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The salivary proteome of *Tetranychus urticae*

Key to its polyphagous nature?

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Jonckheere, W. S. A. (2018). *The salivary proteome of Tetranychus urticae: Key to its polyphagous nature?*

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4

A link between host plant range and the salivary protein repertoire of spider mites

Partly redrafted after:

Jonckheere, W., Dermauw, W., Zhurov, V., Wybouw, N., Van den Bulcke, J., Villaroel, C., Greenhalgh, R., Grbic, M., Schuurink, R.C., Tirry, L., Baggerman, G., Clark, R.M., Kant, M.R., Vanholme, B., Menschaert, G., Van Leeuwen, T. (2016)
The salivary protein repertoire of the polyphagous spider mite *Tetranychus urticae*: a quest for effectors. *Molecular & Cellular Proteomics* 15, 3594–3613

4.0. ABSTRACT

The proteomic composition of secreted saliva from *Tetranychus urticae* lines adapted to different host plant species (bean, maize, soy and tomato) was analyzed using a custom-developed feeding assay coupled with nano-LC tandem mass spectrometry. The mite's salivary protein composition depended on the host plant the mite was adapted to (**Chapter 3**). To supplement the proteomics analysis with statistically supported data, genome-wide expression profiling was performed on *T. urticae* lines adapted to five different hosts (bean, maize, soy, tomato, and additionally, cotton). Indeed, the plant species had an effect on the expression level of several salivary protein genes. Furthermore, the genomes of two closely related spider mite species with a distinct host range – the *Ulex europaeus* specialist *T. lintearius* and the Solanaceae specialist *T. evansi* – were searched for homologs of the salivary proteins identified in *T. urticae*. For some salivary protein families the number of members and the size of the host plant repertoire of the mite was correlated. This study confirms that the salivary protein cocktail delivered in the host depends on the plant species. In addition, it provides support for the hypothesis that the salivary proteome of spider mites co-evolved with the host plant range. This would suggest polyphagous herbivores require a more complex salivary (effector) protein repertoire than monophagous species.

4.1. INTRODUCTION

The cosmopolitan spider mite *Tetranychus urticae* is able to feed on an impressive array of different plant species. With more than 1100 different hosts recorded, belonging to more than 140 different families (Migeon and Dorkeld, 2006–2017), this species can be considered to be one of the most polyphagous arthropods. Distinct host plants represent different challenges for herbivores, as each is characterized by its own nutritional composition and defensive compounds. Furthermore, distinct plant species may respond differently to herbivory (**Chapter 1**). Considering these differences amongst plant species, it does not come as a surprise that mites adjust the expression of genes coding for proteins involved in digestion and detoxification to the host plant they are feeding on (Dermauw et al., 2013; Wybouw et al., 2015; Van Leeuwen and Dermauw, 2016). As such, *T. urticae* is well equipped to deal with the diverse secondary metabolites it encounters when feeding on its various hosts (Dermauw et al., 2013; Van Leeuwen and Dermauw, 2016).

In order to breach plant defensive barriers, herbivores produce effectors which target the products of certain plant susceptibility genes (S genes). Plants have, however, evolved to recognize some of these effectors by means of resistance gene (R gene) encoded receptors, effectively turning these effectors into elicitors of plant defenses. The interaction between effector and R or S gene product is highly specific and the nature of these genes can determine plant-herbivore compatibility (Kaloshian, 2004; Harris et al., 2012; van Schie and Takken, 2014). Reminiscent of digestive enzymes and ABC-transporters, the expression level of herbivore effectors may differ between host plants. The putative aphid effector ACYPI006346, for example, has different transcript levels in aphids adapted to different plants (Pan et al., 2015). Differential expression of effectors after adaptation to a new host plant may also occur in spider mites since mite adaptation has been shown to affect the transcriptional response of the host plant (Wybouw et al., 2015).

With the salivary proteome of *T. urticae* elucidated (**Chapter 3**), and with a host-specific salivary protein composition suggested by this proteomics analysis, we additionally studied the host-dependent expression level of all identified salivary proteins using genome-wide transcriptomics analysis of *T. urticae* lines adapted to five host plants (i.e., bean, maize, soy, tomato, and additionally, cotton).

Furthermore, the occurrence of homologs of *T. urticae* salivary proteins was studied in the related spider mite species *T. lintearius* and *T. evan-*

si. These mites are specialists on *Ulex europaeus* (Fabaceae) and the Solanaceae family respectively (Ireson et al., 2003; Migeon and Dorkeld, 2006-2017; Navajas et al., 2013). A glimpse into their salivary protein repertoire can provide additional insights in the molecular machinery of salivary proteins and will allow a better understanding of the link between salivary proteins and the compatibility of the plant-mite interaction.

4.2. MATERIAL AND METHODS

4.2.1. Establishment of *T. urticae* lines on different host plants

The *T. urticae* London strain has been maintained under laboratory conditions on bean plants (*Phaseolus vulgaris* cv. 'Prelude', Fabaceae) for many years, and the genome of this strain has been sequenced (Grbić et al., 2011). Lines on alternative host plants were established by transferring approximately 250 adult female mites from the London strain on bean to new hosts. These new host plants were cotton (*Gossypium hirsutum*, Malvaceae), maize (*Zea mays*, cv. 'Ronaldinio', Poaceae), soy (*Glycine max* cv. 'Merlin', Fabaceae) and tomato (*Solanum lycopersicum*, cv. 'Moneymaker', Solanaceae). Three independent lines were generated for tomato (Wybouw et al., 2015) and cotton, while four independent lines were obtained for the other hosts. The mite lines were maintained in a climatically controlled environment at 26°C with 60% RH, and a photoperiod of 16:8 h light:dark. Mites were offered fresh plants as needed, and were used in experiments after 5 generations for all hosts, except tomato, where replicate lines were adapted and maintained on tomato for over 30 generations (Wybouw et al., 2015).

4.2.2. Host plant-dependent salivary protein identification in artificial diet probed by *T. urticae*

The list of *T. urticae* salivary proteins, identified in **Chapter 3** using nano-LC-MS/MS analysis of *T. urticae* probed artificial diet, was split up again over the separate host plant lines (i.e., bean, maize, soy and tomato). Host plant specificity of putative *T. urticae* salivary proteins was visualized by means of a venn diagram depicting the number of proteins that are uniquely identified in the samples of a specific host plant line and the number of proteins that are shared by different samples (FIGURE 4.1.A). In addition, a heatmap visualizing the mean rTop3 factor of a selection of candidate salivary proteins (maximum rTop3 value higher than the 30th percentile of all

maximum rTop3 values) for the different investigated host plants was generated (FIGURE 4.1.B).

4.2.3. Transcriptome analysis of *T. urticae* maintained on different host plants

For each host plant adapted mite line, three (tomato and cotton) or four (bean, maize and soy) biologically replicated RNA samples were obtained. Each RNA sample was extracted from 100-150 pooled female adult mites using the RNeasy extraction kit (Qiagen) and was subsequently treated with DNase (Turbo DNA-free kit, Ambion). RNA quantity and integrity was measured using an Agilent TapeStation system. RNA samples were labeled with cyanine dyes following the Low Input Quick Amp Labeling Kit (Agilent Technologies), with 100 ng of total RNA as starting material. RNA samples from mites feeding on the reference bean host plant were labeled with cy3, while cy5-labelling was performed on all other samples. Samples were pooled per host plant transfer and hybridized to a custom-made Sureprint G3 8x60K array (Agilent Technologies, with a GEO platform number of GPL16890) following the standard procedure of the Gene Expression Hybridization Kit (Agilent Technologies). After washing procedures (Gene Expression Wash Buffer kit; Agilent Technologies), raw data was extracted from the 8x60k slides using the GE2_107_Sep09 protocol of the Agilent Feature Extraction Software. The intraspot correlation coefficient per array and the metrics from the arrayQualityMetrics package per host plant line were assessed for optimal background correction and normalization procedures (Kauffmann et al., 2009). Data was background corrected using the 'normexp'-method and normalized by loess and Aquantile (Ritchie et al., 2007). Cyanine intensities were extracted from the processed RG-object and averaged per host plant. Using the normalized MA-object, differential expression was assessed for mites on cotton, maize, soy and tomato against the corresponding ancestral mite population living on bean by an empirical Bayes approach. A heatmap visualizing the expression levels (absolute and relative cyanine intensities) of putative salivary protein-encoding genes in mites adapted to the different hosts was generated (FIGURE 4.2). *Tetranychus urticae* gene expression data have been uploaded to the Gene Expression Omnibus with accession number GSE80337.

A phylogenetic analysis was performed for proteins belonging to OrthoMCL cluster Tu_MCL_35 and Tu_MCL_36. Except for tetur55g00110

(Tu_MCL_36) which is encoded by a pseudogene, proteins from each cluster were aligned using MUSCLE (Edgar, 2004). Model selection was done with ProtTest 2.4 (Abascal et al., 2005) and according to the Akaike information criterion WAG+G and WAG+G+F were optimal for the phylogenetic reconstruction of Tu_MCL_35 and Tu_MCL_36 proteins, respectively. Finally, for each alignment a maximum likelihood analysis was performed using Treefinder (v. 2011) (Jobb et al., 2004) bootstrapping with 1000 pseudoreplicates (LR-ELW). The resulting trees were midpoint rooted and edited with MEGA 6.0 software (Tamura et al., 2013) (FIGURE 4.3).

4.2.4. Mining of the *T. lintearius*, *T. evansi* and *T. urticae* proteomes for homologs of salivary proteins identified in *T. urticae*

The online OrthoMCL software tool (Li et al., 2003, Chen et al., 2006) was used to search the proteomes of *T. urticae*, the monophagous *T. lintearius* and the oligophagous *T. evansi* (versions of 11/08/2016) for homologs of the *T. urticae* salivary proteins identified in **Chapter 3** (TABLE 3.1). The genomes of *T. lintearius* and *T. evansi* are unpublished, yet are available to researchers from the Spider Mite Consortium via the ORCAE website (Sterck et al., 2012).

