the uncertainties regarding localization. Moreover, a few metabolic steps involve several protein units each counted as a reaction. Thus, the total number of reactions is 103 versus the previously cited 102

Table 4: Comparison of the number of genes per reaction between At, Pp and Pm

|  | Arabidopsis thaliana (p-value $=3.8 \times 10^{-4} \pm 6.2 \times 10^{-5}$ ) |  |  |  |  |  | Prumus mume ( p -value $=0 \pm 0$ ) |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Genes/Rx | 0 or 1 | 2 | 3 | $>3$ | Tot | 0 or 1 | 2 | 3 | $>3$ | Tot |
| Prunus persica | 0 or 1 | 27 (18.1) | 18 (17.1) | 2 (6.2) | 2 (7.6) | 49 (49) | 47 (24.7) | 2 (11.9) | 0 (8.1) | 0 (4.3) | 49 (49) |
|  | 2 | 9 (9.2) | 11 (8.7) | 3 (3.2) | 2 (3.9) | 25 (25) | 5 (12.6) | 18 (6.1) | 2 (4.1) | 0 (2.2) | 25 (25) |
|  | 3 | 2 (7.0) | 4 (6.7) | 6 (2.3) | 7 (3.0) | 19 (19) | 0 (9.6) | 4 (4.6) | 13 (3.2) | 2 (1.6) | 19 (19) |
|  | >3 | 0 (3.7) | 3 (3.5) | 2 (1.3) | 5 (1.5) | 10 (10) | 0 (5.1) | 1 (2.4) | 2 (1.6) | 7 (0.9) | 10 (10) |
| Tot |  | 38 (38) | 36 (36) | 13 (13) | 16 (16) | 49 (30.6) | 52 (52) | 25 (25) | 17 (17) | 9 (9) | 85 (34.9) |

Data are based on the same reaction set as for Table 3. Observed values for each combination are provided with the expected values in parenthesis. Numbers in bold indicate reactions with identical numbers of genes. The bolded total is the sum of all reactions with conserved gene numbers. A Multinomial Goodness of Fit test was used to evaluate whether the observed distribution was different from the calculated one. The resulting p-values are provided. The statistical test was performed in R Version 3.1.2 for Windows 64 bit (R Core Team, 2014) with the package XNomial (Engel et al., 2010)
optimal combinations with a mixture of reactions catalyzed by 1-3 (Arabidopsis) or 1-4 (Prunus) enzymes. Both resulting Prunus distributions harbor larger numbers of reactions encoded by single genes then that of Arabidopsis. The real distributions also count a larger number of reactions encoded by single genes in Prunus versus in Arabidopsis.

Next, the observed combinations were used to calculate the probability of two species to harbor identical or different gene numbers for a given reaction. Four categories were listed namely reactions encoded by 0 or $1,2,3$ or $>3$ genes, thus yielding 16 combinations. Those are listed in Table 4 with observed values in the form of Punnett squares for Arabidopsis vs. Prunus persica and for Prunus mume versus Prunus persica. Results for Arabidopsis versus Prunus mume are comparable to those of Arabidopsis versus Prunus persica which is why they are omitted. A total of 49 or 50 reactions ( 48 or $49 \%$ ) have conserved number of genes in Arabidopsis and Prunus persica or Prunus mume, respectively. This is higher than the expected 30.6 or 31.4. The number rises to 85 or $82 \%$ when comparing the two Prunus sp.

Moreover, extreme combinations such as 0 or 1 gene in one species versus over 3 in the second one also show large differences between observed and calculated values. A Multinomial Goodness of Fit test produced $p$ values in the order of $10^{-4}$ or less. Consequently, gene numbers per reaction are more conserved between species than would be expected from a random allocation.

In conclusion, the observed gene distributions within a species result from a combination of biological requirements for large gene families and optimal repartition. Moreover, gene allocation is conserved between species particularly so between Prunus mume and Prunus persica.