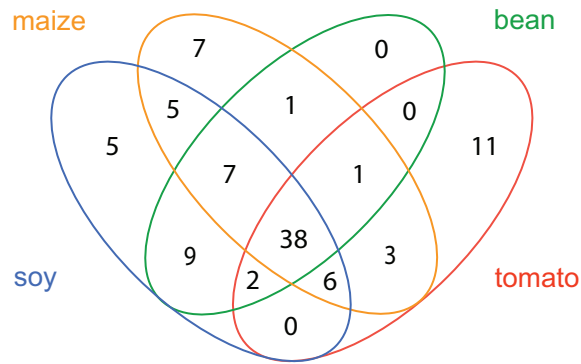
4.3. RESULTS

4.3.1. Host plant-dependent salivary protein composition

Host plant specificity of putative *T. urticae* salivary proteins (FIGURE 4.1.A), obtained by a proteomics analysis of artificial diet fed upon by mite lines

FIGURE 4.1. Overview of nano-LC-MS/MS identified putative *Tetranychus urticae* salivary proteins. (A) Venn diagram depicting overlap between putative *T. urticae* salivary proteins secreted by mites adapted to different host plants (bean, maize, soy, tomato). Only those salivary proteins with a mean PSM of at least two in at least one of the *T. urticae* host plant adapted lines were used for comparison (see TABLES 3.1 and S3.6). (B) Heat map of mean rTop3 values of putative *T. urticae* salivary proteins secreted by mites adapted to different host plants (bean, maize, soy, tomato). Only those salivary proteins (and ‘protein inference groups’) with a mean PSM of at least two in at least one of the *T. urticae* host plant adapted lines and with a maximum rTop3 value higher than the 30th percentile of maximum rTop3 values were used for comparison (see TABLES 3.1 and S3.6). The Euclidean distance metric and Ward’s method were used for clustering of both rows and columns. All putative salivary proteins for which the corresponding genes were shown to be expressed in the salivary glands by ISH (FIGURE 3.3) are shaded grey.

A



B

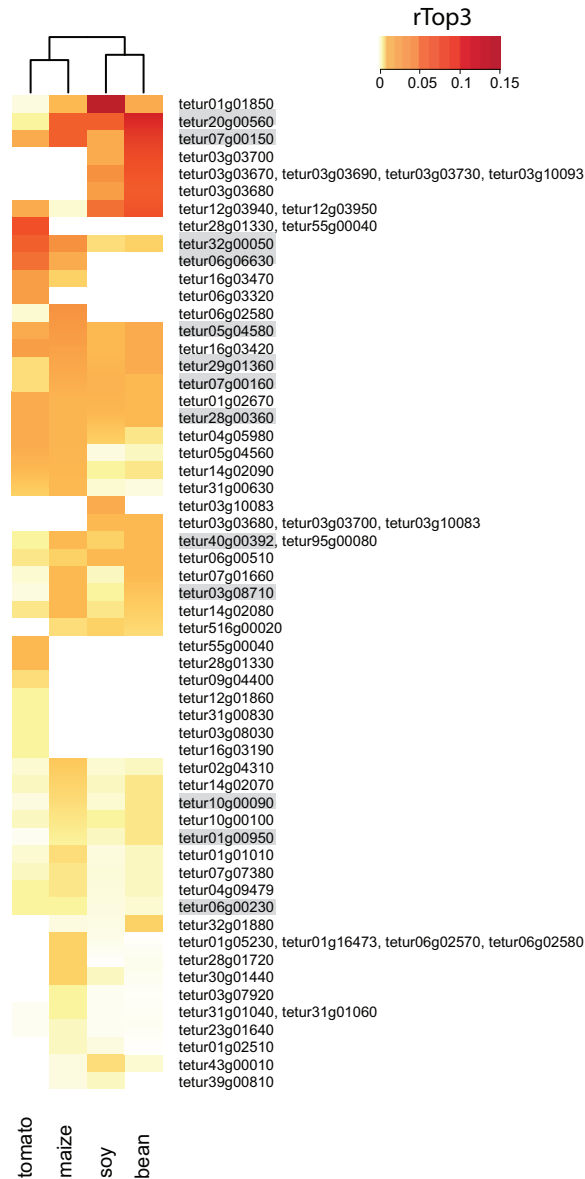


TABLE 4.1. The number of identified *Tetranychus evansi*, *T. lintearius* and *T. urticae* proteins homologous to proteins identified in artificial diet fed upon by *T. urticae*. Only proteins which could be assigned to groups already existing in the OrthoMCL database were accounted for in this list. Detailed information can be found in TABLE S4.1.

OrthoMCL	<i>T. evansi</i>	<i>T. lintearius</i>	<i>T. urticae</i>
OG5_126560	6	3	7
OG5_126583	7	6	10
OG5_126595	2	2	3
OG5_126607	12	9	21
OG5_126738	8	10	10
OG5_126942	6	8	15
OG5_127143	2	2	3
OG5_127584	8	8	8
OG5_127620	2	2	2
OG5_127800	9	10	29
OG5_128075	2	2	3
OG5_128163	3	3	4
OG5_128170	2	3	20
OG5_129300	3	1	4
OG5_129423	15	14	35
OG5_130246	1	1	1
OG5_130527	17	13	52
OG5_131746	2	3	2
OG5_132251	0	6	6
OG5_133467	8	7	15
OG5_134456	1	1	1
OG5_135950	3	5	5
OG5_136350	4	3	6
OG5_141111	3	2	2
OG5_144177	0	0	6
OG5_147492	5	4	5
OG5_149533	3	2	13
OG5_152237	0	1	2
OG5_152337	3	1	1
OG5_152454	3	3	3
OG5_158831	0	0	2
OG5_168371	0	0	5
OG5_176862	0	0	1
OG5_207753	0	0	2

adapted to different host plants (**Chapter 3**) was further illustrated by means of a heat map depicting the mean rTop3 factor of a selection of candidate salivary proteins (maximum rTop3 value higher than the 30th percentile of all maximum rTop3 values) for the different investigated host plants

(FIGURE 4.1.B). The most apparent case of host-specific salivary proteins was *tetur55g00040/tetur28g01330* (WTSP1, belonging to OrthoMCL group TuMCL_36), proteins with an unknown function, which were uniquely and abundantly identified from saliva of the tomato-adapted mites. Based on the clustering analysis, the salivary proteomic repertoire of *T. urticae* feeding on bean plants seemed most similar to the repertoire of soy-adapted mites (both plant species belong to the Fabaceae). However, since only one biological replicate (with two technical replicates) was analyzed with Nano-LC-MS/MS for each host plant specific diet sample, one should be careful to draw conclusions solely based on this comparative proteomics analysis.

4.3.2. Host plant-dependent expression of *T. urticae* salivary protein genes

To additionally validate the host-specific findings based on individual biological replicates, as well as to compare in further detail the expression level of genes coding for *T. urticae* putative salivary proteins across host plant species, we performed a genome-wide expression analysis of the mite lines that were used in the proteomic experiments (i.e., *T. urticae* adapted to maize, soy or tomato relative to the reference line on bean) using an Agilent gene expression microarray. Additionally, a line adapted to cotton during five generations was also investigated. We determined the absolute expression levels using normalized cyanine 3 (cy3) intensity values but also calculated the expression levels relative to the mites feeding on bean as a benchmark (FIGURE 4.2.A and B). We confirmed what we detected previously when comparing the rTop3 values of the proteomics data across different host plant lines (FIGURE 4.1): the salivary composition of *T. urticae* is host plant dependent. For example, in the tomato-adapted mite lines several salivary protein genes were highly expressed relative to bean-adapted mites: *tetur32g00050* (Tu_MCL_35), *tetur28g01330* and *tetur55g00040* (Tu_MCL_36) and *tetur31g00830* (Tu_MCL_25). Alternatively, *tetur03g03670*, *tetur03g10093*, *tetur03g03700*, *tetur03g03680*, *tetur03g03690* and *tetur03g03730* (OrthoMCL cluster OG5_144177; SHOT family, **Chapter 5**) were expressed at much lower levels in mites feeding from all host plants except for soy, compared to bean. The absolute expression level of the latter six genes was very high after feeding on bean or soy (cy3 intensity levels were in the top 1% of most highly expressed genes in mites on either bean or soy) and relatively low after feeding on the non-leguminous host plants under study (fold changes

Host plant dependency of salivary protein composition

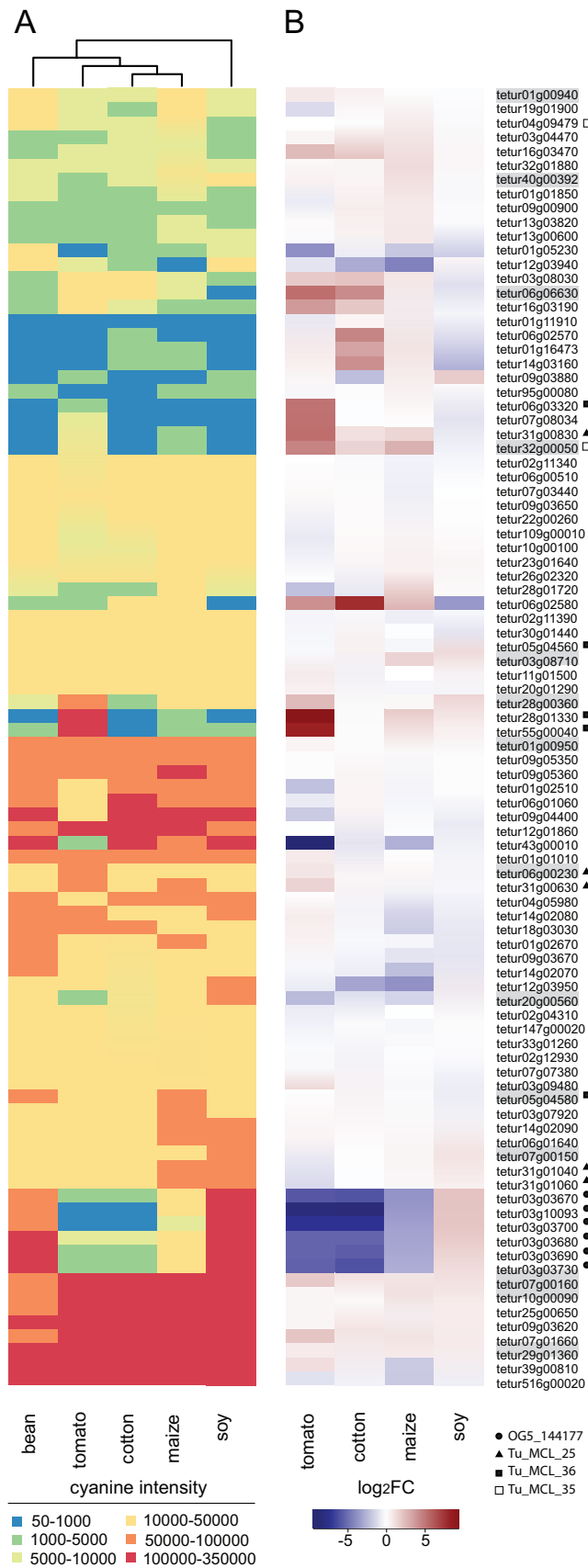


TABLE 4.2. The number of identified spider mite specific *Tetranychus evansi*, *T. lintearius* and *T. urticae* proteins homologous to proteins identified in artificial diet fed upon by *T. urticae*. Detailed information can be found in TABLE S4.2.

OrthoMCL	<i>T. evansi</i>	<i>T. lintearius</i>	<i>T. urticae</i>
Tu_MCL_12	1	1	37
Tu_MCL_25	2	4	17
Tu_MCL_43	2	6	9
Tu_MCL_35	3	0	14
Tu_MCL_36	4	0	12
Tu_MCL_45	0	0	12
Tu_MCL_63	0	2	8
Tu_MCL_74	2	1	5
Tu_MCL_211	2	2	2
Tu_MCL_212	1	2	2
Tu_MCL_153	1	2	2
No group	1	3	1
No group	2	0	2
No group	1	1	1
No group	0	0	3
No group	0	0	2
No group	1	0	1
No group	1	0	1

between mites on bean and mites on tomato, maize or cotton varied between 7 and 289).

4.3.3. Mining of the *T. lintearius*, *T. evansi* and *T. urticae* genomes for homologs of salivary protein genes identified in *T. urticae*

The genomes of *T. urticae*, *T. lintearius* and *T. evansi* were searched for homologs of *T. urticae* salivary protein genes identified in **Chapter 3**

FIGURE 4.2. Heatmap of expression levels of putative *Tetranychus urticae* salivary protein encoding genes and their up- or downregulation in mites adapted to different host plants. (A) Heatmap of cyanine intensities of putative *T. urticae* salivary protein encoding genes. The Euclidean distance metric and Ward's method were used for clustering of both rows and columns. For 92 out of 95 putative salivary protein genes expression data was available. (B) Heatmap of \log_2 FCs of putative salivary protein genes in mites adapted to soy, maize, cotton or tomato as compared to mites adapted to bean. Genes are sorted based on their order in panel A. Genes that were shown to be expressed in the salivary glands by ISH (FIGURE 3.3) are shaded grey. A circle, triangle, filled square and empty square indicates whether a gene belongs to OrthoMCL cluster OG5_144177, Tu_MCL_25, Tu_MCL_36 and Tu_MCL_35, respectively.

(TABLE 3.1). The OrthoMCL analysis grouped the proteins homologous to the identified putative *T. urticae* salivary proteins into 34 groups which already existed in the OrthoMCL database (<http://www.orthomcl.org/orthomcl/>) (TABLES 4.1 and S4.1), and in 18 groups which did not exist in the OrthoMCL database yet. Latter group names start with 'Tu_MCL' and are likely spider mite specific (TABLES 4.2 and S4.2). In what follows, I zoom in on some peculiarities.

The number of identified proteins belonging to an already existing group was, with 304 proteins, highest in *T. urticae*. *Tetranychus evansi* and *T. lintearius* had less than half of this number: 140 proteins for *T. evansi* and 135 proteins for *T. lintearius*. In 91% of these already existing OrthoMCL groups, *T. urticae* had at least as many members as *T. evansi* or *T. lintearius*. The extreme cases were OG5_127800 (Peptidase C1A, papain C-terminal), with 29 members in *T. urticae*, 9 in *T. evansi* and 10 in *T. lintearius* (TABLES 4.1 and S4.1). OG5_128170 (short-chain dehydrogenase/reductase SDR) had 20 members in *T. urticae*, yet only 2 and 3 in *T. evansi* and *T. lintearius* respectively. OG5_129423 (glycosyl hydrolase) had 35 members in *T. urticae*, 15 in *T. evansi* and 14 in *T. lintearius*. OG5_130527 (lipocalins) was, with 52 members, extremely expanded in *T. urticae*, while only 17 and 13 homologs were identified in *T. evansi* and *T. lintearius* respectively.

The number of identified proteins belonging to spider mite specific groups was, with 132 proteins, highest in *T. urticae*, while *T. evansi* and *T. lintearius* had less than one fifth of this amount: 25 proteins for *T. evansi* and 24 proteins for *T. lintearius*. In 89% of these spider mite specific OrthoMCL groups, *T. urticae* has at least as many members as *T. evansi* or *T. lintearius* (TABLES 4.2 and S4.2). The absolute expression levels of the *T. urticae* homologs of mites adapted to different host plants are shown in FIGURE 4.4.

The spider mite specific OrthoMCL group that includes most genes is **Tu_MCL_12**, which is, with 37 representatives, highly expanded in *T. urticae*. Strikingly, *T. evansi* and *T. lintearius* each possess only one homolog (tetev485g00030 and tetli56g00590 respectively). Proteins of Tu_MCL_12 lack homology with other known proteins, and their function remains unknown. Despite the expanded nature of Tu_MCL_12 in *T. urticae*, only tetur20g00560 could reliably be identified from secreted saliva. This protein may be one of the most prominent (putative) effectors in *T. urticae* saliva. Indeed, tetur20g00560 has the highest maximal rTop3 value of all putative salivary proteins identified in artificial diet fed upon by *T. urticae*

(TABLE 3.1). Furthermore, of all *T. urticae* homologs, only the expression of genes *tetur13g04230*, *tetur20g00050*, *tetur20g00090*, *tetur20g00540* and *tetur20g00560* was significantly higher (Benjamini-Hochberg adjusted $p \leq 0.05$ and $FC \geq 8$) in the proterosoma compared to the entire body. Genes encoding other homologs were not differentially expressed (DATA S3.9).

The second most gene-rich mite-specific OrthoMCL group is **Tu_MCL_25**. Again this group is expanded in *T. urticae*, where 17 representatives were found. Eight genes coding for these proteins have a significantly higher expression (Benjamini-Hochberg adjusted $p \leq 0.05$ and $FC \geq 8$) in the proterosoma relative to the entire body (DATA S3.9), while four proteins were detected in the artificial diet fed upon by *T. urticae*. In addition, several of the genes have a host-dependent expression level (Wybouw et al., 2015). In *T. evansi*, two homologs were identified and four could be found in *T. lintearius* (TABLES 4.2 and S4.2). The Tu_MCL_25 cluster includes proteins from ‘family 28’ (Tu28 and Te28), which we have shown to be effectors (**Chapter 2**).

The next spider mite specific OrthoMCL group that catches the eye is **Tu_MCL_43**. This group is expanded in *T. urticae* (nine members), has two members in *T. evansi*, and is with six members relatively well represented in *T. lintearius*. Seven out of the nine homologs in *T. urticae* have a high gene expression level in the proterosoma (Benjamini-Hochberg adjusted $p \leq 0.05$ and $FC \geq 8$) (DATA S3.9).

Tu_MCL_35 is expanded (14 members) in *T. urticae*, yet only *tetur04g09479* and *tetur32g00050* are significantly upregulated in the proterosoma, while *tetur12g00700* is even strongly downregulated in this head region (Benjamini-Hochberg adjusted $p \leq 0.05$ and $|FC| \geq 8$). Host dependent expression levels are evident (FIGURE 4.4). Tu_MCL_35 has three homologs in *T. evansi* and appears to lack representatives in *T. lintearius* (TABLES 4.2 and S4.2).

Some genes of the **Tu_MCL_36** group (WTSP) are highly expressed when *T. urticae* is feeding on tomato plants (FIGURE 4.3). Furthermore, two proteins of this group (*tetur28g01330* and *tetur55g00040*) were uniquely and abundantly identified from saliva of *T. urticae* adapted to tomato plants. This points to a tomato- or Solanaceae-specific function for these two proteins. Intriguingly, the Solanaceae specialist *T. evansi* possesses four genes coding for Tu_MCL_36 homologs, while homologs could not be found in the gorse-specialist *T. lintearius* (TABLES 4.2 and S4.2).