Several Prunus genes harbor two or more counterparts in Arabidopsis: Homologues in Prunus were searched for all Arabidopsis genes identified based on phenotype. Most of the time, BLAST provided several hits with relatively close scores. However, the objective was to determine the most likely Prunus counterpart for a given Arabidopsis gene. As a result, a multiple alignment was performed for all selected BLAST hits and identity

Table 5: Percent identity matrix generated for the leucine rich repeat receptor kinase AT4G39400 and similar genes

| Identify | ppa000438m | XM_8236051 | AT2G01950 | ppa022290m | XM_8222737 | AT4G39400 | ppa000566m | XM_8234124 | ppa000552m | XM_8248236 | AT3G13380 | AT1G55610 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ppa000438m | 100 | - | - | - | - | - | - | - | - | - |  | - |
| XM_8236051 | 99 | 100 | - | - | - | - | - | - | - | - | - | - |
| AT2G01950 | 46 | 46 | 100 | - | - | - | - | - | - | - | - | - |
| ppa022290m | 46 | 46 | 66 | 100 | - | - | - | - | - | - | - | - |
| XM_8222737 | 46 | 46 | 65 | 99 | 100 | - | - | - | - | - | - | - |
| AT4G39400 | 46 | 45 | 53 | 51 | 51 | 100 | - | - | - | - | - | - |
| ppa000566m | 46 | 46 | 50 | 52 | 51 | 67 | 100 | - | - | - | - | - |
| XIM_8234124 | 46 | 46 | 50 | 52 | 51 | 66 | 98 | 100 | - | - | - | - |
| ppa000552m | 47 | 46 | 52 | 52 | 52 | 53 | 54 | 54 | 100 | - | - | - |
| XM_8248236 | 47 | 46 | 52 | 52 | 51 | 52 | 53 | 52 | 98 | 100 | - | - |
| AT3G13380 | 46 | 45 | 53 | 51 | 51 | 54 | 54 | 54 | 66 | 66 | 100 | - |
| AT1G55610 | 45 | 45 | 54 | 52 | 52 | 54 | 54 | 53 | 65 | 66 | 79 | 100 |

Results were derived by Clustal Omega (Goujon et al., 2010; Sievers et al, 2011) based on the alignment performed with default parameters. Percent identity numbers are bolded for Pp and Pm homologues as well as for the best aligning At and Pp, Pm homologues. Prumusmume gene names (XIM_..) were shortened by omitting the two leading zeros and deleting the final ". 1 "
matrixes obtained. As an example, Table 5 provides the identity matrix obtained for the leucine rich repeat receptor kinase AT4G39400 and hits. As can be seen, AT4G39400 aligns best with ppa000566m and XM_008234124.1 with an identity around $65 \%$. However, both AT3G13380 and AT1G55610 show preferential alignment for the same hypothesized that ppa000566m and XM_008234124.1 intervene in the same signaling network as AT4G39400 while ppa000438m and XM_008236051.1 share roles specific to Prunus. Finally, identity between Prunus homologues is at 98 or $99 \%$ showing tremendous gene conservation in the species. The approach is valid provided all genes with a significant alignment to the query are selected. This is why a conservative threshold of $16 \%$ alignment was set for the selection of BLAST hits. The number separated random hits from those of potential interest.

Counterparts were determined for all 44 Arabidopsis genes with hits. A total of 15 preferentially aligned with single Prunus genes. Matches to AT3G26744 were treated as clear counterparts although identity was significant for half of the sequence. Two genes, namely AT2G43790 and AT1G74520 were considered to have clear counterparts. Indeed, though two Arabidopsis sequences aligned with each Prunus hit, identity differed by $8-10 \%$. Finally, a protein alignment identified a preferential counterpart for AT1G50720. The remaining 26 assignments were ambiguous for Arabidopsis, Prunus persica and Prunus mume 22, nine and ten times, respectively.

Clear counterparts are listed in Table 6 with the identity between Prunus and Arabidopsis as well as that between the Prunus genes. Sequence identity between Arabidopsis and Prunus ranges from $60-76 \%$ while that between two Prunus counterparts lies at $95-99 \%$. The numbers confirms the high conservation of Prunus sequences. Moreover, genes originate from all databases in similar proportions. This shows the necessity of using several sources for a gene search based on phenotype.

Localization may vary between counterparts: Localization was determined for all genes based on literature, TAIR annotation, protein sequence alignment and prediction. Information was fairly consistent for proteins found in the cytoplasm, the chloroplast, the mitochondria or the nucleus. However, targeting predictions produced contradicting results for proteins located elsewhere. Hence, the compartment for the genes identified based on phenotype remains for the most part unclear. Moreover, a certain number of aspartate aminotransferase proteins could not be clearly assigned. Localization did differ between a few counterparts. Consequently, protein sequences were aligned to pinpoint consistencies between compartment assigned and sequence length. Three examples are provided in Table 7.