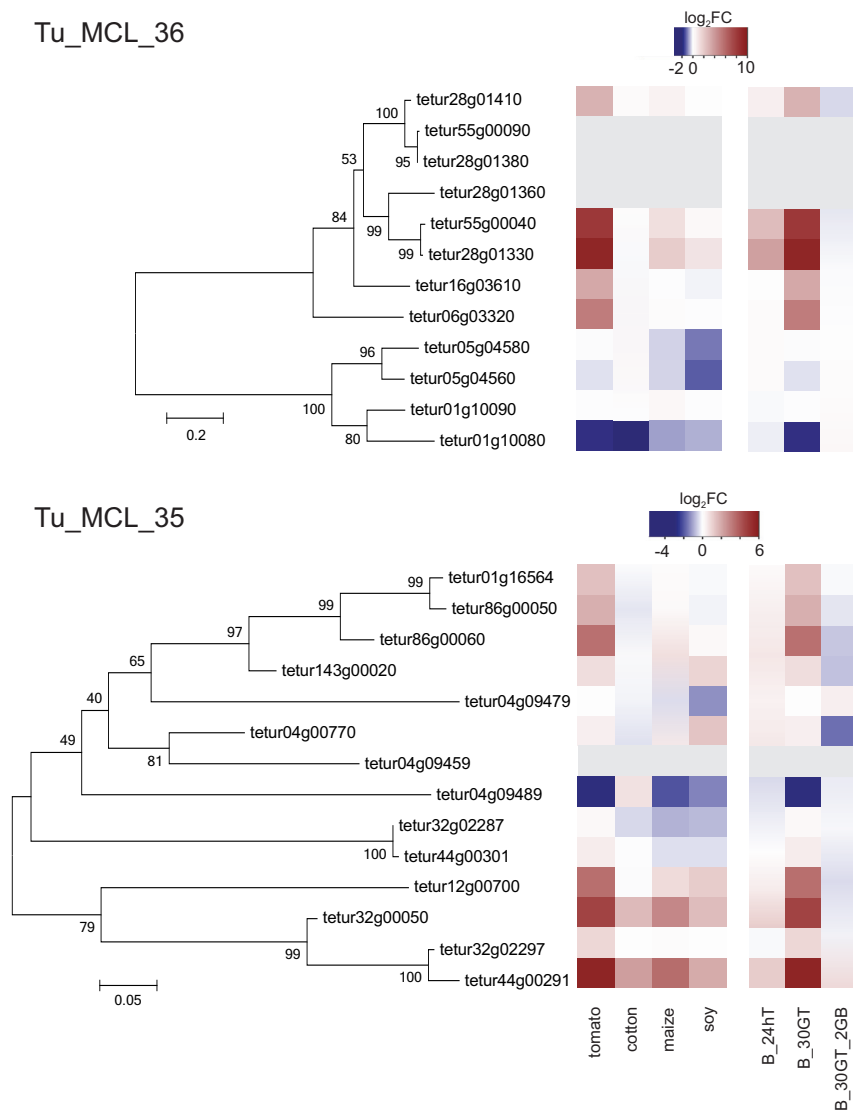
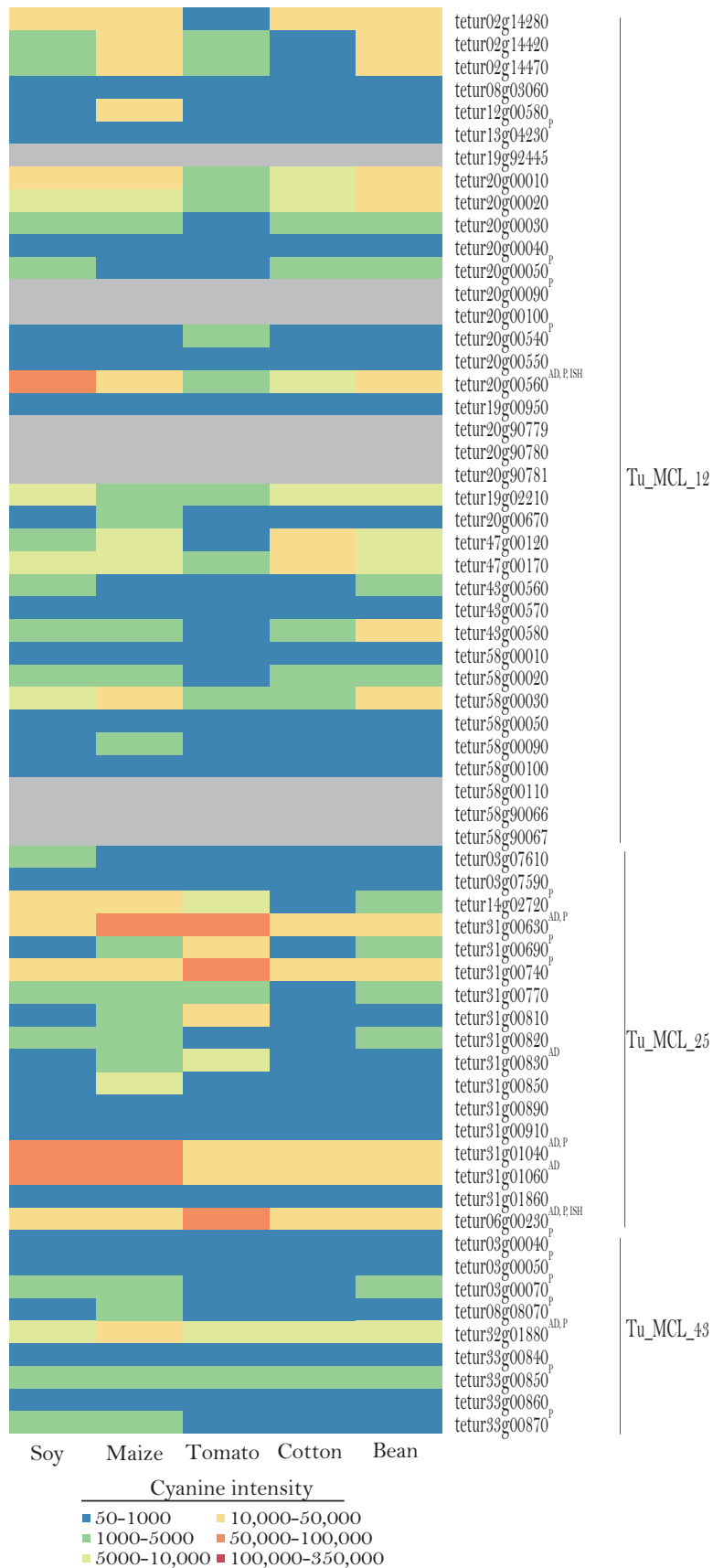


FIGURE 4.3. Expression profiles of genes coding for members of the expanded Tu_MCL_36 and Tu_MCL_35 protein families in *Tetranychus urticae*. Phylogenetic analyses of Tu_MCL_36 and Tu_MCL_35 proteins are shown next to heatmaps depicting relative gene expression of Tu_MCL_35 and Tu_MCL_36 genes in *T. urticae* subjected to different host plant regimes (\log_2 FCs, relative to the expression level on bean). The heatmap on the left represents \log_2 FCs of Tu_MCL_35 and Tu_MCL_36 genes in mites adapted to soy, maize, cotton or tomato compared to bean, while the heatmap on the right represents \log_2 FCs of Tu_MCL_35 and Tu_MCL_36 genes in *T. urticae* after host shifts between bean and tomato (B_24hT, mites from the London reference strain on bean transferred to tomato for 24h; B_30GT, mites from the bean strain grown on tomato for 30 generations; B_30GT_2GB, mites from the bean strain grown on tomato for 30 generations and transferred back to bean for two generations; Wybouw et al., 2015). Grey boxes indicate that for a specific gene no probes were included in the *T. urticae* microarray design, and hence expression could not be assayed.



Host plant dependency of salivary protein composition

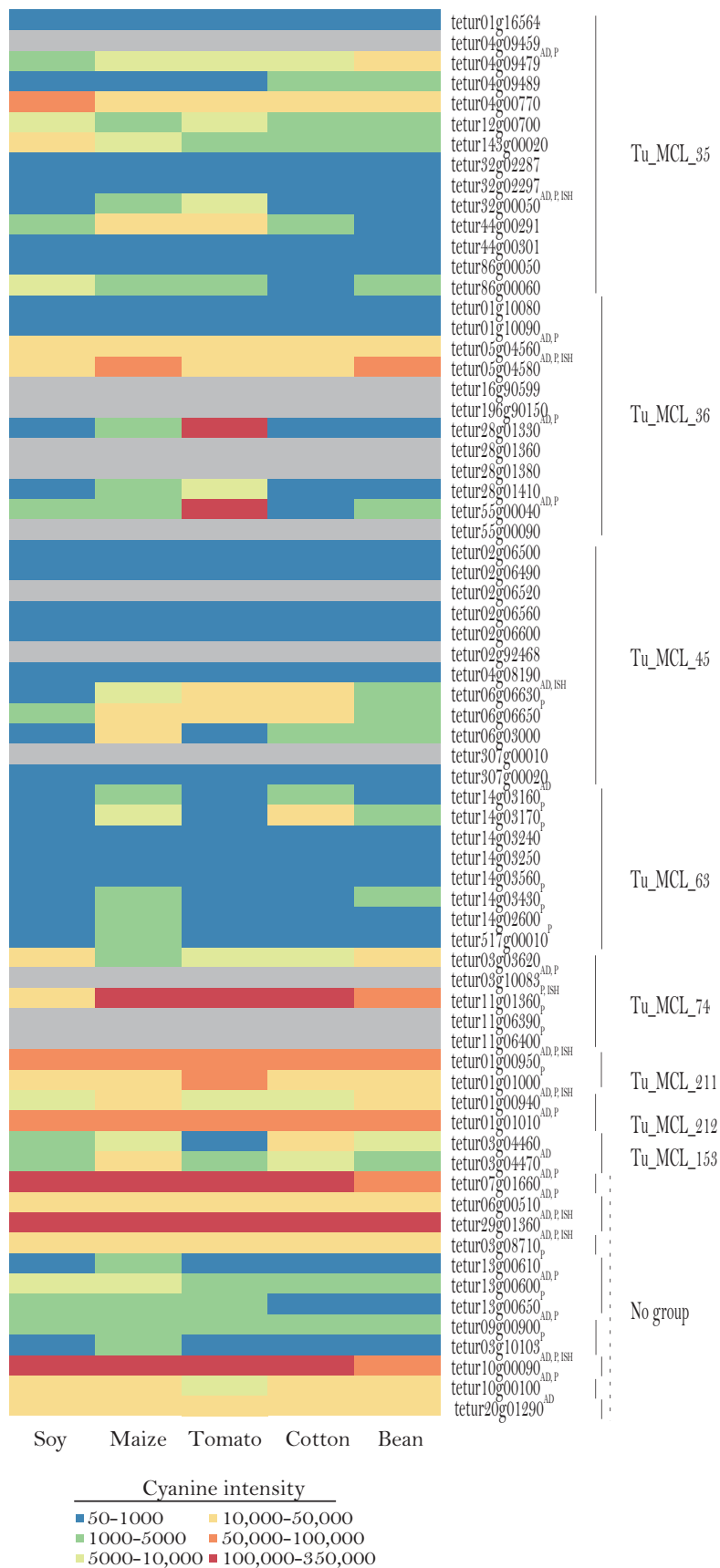


FIGURE 4.4 Heatmap of the absolute expression levels of a selection of genes of *Tetranychus urticae* adapted to different host plants. The selection consists of homologs of spider mite-specific genes encoding putative salivary proteins identified in artificial diet after feeding by *T. urticae*. The genes are grouped in ‘Tu_MCL’ clusters. The absolute expression level is shown as cyanine intensity, while genes for which no expression data are available are depicted as gray boxes. Superscripts supply additional info: ^{AD} protein identified in artificial diet fed upon by *T. urticae*; ^P expression level is significantly higher (FC \geq 8, adj. $p\leq$ 0.05) in the proterosoma versus the entire body of *T. urticae* adapted to bean plants; ^{ISH} gene confirmed to be expressed in the salivary glands using whole-mount *in situ* hybridizations.

Tu_MCL_45 has representatives in *T. urticae*, yet not in *T. evansi* and *T. lintearius* (TABLES 4.2 and S4.2). Protein tetur06g06630 was detected in artificial diet fed upon by *T. urticae*, while its encoding gene was shown to be expressed in the salivary glands (**Chapter 3**). Only tetur06g06650 appears to be significantly upregulated in the proterosoma (Benjamini-Hochberg adjusted $p\leq$ 0.05 and FC \geq 8).

Tu_MCL_63 has eight members in *T. urticae*, five of which are upregulated in the proterosoma (Benjamini-Hochberg adjusted $p\leq$ 0.05 and FC \geq 8). No representatives could be found in *T. evansi*, while two were present in *T. lintearius* (TABLES 4.2 and S4.2).

Proteins belonging to the spider mite-specific group **Tu_MCL_74** are homologous to proteins belonging to OG5_144177 and are further discussed in **Chapter 5** (SHOT family) (TABLES 4.2 and S4.2).

The other **Tu_MCL** groups (TABLES 4.2 and S4.2, Figure 4.4) include just a few proteins each, of which several were confirmed to be produced in the salivary glands using whole-mount *in situ* hybridizations (**Chapter 3**). Most of the genes coding for these proteins are significantly upregulated in the proterosoma. Many lack homologs in *T. evansi* or *T. lintearius* (TABLES 4.2 and S4.2).

4.4. DISCUSSION

4.4.1. The salivary protein composition of *T. urticae* is host plant-dependent

A host plant-dependent *T. urticae* salivary protein composition was evident from the proteomics data in **Chapter 3** and was studied in more detail in this chapter. To this purpose, genome wide expression analyses were per-

formed on the mite lines used for the salivary proteome study. Additionally, cotton-adapted lines were also investigated.

Secreting the full salivary protein repertoire, independent of the host plant, might come with an enhanced risk that some salivary proteins are perceived as elicitors. In addition, it is unlikely that all proteins will function optimally in all hosts as targets may differ across plants or may be absent. Therefore it would be beneficial for polyphagous herbivores to alter the composition of their saliva according to the nature of the host plant. Differences between the salivary gland transcriptomes of two populations of *Nilaparvata lugens*, a rice specialist, maintained on either a resistant or susceptible rice variety, were proposed to be related to different virulence traits of these brown planthoppers (Ji et al., 2013; Wang et al., 2015). Furthermore, drastically different salivary protein profiles were found when the western tarnished plant bug *Lygus hesperus* was fed artificial diet, cotton or pinto bean (Habibi et al., 2001). Hence, we investigated host plant specific secretion and expression of the mite's salivary genes. As suggested by the proteomics data (FIGURE 4.1), transcriptome analysis revealed that the expression of *T. urticae* salivary protein genes is strongly influenced by the host plant species on which the mites had been feeding (FIGURE 4.2). For example, proteins tetur28g01330 and tetur55g00040 were uniquely and abundantly identified from diet fed upon by tomato-adapted mites (FIGURE 4.1), while expression of the corresponding gene was also extremely high when feeding on tomato, relative to mites feeding on the other tested plants (FIGURE 4.2). Next to individual genes, we also studied OrthoMCL gene family groups. Tu_MCL_25, Tu_MCL_35 and Tu_MCL_36 (TABLES 3.1 and S3.7) all have members that are highly expressed when feeding on tomato, while the expression of other members of these groups was not influenced by the host plant (FIGURES 4.2 and 4.3). Future experiments should point out whether this is due to the fact that mites from the tomato-adapted lines had been maintained on tomato for many more generations than mites on the other hosts or whether this is due to the specific allelochemicals of tomato posing digestive or defensive challenges. Members of the OrthoMCL group OG5_144177 (SHOT family, **Chapter 5**) (TABLES 3.1 and S3.7) were, relative to mites on bean, expressed at lower levels in mites feeding from maize, cotton and tomato (FIGURE 4.2). When feeding on soy, however, these genes were expressed at slightly higher levels. A similar pattern can also be deduced from the proteomics data of mites adapted to bean and

soy versus mites adapted to maize and tomato. Bean and soy both are legume species (Fabaceae) and mites feeding on these plants probably encounter analogous plant secondary compounds that select for or induce a similar repertoire of salivary proteins. The observation that different host plant species can differentially affect expression levels of genes coding for salivary proteins – such as effectors – has been reported for aphids as well (Elzinga et al., 2014; Pan et al., 2015; Wang et al., 2015). Elzinga et al. (2014) suggested that the differential expression of salivary effector genes represents a strategy to avoid activation of defenses and to facilitate feeding.

In the scope of this discussion it is important to realize that only one *T. urticae* strain (London) was used. Although mites from this strain had been maintained on bean for many years, previous studies have shown that this population is both not fully inbred (Van Leeuwen et al., 2012) and capable of extensive transcriptional plasticity upon transfer to new hosts (Grbić et al., 2011; Dermauw et al., 2013; Wybouw et al., 2014; Zhurov et al., 2014), the latter of which has been further confirmed in this study. We believe therefore that this study has captured much of the repertoire of *T. urticae*'s biologically relevant salivary proteins. However, as marked variation of genotypes exists between *T. urticae* populations and across spider mite species (Kant et al., 2008; Alba et al., 2015), the use of additional *T. urticae* strains or different spider mite species may deliver additional salivary proteins. Identification via shotgun proteomics will, however, be less straightforward without reference genomes for these strains or species.

Due to the highly specific function of effectors, variable expression levels depending on the host plant could be indicative of such context-dependent function (Pan et al., 2015). The presence of R genes, which turn effectors effectively into elicitors, can differ among plant species or varieties (Stuart et al., 2012; Kanvil et al., 2014; Stuart, 2015), as does the presence of S genes (van Schie and Takken, 2014), the target of the effector. As such, the specific R and S gene composition of a specific host may determine the specific transcriptional response of a herbivore's secretome and thus its effector repertoire.

4.4.2. A glimpse into the salivary protein composition of *T. evansi* and *T. lintearius*

Tetranychus urticae and *T. lintearius* are closely related species which are more distantly related to *T. evansi* (Toni Gabaldon, personal communication

in Cao, 2014). Despite the close phylogenetic relationship of these three spider mites, their host ranges differ dramatically. *Tetranychus lintearius* is a specialist on *U. europaeus* (common gorse, Fabaceae), while the polyphagous *T. urticae* has been reported to feed on plants belonging to 140 different families (Ireson et al., 2003; Migeon and Dorkeld, 2006-2017). *Tetranychus evansi* is an oligophagous mite which is specialized in feeding on solanacean host plants such as tomato, although it has been recorded on other plant families as well (Migeon and Dorkeld, 2006-2017; Navajas et al., 2013). With the annotated genomes of these three species available to our research group, the occurrence of homologs of genes coding for (putative) *T. urticae* salivary proteins (**Chapter 3**, TABLE 3.1) in the genomes of *T. lintearius*, *T. evansi* and *T. urticae* could be investigated. It is important to keep in mind that proteins identified in the secreted saliva of *T. urticae* were used as a query. As such, ‘new’ salivary protein families, unique to *T. lintearius* and/or *T. evansi*, could not be identified, and we can only compare the number of members within the query families. With an efficient proteomics approach to identify spider mite salivary proteins reported in this PhD thesis (**Chapter 3**), it should be straightforward to determine the salivary proteome of *T. lintearius* and *T. evansi* as well, particularly because their genomes will be publically available soon. Encouragingly, preliminary proteomics experiments with *T. evansi* saliva have proven to be successful (data not shown). Not all NanoLC-MS/MS identified proteins used as search query (**Chapter 3**, TABLE 3.1) are true salivary proteins as some contamination is present. The proterosoma fold change (expression relative to the entire mite body) is an important criterion to evaluate whether these proteins are produced in the salivary glands or not. Of further notice, the proterosomal transcriptome (**Chapter 3**, DATA S3.9) was constructed using a *T. urticae* line (London) adapted to bean (*P. vulgaris*), and as such, these relative gene expression data may not be representative for genes with a strong host-dependent expression level (e.g., a gene that has a low expression level when feeding on bean will have a relatively low proterosoma to body expression ratio, which might have been high if a tomato-adapted line was used to generate the proterosoma transcriptome). Furthermore, proteins likely to be contamination – based on the proterosoma FC – were not omitted from TABLE 3.1 because these proteins, although contamination from, e.g., gut and cuticle, may still induce defenses when they end up in the plant tissue. As such, they may be relevant for other studies.

The identification of salivary protein orthologs among spider mite species is an important step in determining the protein's function during the plant-mite interaction. In aphids, for example, the functionality of *Myzus persicae* and *Acyrtosiphon pisum* orthologs of the effectors c002, PIntO1 (MP1) and PIntO2 has been evaluated (Pitino and Hogenhout, 2013). *Myzus persicae* is polyphagous and can count *Arabidopsis* amongst its host plants, while *A. pisum* specializes on colonizing fabacean hosts. Transgenic *Arabidopsis* lines producing the *M. persicae* orthologs of latter three effectors were found to increase the reproduction of *M. persicae* feeding on them. Reproduction of this aphid species did, however, not increase when they fed on *Arabidopsis* lines expressing the *A. pisum* orthologs of C002, PIntO1 or PIntO2 (Pitino and Hogenhout, 2013). This suggests that effector orthologs have evolved to enhance compatibility between the herbivore and a specific group of host plant species. This adaptation may, however, compromise the effector function in other plant species, potentially excluding them as hosts. Analogously, the host plant range of *T. urticae*, *T. lintearius* and *T. evansi* may be mirrored in the diversity of their effectors. For example, proteins encoded by the genes *tetur28g01330* and *tetur55g00040* (Tu_MCL_36) were abundantly and uniquely identified in salivary samples from tomato-adapted *T. urticae* lines. This was accompanied by a high gene expression level. The Solanaceae specialist *T. evansi* possesses four Tu_MCL_36 genes, while in the *Ulex* specialist *T. lintearius*, homologs were absent. This may be indicative for a Solanaceae-specific function of (most) Tu_MCL_36 members. However, a more detailed study is clearly needed to vouch for this hypothesis. Another example (Tu_MCL_74 & OG5_144177, 'SHOT' family) of host-dependent expression in *T. urticae*, and presence or absence of homologs in closely related spider mites species with a limited host range is studied in more detail in **Chapter 5**.