Methionine synthase is encoded by three genes in Arabidopsis and two in the Prunus species. The enzyme is located in the chloroplast and cytosol. However, targeting programs failed to find a chloroplastic isoform in Prunus persica. A protein sequence alignment shows that the results are consistent (Table 7). Indeed, all sequences harbor a methionine at the start of the corresponding cytosolic sequence. Only two proteins exhibit clear localization sequences, namely AT5G20980 and XM-008232732.1. No other sequences were detected for persica. Therefore, Prunus persica may harbor purely cytosolic forms of methionine synthase.

The two other examples are branched-chain amino acid transaminase and acetylserine lyase. Once again, Prunus proteins align with Arabidops is isozymes targeted to an organelle but are predicted to be in different compartments. The sequence alignment shows that the results are consistent. Indeed, the protein supposedly located in the cytoplasm displays a shorter sequence than its organelle homologue (Table 7). Surprisingly, the cytoplasmic arabidopsis sequences are slightly longer with an appearance of a signaling peptide. However, the targeting data is based on experimental results.

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| At gene | Description | Pp counterpart | Pm counterpart | $\mathrm{Pp} / \mathrm{Pm}$ |
| :---: | :---: | :---: | :---: | :---: |
| AT1G10760 ${ }^{\text {a }}$ | Pyruvate phosphate dikinase | ppa000209m (71) | XM_008247257.1 (69) | 98 |
| AT2G40840 ${ }^{\circ}$ | Disproportionating enzyme 2 | ppa000782m (72) | XM_008245862.1 (72) | 98 |
| AT3G52180 ${ }^{\circ}$ | Protein phosphatase | ppa007299m (67) | XM_008233843.1 (67) | 98 |
| AT1G34790 ${ }^{\circ}$ | Zinc finger protein | ppa016755m (64) | XM_008221233.1 (63) | 99 |
| AT5G23630 ${ }^{\text {b }}$ | ATPase cation pumps | ppa000424m (74) | XM_008220607.1 (74) | 99 |
| AT1G69180 ${ }^{\text {b }}$ | Transcription factor | ppa014900m (65) | XM_008245598.1 (64) | 99 |
| AT1G68560 ${ }^{\text {b }}$ | $\alpha-1$-arabinofuranosidase/ $\beta$-D-xylosidase | ppa001168m (70) | XM-008244999.1 (70) | 99 |
| AT5G49360 ${ }^{\text {b }}$ | $\alpha$-1-arabinofuranosidase/ $/$-D-xylosidase | ppa001718m (68) | XM_008225083.1 (67) | 98 |
| AT2G01390 ${ }^{\text {b }}$ | (TPR)-like protein | ppa004294m (62) | XM_008243399.1 (61) | 98 |
| AT4G39400 ${ }^{\circ}$ | Leucine-rich repeat receptor kinase | ppa000566m (67) | XM_008234124.1 (66) | 98 |
| AT3G26744 ${ }^{\circ}$ | Transcription activator | ppa005038m (75) | XM_008241330.1 (74) | 9899* |
| AT2G39810 ${ }^{\text {a }}$ C | Novel protein | ppa000974m (65) | XM_008240210.1 (64) | 97 |
| AT3G59770 ${ }^{\text {c }}$ | Phosphoinositide phosphatase | ppa000157m (72) | XM_008244816.1 (71) | 99 |
| AT4G04920 ${ }^{\circ}$ | Nuclear targeted protein | ppa000947m (67) | XM_008226829.1 (74) | 86 |
| AT5G13650 ${ }^{\circ} \mathrm{C}$ | Srv3 | ppa002327m (76) | XM_008242662.1 (76) | 99 |
| AT5G50720 ${ }^{\circ}$ | AtHVA22e | ppa013097m (68) | XM_008241283.1 (68) | 95 |
| AT1G74520 ${ }^{\circ}$ | AtHVA22a | ppa012417m (71) | XM_008241585.1 (71) | 97 |
| AT2G43790 ${ }^{\circ}$ | MPK6 | ppa006536m (76) | XM-008244822.1 (76) | 99 |
| AT3G54300 ${ }^{\text {d }}$ | Synaptobrevin-likeproteinfamily | ppa010737m (76) | XM_008246668.1 (76) | 98 |
| AT5G47910 ${ }^{\text {d }}$ | Respiratory burst oxidase protein D | ppa000883m (67) | XM_008224070.1 (67) | 97 |