The comparative OrthoMCL study revealed that *T. urticae* possesses most proteins (436 proteins) homologous to proteins identified in artificial diet fed upon by *T. urticae*. Further, we identified 165 homologs in *T. evansi* and 159 in the closely related *T. lintearius*. As such, there is a correlation between the number of recorded host plants and the (predicted) number of putative salivary proteins. Whether this represents a causal relationship remains to be elucidated. But it would not come as a surprise that salivary protein homologs provide functional variability, allowing interaction with

components of distinct host plants. Alternatively, qualitative difference between the annotated genomes of the three mite species may be at the basis of the difference between the identified number of homologs. Indeed, the genome of *T. urticae* was sequenced using whole genome shotgun sequencing with Sanger technology (Grbić et al., 2011), while the genomes of *T. lintearius* and *T. evansi* were sequenced using Illumina technology (2012) (Cao, 2014). However, the EuGene gene prediction platform (Foissac et al., 2008) was used for genome annotation of all three species, using identical parameters. Furthermore, the size of the genomes and proteomes of all three mite species appeared to be about the same (Cao, 2014). So, despite the fact that the proteomes were not obtained by an identical procedure, a comparison seemed to be justified. In addition, a comparative analysis on the gustatory receptors of *T. urticae*, *T. lintearius* and *T. evansi* was based on the same genomic databases used in current study. It revealed an analogous correlation between the size of the host range and the number of these chemoreceptors, which have a function in the detection of chemical components in their environment, including those of plants (Ngoc et al., 2016).

Interestingly, genes encoding (predicted) salivary proteins belonging to the same (expanded) OrthoMCL cluster are often in close proximity of each other on the genome scaffold [e.g., *tetur20g00010-tetur20g00020-tetur20g00030-tetur20g00040-tetur20g00050* (Tu_MCL_12) and *tetli84g01040-tetli84g01050-tetli84g01060-tetli84g01070* (Tu_MCL_43)]. This may be a signature of (tandem) gene duplication followed by diversification, showing the effect of diversifying selection (Chen et al., 2010; Johnson et al., 2015; Zhao et al., 2015). The ubiquitous proliferations in salivary protein families in *T. urticae* may correlate with the polyphagous nature of this pest. One member of a family may for example be active in one host plant or group of related plants, while another member is active in another host plant (group). The few salivary protein families that are expanded in *T. evansi* and *T. lintearius* may reflect proliferation of a salivary protein family that is advantageous or even essential when feeding on the host plant group these mites have specialized to feed on. However, additional research is needed. Next to the identification of protein family expansions, an additional source of information about the ecological adaptation of the three spider mite species to their host plant range may be to look at pseudogenisation events (Ngoc et al., 2016). These pseudogenes are relics of former genes that no longer possess biological functions and can be

regarded as fossilized footprints of past gene expression. Most pseudogenisation events occur through random accumulation of mutations in genes on which the functional constraints are relaxed (Podlaha and Zhang, 2010). For example, *T. lintearius* may have lost salivary protein members since its divergence from the common ancestor with *T. urticae* because these genes are no longer needed due to the reduced host arsenal. An in-depth study of one family (SHOT) of salivary proteins (**Chapter 5**) further supports this hypothesis and we conclude that a spider mite's salivary protein repertoire is a reflection of its host plant range.

4.5. Conclusions

The saliva of *T. urticae* contains many proteins which appear to be spider mite specific and as such are likely to have a unique function. Several of these proteins are encoded by genes belonging to expanded families. In addition, the expression level of some of these putative salivary protein genes is influenced by the plant species to which the mite lines were adapted. This host dependent expression may be an indication of a host plant specific mode of action, which may be characteristic for effectors. Furthermore, *in silico* prediction of *T. evansi* and *T. lintearius* homologs of salivary proteins identified in *T. urticae* has shed some light on the salivary proteome of spider mite species with a more limited host range. It further supports our hypothesis that the host range of spider mites is linked to the salivary protein composition, and as such, expansions in *T. urticae* salivary protein gene families are likely key to *T. urticae*'s polyphagous nature.

Funding information

This project was supported by the European Commission (EC contract 618105) via FACCE ERA-NET Plus and FACCE-JP (Genomite, project ID 137), the Fund for Scientific Research Flanders (FWO) (grant G009312N to L.T. and T.V.L. and grant G053815N to L.T., T.V.L. and W.D.), The Special Research Fund (DOZA 01J131711 to W.J.) and NSF award DEB 1457346 (to R.M.C). R.G. was supported by National Institutes of Health Genetics Training Grant T32 GM07464. C.A.V. was supported by CONICYT BECAS (Chile) and M.R.K. by NWO (The Netherlands) (STW-VIDI 13492). N.W. was supported by a Marie Skłodowska-Curie Action (MSCA) Individual fellowship (658795-DOGMITE) of Horizon 2020. M.G. acknowledges funding by the Government of Canada through Genome

Canada and the Ontario Genomics Institute (OGI-046) and Ontario Research Fund- Global Leadership in Genomics and Life Sciences GL2-01-035. W.D. and G.M. are postdoctoral fellows of the Fund for Scientific Research Flanders (FWO). Some of the computational resources (Stevin Supercomputer Infrastructure) and services used in this work were provided by the VSC (Flemish Supercomputer Center), funded by Ghent University, the Hercules Foundation and the Flemish Government – department EWI.

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TABLE S4.1. *Tetranychus evansi*, *T. lintearius* and *T. urticae* proteins homologous to proteins identified in artificial diet fed upon by *T. urticae*. Only proteins which could be assigned to groups already existing in the OrthoMCL database were accounted for in this list.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_126560		tetev174g00270 tetev38g02260 tetev50g00620 tetev54g000330 tetev59g00640 tetev73g000390	tetli08g00930 tetli38g00560 tetli78g00410	tetur01g09300 tetur02g10210 tetur03g08060 ^P tetur05g07800 tetur12g01140 tetur12g02880 tetur24g01640
OG5_126583	IPR023210 (NADP-dependent oxidoreductase domain), IPR018170 (Aldo/keto reductase, conserved site), IPR020471 (Aldo/keto reductase)	tetev275g00070 tetev33g00180 tetev43g00610 tetev43g00640 tetev43g00670 tetev43g00690 tetev89g00430	tetli105g00100 tetli154g00030 tetli216g00070 tetli231g00220 tetli23g02180 tetli333g00010	tetur02g07450 tetur02g07550 tetur02g07590 tetur02g11330 tetur02g11340 ^{AD,P} tetur02g11390 tetur02g11410 tetur02g11530 tetur02g90723 tetur30g00660
OG5_126595	IPR004000 (Actin family), IPR004001 (Actin, conserved site), IPR020902 (Actin/actin-like conserved site)	tetev109g00360 tetev26g00150	tetli04g00180 tetli246g00020	tetur03g09480 ^{AD,P} tetur09g05350 tetur09g05360
OG5_126607	IPR013201 (Cathepsin propeptide inhibitor domain; I29), IPR025660 (Cysteine peptidase, histidine active site), IPR025661 (Cysteine peptidase, asparagine active site), IPR000668	tetev02g00220 tetev28g00730 tetev321g00030 tetev36g00820 tetev36g00830	tetli09g00600 tetli33g01670 tetli344g00080 tetli38g01500 tetli38g01510	tetur05g92633 tetur06g03520 tetur09g04400 ^{AD} tetur09g04420 tetur09g04470

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
	(Peptidase C1A, papain C-terminal), IPR000169 (Cysteine peptidase, cysteine active site)	tetev36g00840 tetev36g00850 tetev36g00860 tetev36g00870 tetev36g00880 tetev38g02400 tetev85g00370	tetli38g01520 tetli38g01530 tetli38g01540 tetli38g01550	tetur123g00050 tetur12g01810 tetur12g01820 tetur12g01830 tetur12g01840 tetur12g01850 ^P tetur12g01860 ^{AD} tetur12g04631 tetur16g03680 tetur16g03770 tetur23g00050 tetur23g00860 tetur23g00880 tetur23g00890 tetur23g01290 tetur25g00650 ^{AD}
OG5_126738	IPR013783 (Immunoglobulin-like fold), IPR007110 (Immunoglobulin-like domain), IPR003961 (Fibronectin type III), IPR000719 (Protein kinase domain), IPR008271 (Serine/threonine-protein kinase, active site), IPR017441 (Protein kinase, ATP binding site), IPR013098 (Immunoglobulin I-set), IPR002290 (Serine/threonine/dual specificity protein kinase, catalytic domain), IPR003599 (Immunoglobulin subtype), IPR003598	tetev04g01160 tetev101g00190 tetev101g00220 tetev124g00030 tetev124g00040 tetev169g00060 tetev169g00210 tetev169g00220	tetli260g00310 tetli260g00320 tetli26g02110 tetli26g02120 tetli26g02130 tetli302g00020 tetli321g00010 tetli46g00640 tetli478g00030 tetli69g00060	tetur01g01810 tetur01g01830 tetur01g01850 ^{AD} tetur04g02800 tetur04g02880 tetur05g03810 tetur06g00290 tetur114g00010 tetur30g00590 tetur31g01290

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_126942	(Immunoglobulin subtype 2), IPR011009 (Protein kinase-like domain) IPR012336 (Thioredoxin-like fold), IPR010987 (Glutathione S-transferase, C-terminal-like), IPR004045 (Glutathione S-transferase, N-terminal), IPR004046 (Glutathione S-transferase, C-terminal)	tetev125g00360 tetev125g00410 tetev25g01600 tetev25g01620 tetev52g01110 tetev88g01280	tetli118g00150 tetli118g00180 tetli118g00190 tetli118g00200 tetli16g01140 tetli16g01220 tetli26g01650 tetli29g00100	tetur01g02230 tetur01g02470 tetur01g02480 tetur01g02490 tetur01g02500 tetur01g02510 ^{AD,P} tetur03g07920 ^{AD} tetur26g01450 tetur26g01460 tetur26g01490 tetur26g01500 tetur26g02801 tetur26g02802 tetur29g00220 tetur31g01330 tetur07g03440 ^{AD} tetur109g00010 tetur109g00020 tetur01g16010 tetur02g04630 tetur02g11180 tetur04g09509 ^P tetur07g03760 tetur09g01550 tetur26g02320 ^{AD}
OG5_127143	IPR013785 (Aldolase-type TIM barrel), IPR029768 (Fructose-bisphosphate aldolase class-I active site), IPR000741 (Fructose-bisphosphate aldolase, class-I)	tetev258g00130 tetev258g00160	tetli162g00220 tetli162g00240	
OG5_127584	IPR001424 (Superoxide dismutase, copper/ zinc binding domain), IPR018152 (Superoxide dismutase, copper/zinc, binding site)	tetev11g00600 tetev11g00760 tetev14g00420 tetev16g00390 tetev204g00040 tetev38g02250 tetev43g00880	tetli07g00150 tetli125g00040 tetli147g00210 tetli147g00500 tetli149g01060 tetli168g00290 tetli52g00820	