Table 7: Examples of counterparts with potentially different alignments. Homologue names and methionines are bolded. Arabidopsis sequences were included for all compartments as a reference. Localization abbreviations are as for Table 1. All alignments were performed with Clustal Omega (Goujon et al., 2010; Sievers et al., 2011) using default parameters

| Protein | Alignment |
| :---: | :---: |
| Methionine synthase | AT5G20980 (Cl) MGQLALQRLQPLASLPRRPPSLPPPSSATPSLPCATASRRPRFYVARAMSSHIVGYPRIG |
|  | ppa021650m (Cy) ----------------------------------------------------------------------------------------MASHVIV |
|  | XM_8232732 (Cl) MKQ---------------VSSITFGP-CYGSLCFSAKRPTLLRFTHHFKFHSTRAMASHIVGLPRIG |
|  |  |
|  |  |
|  |  |
|  | AT3G03780 (Cy) |
| Branched-chain amino acid transaminase |  |
|  | AT1G10060 (Mt) ------------MALRRCLPQYSTTSSYLSKIWGFRMH---------GTKAAASVVEEHVSGAEREDEE |
|  | AT1G10070 (Cl) MIKTITSLRKTL--------VL--------PLHLHRTLQTFAKYNAQAASALREERKKPLYQNGDDV |
|  | ppa008826m (Cy) |
|  | XM_8222198 (Mt) MIQRTTRLHKLVRSIGVGSSLSSSSKQLRVHRCFSSVAASNA-EQACEQSVESYNVKKNE |
|  | AT1G50110 (Cy) YANVKWEELGFALTPIDYMYVAKCRQGESFTQGKIVPYGDISISPCSPILNYGQGLFEGL |
|  | AT1G10060 (Mt) YADVDWDNLGFSLVRTDFMFATKSCRDGNFEQGYLSRYGNIELNPAAGILNYGQGLIEGM |
|  | AT1G10070 (Cl) YADLDWDNLGFGLNPADYMYVMKCSKDGEFTQGELSPYGNIQLSPSAGVLNYGQAIYEGT |
|  | ppa008826m(Cy)-----------------------------------MYVMKCSNNGTFEKGQLNRYGNIELNPAAGVLNYGQGLYEGT |
|  | XM_8222198(Mt) YADVDWDNLGFGLTPTDYMYVMKCSNNGTFEQGQLNRYGNIELNPAAGVLNYGQGLYEGT |
| Acetylserinelyase | AT3G04940 (Cy) |
|  | AT3G03630 (Cl) MAFASPSLRLLPQSPLGRITSKLHRFSTAKLSLFSFHHDSSSSLAVRTPVSSFVVGAISG ppa007201m (Cl) MAILSAPLLLSLPHPF-SFPSKRHRFGTFKVSSSILS |
|  | XM_8242347 (Cy) |
|  | AT3G04940 (Cy) -------------------------------------------------------------MEEDRCSIKDDATQLIGNTPMVYLNNV |
|  | AT3G03630 (Cl) KSSTGTKS-KSKTKRKPPPPPPVTTVAEEQHIAESETVNIAEDVTQLIGSTPMVYLNRVT |
|  | ppa007201m (Cl) -----TNGALLRRQFTQRYPLVFAK---ASSVYATREDLDTVNIAEDVTQLIGSTPMVYLNKVT |
|  |  |

## DISCUSSION

Prunus genes show a strong degree of conservation: All species show a similar gene distribution per function with
the majority of catalytic steps being performed by 1-3 genes while a few reactions require larger numbers. Moreover, a given reaction is more likely to harbor the same number of genes in a second species than would be
expected from a random distribution. Hence, there is a general conservation in gene distribution patterns between organisms. However, the degree of conservation is particularly pronounced between the two Prunus sp. with over $80 \%$ of the reactions harboring similar gene numbers. The results are confirmed at the sequence level as gene identities between Prunus counterparts are over $90 \%$. This reflects previous data as microarrays designed for Prunus persica function with apricot extracts (Manganaris et al., 2011). The surprise comes from the extent of conservation.

The results have to be contrasted with the diversity observed within a species. As mentioned in the introduction, polymorphism studies identified significant variation in apricot varieties (Takeda et al., 1998; Hagen et al., 2002; Yuan et al., 2007; Krichen et al., 2008; Yilmaz et al., 2009). Moreover, close to a million informative SNPs were found in Prunus persica accessions (Verde et al., 2013). The polymorphism is reflected at the phenotypic level with significant variations in phenolic compounds, carotenoids, sugar and acid levels (Ruiz et al., 2005a, b; Drogoudi et al., 2008; Engel et al., 2010; Schmitzer et al., 2011; Gundogdu et al., 2013) as well as fruit ripening times (Asthma, 2012). Consequently, a potential for diversity exists in the Prunus sp. in spite of the strong genome conservation.