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_127620	IPR013780 (Glycosyl hydrolase, family 13, all-beta), IPR002241 (Glycoside hydrolase, family 27), IPR013785 (Aldolase-type TIM barrel), IPR000111 (Glycoside hydrolase family 27/36, conserved site), IPR017853 (Glycoside hydrolase super-family)	tetev44g01030 tetev05g00340 tetev98g00090	tetli80g00540 tetli02g00890 tetli94g00190	tetur26g02520 tetur02g04310AD tetur04g00130
OG5_127800	IPR000668 (Peptidase C1A, papain C-terminal)	tetev12g02780 tetev176g00050 tetev183g00220 tetev32g00870 tetev33g00190 tetev407g00010 tetev488g00020 tetev650g00030 tetev92g00890	tetli105g00130 tetli179g00030 tetli179g00040 tetli179g00350 tetli179g00580 tetli256g00020 tetli256g00230 tetli256g00240 tetli268g00120 tetli48g01490	tetur01g05230AD tetur01g05480 tetur01g16463 tetur01g16473 tetur02g11420 tetur03g07930 tetur03g07950 tetur03g08010 tetur03g08020 tetur03g08030AD tetur03g09997 etur06g02570 tetur06g02580AD tetur08g05010 tetur08g05020 tetur08g05030 tetur08g05310 tetur09g00570 tetur09g00600

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_128075	IPR002347 (Glucose/ribitol dehydrogenase), IPR002198 (Short-chain dehydrogenase/reductase SDR), IPR020904 (Short-chain dehydrogenase/reductase, conserved site), IPR016040 (NAD(P)-binding domain)	tetev61g00450 tetev61g00560	tetli84g00100 tetli84g00140	tetur24g00270 tetur24g00280 tetur28g01340 tetur28g01390 tetur28g01400 tetur28g01420 tetur28g01430 tetur55g00100 tetur55g00120 tetur55g00130P tetur32g01960 tetur32g02180 tetur32g02327AD
OG5_128163	IPR001944 (Glycoside hydrolase, family 35), IPR008979 (Galactose-binding domain-like), IPR017853 (Glycoside hydrolase superfamily), IPR019801 (Glycoside hydrolase, family 35, conserved site), IPR013781 (Glycoside hydrolase, catalytic domain), IPR031330 (Glycoside hydrolase 35, catalytic domain), IPR026283 (Beta-galactosidase 1-like)	tetev10g00930 tetev15g00040 tetev178g00370	tetli108g00380 tetli24g00700 tetli31g00290	tetur07g07380AD tetur09g05870 tetur13g04220 tetur13g04678
OG5_128170	IPR016040 (NAD(P)-binding domain), IPR002347 (Glucose/ribitol dehydrogenase)	tetev38g02000 tetev92g00590	tetli04g01370 tetli287g00100	tetur06g04880 tetur06g04890

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
	nase), IPR020904 (Short-chain dehydrogenase/reductase, conserved site), IPR002198 (Short-chain dehydrogenase/reductase SDR)		tetli29g01000	tetur06g04900 tetur06g04970 tetur06g04980 tetur06g04990 tetur06g05070 tetur06g05090 tetur07g07300 tetur07g07310 tetur106g00010 tetur106g00040 tetur13g04450 tetur28g01070 tetur28g01140 tetur28g01710 tetur28g01720 ^{AD} tetur28g01730 tetur28g01740 tetur29g00870 ^P tetur02g02400 tetur16g03420 ^{AD,P} tetur16g03430 ^P tetur28g00360 ^{AD,ISH,P}
OG5_129300	IPR013781 (Glycoside hydrolase, catalytic domain), IPR008979 (Galactose-binding domain-like), IPR006104 (Glycosyl hydrolases family 2, sugar binding domain), IPR017853 (Glycoside hydrolase superfamily)	tetev16g01180 tetev16g01190 tetev69g00510	tetli94g00590	
OG5_129423	IPR013780 (Glycosyl hydrolase, family 13, all-beta), IPR001139 (Glycoside hydrolase, family 30), IPR013781 (Glycoside hydrolase, catalytic domain), IPR017853 (Glyco-	tetev10g00050 tetev193g00310 tetev213g00090 tetev246g00020	tetli112g00490 tetli112g00550 tetli134g00100 tetli164g00130	tetur02g07230 tetur02g07250 tetur02g07770 tetur02g07780

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
	side hydrolase superfamily)	tetev291g00040	tetli168g00350	tetur02g10940
		tetev299g00050	tetli16g01210	tetur02g12930 ^{AD}
		tetev299g00100	tetli186g00110	tetur02g90714
		tetev370g00030	tetli236g00110	tetur04g00790
		tetev39g00060	tetli237g00180	tetur05g06960
		tetev52g00390	tetli41g00220	tetur05g07000
		tetev59g00540	tetli41g00260	tetur06g01570
		tetev689g00010	tetli70g01270	tetur06g01580
		tetev689g00020	tetli87g00560	tetur06g01590
		tetev85g00380	tetli95g00230	tetur06g01660
		tetev89g00470		tetur06g01690
				tetur06g01700
				tetur09g06680 ^P
				tetur147g00020 ^{AD}
				tetur147g00030
				tetur147g00040
				tetur17g00790
				tetur17g00830
				tetur17g00850
				tetur17g00870
				tetur187g00010
				tetur187g00020
				tetur24g02260 ^P
				tetur30g02260
				tetur31g01370
				tetur32g00060
				tetur33g01260

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_130246	IPR029058 (Alpha/Beta hydrolase fold), IPR008758 (Peptidase S28)	tetev75g00330	tetli03g01710	tetur33g01270 tetur33g01290 tetur72g00010 ^P tetur72g00020 ^P tetur19g01900 ^{AD}
OG5_130527	IPR012674 (Calycin), IPR022271 (Lipo- calin, ApoD type), IPR011038 (Calycin- like), IPR000566 (Lipocalin/cytosolic fatty- acid binding domain), IPR003057 (Invertebrate colouration protein)	tetev01g00490 tetev06g00100 tetev124g00310 tetev124g00320 tetev124g00340 tetev147g00020 tetev14g00610 tetev14g00630 tetev14g00640 tetev164g00040 tetev164g00070 tetev210g00200 tetev210g00210 tetev210g00220 tetev299g00070 tetev52g00260 tetev83g000450	tetli07g01010 tetli07g01020 tetli102g00300 tetli112g00400 tetli139g00390 tetli13g00660 tetli145g00130 tetli145g00300 tetli208g00170 tetli358g00010 tetli41g00210 tetli80g00390 tetli80g00400	tetur01g01510 tetur01g05730 tetur01g05740 tetur01g05750 tetur01g05770 tetur01g05880 tetur04g05970 tetur04g05980 ^{AD} tetur04g06000 tetur04g06010 tetur05g07070 tetur06g01640 ^{AD} tetur06g02130 tetur06g02140 tetur06g02670 tetur06g03020 tetur06g03090 tetur06g03100 tetur06g03340 tetur06g03350 tetur06g03360

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lincolni</i> protein ID	<i>T. urticae</i> protein ID
				tetur06g03370
				tetur06g03440
				tetur06g03550
				tetur06g03860 ^P
				tetur06g06691
				tetur06g91377
				tetur06g91385
				tetur07g03970
				tetur09g04720
				tetur09g04730
				tetur09g04920
				tetur11g05210
				tetur11g05230
				tetur16g03410
				tetur16g03450
				tetur16g03460
				tetur174g00050
				tetur18g00900
				tetur282g00020
				tetur30g01430
				tetur31g00670
				tetur31g00680
				tetur31g00710
				tetur31g00730
				tetur31g00780
				tetur31g00800
				tetur31g00880

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_131746	IPR016040 (NAD(P)-binding domain)	tetev116g00460 tetev22g00600	tetli15g00980 tetli33g01300 tetli93g00560	tetur31g00900 tetur31g00920 tetur31g01880 tetur37g00940 ^P tetur01g12240 tetur23g01640 ^{AD}
OG5_132251	IPR001846 (von Willebrand factor, type D domain)		tetli04g00850 tetli04g00860 tetli07g01670 tetli07g01700	tetur07g05480 tetur39g00700 tetur39g00720 tetur39g00810 ^{AD}
OG5_133467	IPR003172 (MD-2-related lipid-recognition domain), IPR014756 (Immunoglobulin E-set)	tetev107g00370 tetev107g00380 tetev107g00390 tetev107g00400 tetev13g01150 tetev25g01430 tetev52g01260 tetev95g00110	tetli1323g00010 tetli571g00010 tetli01g01890 tetli132g00590 tetli16g00920 tetli25g00680 tetli318g00040 tetli51g00500 tetli85g00410	tetur43g00010 ^{AD} tetur516g00020 ^{AD} tetur03g01850 tetur04g08680 tetur07g02990 tetur13g92931 tetur140g00020 tetur14g01690 tetur14g01700 tetur14g02060 tetur14g02070 ^{AD} tetur14g02080 ^{AD} tetur14g02090 ^{AD} tetur17g03530 tetur17g03550 tetur28g01370

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_134456	IPR002223 (Pancreatic trypsin inhibitor Kunitz domain), IPR020901 (Proteinase inhibitor I2, Kunitz, conserved site)	tetev232g00090	tetli06g00480	tetur55g00080 tetur13g03820AD
OG5_135950	IPR001254 (Serine proteases, trypsin domain), IPR009003 (Peptidase S1, PA clan), IPR018114 (Peptidase S1, trypsin family, active site), IPR001314 (Peptidase S1A, chymotrypsin-type)	tetev13g01730 tetev30g00470 tetev83g00070	tetli06g01170 tetli115g00180 tetli154g00160 tetli28g00930 tetli79g00510	tetur06g06761P tetur07g00150AD,ISH,P tetur09g00280P tetur35g00910 tetur35g01000
OG5_136350	IPR019826 (Carboxylesterase type B, active site), IPR029058 (Alpha/Beta hydro-lase fold), IPR002018 (Carboxylesterase, type B)	tetev492g00030 tetev56g01620 tetev59g00690 tetev97g00030	tetli72g00640 tetli72g00650 tetli72g00720	tetur02g09310 tetur02g14551 tetur02g15189 tetur11g01500AD tetur207g00020 tetur29g00950
OG5_141111	IPR001254 (Serine proteases, trypsin domain), IPR009003 (Peptidase S1, PA clan)	tetev136g00120 tetev18g00990 tetev194g00150	tetli10g00400 tetli51g00300	tetur13g00200 tetur16g03190AD,P
OG5_144177	'Secreted host responsive protein in Tetranychidae' (SHOT) family (Chapter 5)			tetur03g03670AD,P tetur03g03680AD,P tetur03g03690ISH,P tetur03g03700AD,ISH,P tetur03g03730P tetur03g10093P
OG5_147492	IPR000010 (Cystatin domain)	tetev08g00660 tetev08g00670	tetli144g00040 tetli145g00170	tetur06g01060AD tetur06g06610

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_149533	IPR009003 (Peptidase S1, PA clan), IPR001254 (Serine proteases, trypsin domain), IPR001314 (Peptidase S1A, chymotrypsin-type), IPR018114 (Peptidase S1, trypsin family, active site)	tetev100g00570 tetev116g00510 tetev292g00240 tetev37g01130 tetev37g01260 tetev49g00470	tetli37g00030 tetli38g00430 tetli18g00950 tetli50g00280	tetur09g03620 ^{AD} tetur09g03650 tetur09g04770 tetur05g08340 tetur07g08034 ^{AD} tetur35g00970 tetur35g01050 tetur35g01250 tetur35g01260 tetur35g01270 tetur35g01310 tetur35g01540 tetur35g01600 tetur35g01610 tetur35g01630 tetur441g00030 tetur07g03160 tetur16g03470 ^{AD,P} tetur30g01440 ^{AD}
OG5_152237	IPR009003 (Peptidase S1, PA clan), IPR001254 (Serine proteases, trypsin domain)	tetev229g00140	tetli51g00220	
OG5_152337	IPR009003 (Peptidase S1, PA clan), IPR018114 (Peptidase S1, trypsin family, active site), IPR001254 (Serine proteases, trypsin domain), IPR001314 (Peptidase S1A, chymotrypsin-type)	tetev38g00570 tetev83g00100	tetli208g00160	
OG5_152454	IPR002557 (Chitin binding domain), IPR029070 (Chitinase insertion domain), IPR01583 (Chitinase II), IPR001579 (Glycoside hydrolase, chitinase active site),	tetev08g01790 tetev22g00350 tetev31g00080	tetli179g00520 tetli184g00230 tetli21g00780	tetur01g11910 ^{AD,P} tetur08g05470 tetur11g03440

TABLE S4.1. Continued.

OrthoMCL	Functional description	<i>T. evansi</i> protein ID	<i>T. lintearius</i> protein ID	<i>T. urticae</i> protein ID
OG5_158831	IPR013781 (Glycoside hydrolase, catalytic domain), IPR017853 (Glycoside hydrolase superfamily), IPR001223 (Glycoside hydrolase family 18, catalytic domain)			tetur01g02650 tetur01g02670 ^{AD}
OG5_168371	IPR003172 (MD-2-related lipid-recognition domain), IPR014756 (Immunoglobulin E-set) IPR001254 (Serine proteases, trypsin domain), IPR001314 (Peptidase S1A, chymotrypsin-type), IPR018114 (Peptidase S1, trypsin family, active site), IPR009003 (Peptidase S1, PA clan)			tetur09g03880 ^{AD,P} tetur12g03940 ^{AD,P} tetur12g03950 ^P tetur66g00060 tetur66g00070 tetur22g00260 ^{AD}
OG5_176862	IPR002919 (Trypsin Inhibitor-like, cysteine rich domain)			tetur22g00290 ^P
OG5_207753	IPR002919 (Trypsin Inhibitor-like, cysteine rich domain)			tetur40g00392 ^{AD,ISH,P}

^{AD} Protein identified in artificial diet fed upon by *T. urticae* (**Chapter 3**), ^{ISH} Gene coding for the protein is confirmed to be expressed in the salivary glands using whole-mount *in situ* hybridizations (**Chapter 2, 3 & 5**), ^P Expression level of the gene coding for the protein is significantly higher (FC \geq 8, adj. $p\leq$ 0.05) in the proterosome versus the entire body of *T. urticae* adapted to bean plants (**Chapter 3**, TABLE S3.9)

TABLE S4.2. Spider mite-specific *Tetranychus evansi*, *T. lintearius* and *T. urticae* proteins homologous to proteins identified in artificial diet fed upon by *T. urticae*.

OrthoMCL group	<i>T. evansi</i>	<i>T. lintearius</i>	<i>T. urticae</i>
Tu_MCL_12	tetev485g00030	tetli56g00590	tetur02g14280
			tetur02g14420
			tetur02g14470
			tetur08g03060
			tetur12g00580
			tetur13g04230 ^P
			tetur19g92445
			tetur20g00010
			tetur20g00020
			tetur20g00030
			tetur20g00040
			tetur20g00050 ^P
			tetur20g00090 ^P
			tetur20g00100
			tetur20g00540 ^P
			tetur20g00550
			tetur20g00560 ^{AD, P, ISH}
			tetur19g00950
			tetur20g90779
			tetur20g90780
			tetur20g90781
			tetur19g02210
			tetur20g00670
			tetur47g00120
			tetur47g00170
			tetur43g00560
			tetur43g00570
			tetur43g00580
			tetur58g00010
			tetur58g00020
			tetur58g00030
			tetur58g00050
			tetur58g00090
tetur58g00100			
tetur58g00110			
tetur58g90066			
tetur58g90067			
Tu_MCL_25	tetev52g01490	tetli16g00750	tetur03g07610
			tetur03g07590
			tetur14g02720 ^P
			tetur31g00630 ^{AD, P}
			tetur31g00690 ^P
			tetur31g00740 ^P
			tetur31g00770

TABLE S4.2. Continued.

OrthoMCL group	<i>T. evansi</i>	<i>T. lintearius</i>	<i>T. urticae</i>
			tetur31g00810
			tetur31g00820
			tetur31g00830 ^{AD}
			tetur31g00850
			tetur31g00890
			tetur31g00910
			tetur31g01040 ^{AD, P}
			tetur31g01060 ^{AD}
			tetur31g01860
Tu_MCL_25	tetev219g00200	tetli16g00760	tetur06g00230 ^{AD, P, ISH}
		tetli174g00060	
		tetli174g00050	
Tu_MCL_43	tetev148g00350	tetli20g03110	tetur03g00040 ^P
	tetev102g00780	tetli84g01040	tetur03g00050 ^P
		tetli84g01050	tetur03g00070 ^P
		tetli84g01060	tetur08g08070 ^P
		tetli84g01070	tetur32g01880 ^{AD, P}
		tetli84g00020	tetur33g00840
			tetur33g00850 ^P
			tetur33g00860
			tetur33g00870 ^P
Tu_MCL_35	tetev446g00020		tetur01g16564
	tetev370g00040		tetur04g09459
	tetev54g00740		tetur04g09479 ^{AD, P}
			tetur04g09489
			tetur04g00770
			tetur12g00700
			tetur143g00020
			tetur32g02287
			tetur32g02297
			tetur32g00050 ^{AD, P, ISH}
			tetur44g00291
			tetur44g00301
			tetur86g00050
			tetur86g00060
Tu_MCL_36	tetev13g01130		tetur01g10080
	tetev187g00210		tetur01g10090
	tetev192g00290		tetur05g04560 ^{AD, P}
	tetev317g00050		tetur05g04580 ^{AD, P, ISH}
			tetur16g90599
			tetur196g90150
			tetur28g01330 ^{AD, P}
			tetur28g01360
			tetur28g01380
			tetur28g01410

TABLE S4.2. Continued.

OrthoMCL group	<i>T. evansi</i>	<i>T. lintearius</i>	<i>T. urticae</i>
			tetur55g00040 ^{AD, P}
Tu_MCL_45			tetur55g00090
			tetur02g06500
			tetur02g06490
			tetur02g06520
			tetur02g06560
			tetur02g06600
			tetur02g92468
			tetur04g08190
			tetur06g06630 ^{AD, ISH}
			tetur06g06650 ^P
			tetur06g03000
			tetur307g00010
			tetur307g00020
Tu_MCL_63		tetli115g00430	tetur14g03160 ^{AD}
		tetli115g00400	tetur14g03170 ^P
			tetur14g03240 ^P
			tetur14g03250
			tetur14g03560
			tetur14g03430 ^P
			tetur14g02600 ^P
			tetur517g00010 ^P
Tu_MCL_74	tetev340g00035	tetli74g00350	tetur03g03620
	tetev340g00030		tetur03g10083 ^{AD, P}
			tetur11g01360 ^{P, ISH}
			tetur11g06390 ^P
			tetur11g06400 ^P
Tu_MCL_211	tetev132g00350	tetli11g00110	tetur01g00950 ^{AD, P, ISH}
	tetev132g00300	tetli11g00160	tetur01g01000 ^P
Tu_MCL_212	tetev132g00290	tetli11g00100	tetur01g00940 ^{AD, P, ISH}
		tetli11g00170	tetur01g01010 ^{AD, P}
Tu_MCL_153	tetev252g00100	tetli223g00110	tetur03g04460
		tetli223g00120	tetur03g04470 ^{AD}
No group	tetev06g00640	tetli23g01520	tetur07g01660 ^{AD, P}
		tetli23g01540	
		tetli66g00240	
No group	tetev223g00110		tetur06g00510 ^{AD, P}
	tetev01g02160		tetur29g01360 ^{AD, P, ISH}
No group	tetev117g00420	tetli101g00190	tetur03g08710 ^{AD, P, ISH}
No group			tetur13g00610 ^P
			tetur13g00600 ^{AD, P}
			tetur13g00650 ^P
No group			tetur09g00900 ^{AD, P}
			tetur03g10103 ^P
No group	tetev168g00210		tetur10g00090 ^{AD, P, ISH}

TABLE S4.2. Continued.

OrthoMCL group	<i>T. evansi</i>	<i>T. lintearius</i>	<i>T. urticae</i>
No group	tetev168g00220		tetur10g00100 ^{AD, P}
OG5_126560	tetev64g00950		tetur20g01290 ^{AD}

^{AD} Protein identified in artificial diet fed upon by *T. urticae* (**Chapter 3**)

^P Expression level of the gene coding for the protein is significantly higher (FC \geq 8, adj. $p\leq$ 0.05) in the proterosoma versus the entire body of *T. urticae* adapted to bean plants (**Chapter 3**, TABLE S3.9)

^{ISH} Gene coding for the protein is confirmed to be expressed in the salivary glands with whole-mount *in situ* hybridizations (**Chapter 3 & Chapter 5**)