Variation is also observed in protein length. At least three cases are linked to differences in localization. For instance, Prunus persica does not seem to harbor a plastidic methionine synthase. Results are based on predictions which assume a single compartment while several proteins have dual locations (Carrie et al., 2009). Hence, the chloroplastic location may have been overlooked. In fact, the prediction programs suggest a second targeting to the mitochondria. The cytosolic methionine synthase is definitely essential in regenerating the methyl group of S -adenosyl-L-methionine, an intermediate in the methionine salvage pathway (Ravanel et al., 1998). Conversely, the chloroplastic form ensures autonomy of the organelle for methionine synthesis (Ravanel et al., 2004). The single mutant presented in the Chloroplast 2010 database looks smaller but otherwise fairly normal. Consequently, the peculiar localization distribution in Prunus persica appears to be possible. The other two cases merely change the repartition of proteins between compartments. The three cases are an underestimate of the potential localization differences between counterparts. Moreover, when related to the number of proteins with several compartments, the ratio appears quite significant. Hence, at the genome scale a noteworthy number of counterparts in different species may encode proteins with different localizations.

In conclusion, gene to gene correspondence seems fairly straight forward to pinpoint in the Prunus sp. This opens many possibilities in terms of combining transcriptomic data from different organisms. However, polymorphic variations and potential differences in localization suggest that caution must still be exercised when assessing the fine details of a gene function.

Arabidopsis data is potentially applicable to Prunus:
While Arabidopsis is clearly different from a tree, a surprising number of functional elements appear to be conserved. Thus, Prunus homologues were identified for 44/48 Arabidopsis genes. Moreover, an Arabidopsis pseudogene listed in amino acid metabolism matched sequences in both Prunus sp. Genes are potentially conserved for at least three reasons: a similar role, a similar function but with different applications in each organism, a similar gene origin. Clearly, differentiation between these mechanisms is essential in correctly transferring data from Arabidopsis to Prunus. Precious information is provided by sequence alignments. As an example, identity matrixes show clear examples of Prunus genes sharing similar conservation with all genes of the corresponding Arabidopsis family. This would point to a shared function or an evolutionary relic rather than to a similar role.

A second issue is the preferential alignment of a given gene with several counterparts. For instance, only 18 Arabidopsis genes harbor a clear Prunus counterpart with correspondence assignment being ambiguous in the other cases. Three Arabidopsis genes involved in cold response aligned with a single Prunus counterpart thus causing complexities in the comparison of the cold response signaling pathway. A potential answer may be found in the studies on gene duplication. Genome comparison shows clear examples of the enlargement of specific gene families in given organisms. For instance, Prunus persica harbors a large number of genes devoted to fruit quality (Verde et al., 2013). Moreover, several families important for the production of a lignified seed are enlarged. The increase is attributed to gene duplication as an adaptive process to deal with a lignified stone. In Arabidopsis over $80 \%$ of the genome represents duplicated regions (Briggs et al., 2006). Duplicates are often maintained albeit with a reduced function. Therefore, the evolutionary information provided by sequencing data is essential for understanding gene to gene correspondence between species.

Databases are a key tool for the modern biologist: As mentioned, moving from model plants to crops requires a sensible perusal of the massive data currently being
generated. As a result, databases of all type are an essential tool for the modern biologist. Unfortunately, progress in sequencing and other data production has exponentially increased the cost and manpower necessary to maintain databases. The funding situation encountered by TAIR a few years ago shows that even major players are at risk. Thankfully, they are now "thriving" thanks to donations. But what is the status of less frequently used databases? Sadly, the issue is not easy to resolve. Indeed, this type of resource needs to generate biological data and publications to justify academic funding. This is usually possible in the early stages of the project but less so in the maintenance phase. Hence, there is a risk of information being lost through lack of funding. Moreover, as the number of available databases grows, it becomes increasingly important to maintain their low cost. Otherwise, the average researcher will be deprived of essential resources in case of insufficient funding. In short, data management is an important and difficult issue to resolve. Currently, the massive amounts of data produced are underused.

## CONCLUSION

The results point to a remarkable conservation of the Prunus genes both in terms of number of enzymes per family and of sequence alignment. Second, the basis of comparison across species lies in the existence and correct assignment of homologues. The data show that though homologous genes are detected most of the time the assignment of a clear gene to gene correspondence is often ambiguous between Arabidopsis and Prunus. Finally, clear protein counterparts do not necessarily share the same cellular localization. In short, Prunus data will likely apply to Prunus armeniaca thanks to the remarkable conservation of the genus. However, Arabidopsis information may be valid to some extent.

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