A proteomic insight into vitellogenesis during tick ovary maturation

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Ticks are arthropod ectoparasites of importance for public and veterinary health. The understanding of tick oogenesis and embryogenesis could contribute to the development of novel control methods. However, to date, studies on the temporal dynamics of proteins during ovary development were not reported. In the present study we followed protein profile during ovary maturation. Proteomic analysis of ovary extracts was performed by liquid chromatography-tandem mass spectrometry (LC-MS/MS) using shotgun strategy, in addition to dimethyl labelling-based protein quantification. A total of 3,756 proteins were identified, which were functionally annotated into 30 categories. Circa 80% of the annotated proteins belong to categories related to basal metabolism, such as protein synthesis and modification machineries, nuclear regulation, cytoskeleton, proteasome machinery, transcriptional machinery, energetic metabolism, extracellular matrix/cell adhesion, immunity, oxidation/detoxification metabolism, signal transduction, and storage. The abundance of selected proteins involved in yolk uptake and degradation, as well as vitellin accumulation during ovary maturation, was assessed using dimethyl-labelling quantification. In conclusion, proteins identified in this study provide a framework for future studies to elucidate tick development and validate candidate targets for novel control methods.

Ticks are hematophagous ectoparasites of importance for human and veterinary health, due to direct effects of the parasitism, and the pathogenic agents they can transmit1. During development, ticks go through four stages of life: egg, larva, nymph and adult. Members of the family Ixodidae undergo either one-host, two-host or three-host life cycles. Particularly, Rhipicephalus microplus, a major cattle parasite2 is a one-host tick; it remains on the same host throughout larval, nymph and adult stages, only leaving the host prior to laying eggs3. R. microplus attaches to the cattle host as unfed larvae, and then proceeds to feed and moult through nymphal and immature adult stages, during a period of about 12 days. After mating, adult females start the slow feeding phase, when they swallow moderate amounts of blood for about 6 days3. The rapid engorgement phase takes place on the last 3 days on the host, when most of the blood is ingested. After completing the blood meal, the fully engorged females detach from the host and perform oviposition on the soil4.

Oogenesis is the development of female sex cells (oocytes) in the ovaries, while vitellogenesis is the process whereby ovarian and extracellular tissues produce protein precursors and other molecules which are transported to and accumulated inside the oocytes5. In Ixodidea ticks, vitellogenin (Vg) is synthesised as a high molecular mass-precursor in fat body, gut and ovaries6–9. Vg is released into the haemolymph to be taken up by oocytes via a receptor mediated endocytosis10, and then accumulates inside the oocytes yolk granules, being thereafter named vitellin (Vt). Together with other yolk components, Vt is a source of nutrients for embryo development11.
Biological function depends on the interplay of gene and protein regulation, influenced by extracellular factors. Therefore, a comprehensive understanding of the profile of gene expression and the proteins present throughout ovary development is necessary to elucidate the events occurring during arthropod development. Transcriptional profile analysis has been widely used in insect research\textsuperscript{13} and the gene expression profiles of several insect embryos during the early developmental stages have been characterised\textsuperscript{13,14}. Nevertheless, there are few proteomic studies addressing the molecular basis of developmental regulation in arthropods. Moreover, despite its importance, tick ovary has not been studied in detail and, unfortunately, there is a paucity of reports showing the successive alterations in protein profile during tick ovary maturation. In this sense, various aspects of *R. microplus* ovary maturation have been addressed at the level of individual genes and/or proteins, which is insufficient to uncover the underlying mechanisms of ovary development\textsuperscript{15}. In arthropods, comparative proteomic approaches were applied to analyse reproduction strategies used by the parasitoid wasp *Venturia canescens*\textsuperscript{16}, to compare activated and inactivated ovaries from workers bees *Apis mellifera*\textsuperscript{17}, and different ovary stages in the shrimp *Metapenaeus ensis*\textsuperscript{18}, and also to study differences in ovaries of domestic and wild shrimps *Penaeus monodon*\textsuperscript{19}.

The study of *R. microplus* embryogenesis\textsuperscript{20} can contribute to a better understanding of embryo metabolism and development in ticks, as well as in other arthropods. Despite the importance of this parasite, no proteomic study on ovary development of *R. microplus* has been reported so far. Until now, there are only two comparative studies addressing tick ovary using proteomics\textsuperscript{21} or transcriptomics\textsuperscript{22}, but these studies are focused on *Babesia bovis* infection. In the present study, proteomic profiles were analysed in whole tissue extracts of tick ovary representing sequential stages of ovary maturation. Samples were analysed by liquid chromatography-tandem mass spectrometry (LC-MS/MS) based on shotgun strategy. In addition, dimethyl labelling-based protein quantification was used to identify differences in protein variation. In summary, the results shown here provide a clearer view on the post-transcriptional events in *R. microplus* ovary maturation, pointing out protein modulation during ovary development. This work provides a better understanding of how tick ovary develops, and how protein synthesis is regulated during embryogenesis. Consequently, this knowledge offers opportunities for advancing novel targets of tick control methods, since the inhibition of protein function may result in a decreased viability of the eggs, or even in premature mortality.

**Results and Discussion**

**Ovary development.** Oocyte formation is the initial step in embryonic development and, like ovary maturation, is heavily affected by blood ingestion and consequently by female weight. Balashov\textsuperscript{23} proposed a categorisation of tick oogenesis according to individual oocyte development. However, oocyte asynchrony is determined from the beginning of yolk uptake, and it is possible to find all Balashov-oocyte stages occurring at the same time during ovarian maturation\textsuperscript{24}. Thus, using *Ambyomma hebraeum* tick as a model, Seixas et al.\textsuperscript{25} proposed a new classification that comprises the degree of oocyte development and the size of the ovary. According to this classification, ovarian growth phase (OGP) 1 is characterised by a thin and translucent ovary with visible oocyte nuclei. In OGP 2, ovaries are longer and thicker than in OGP 1, and oocytes are opaque with no visible nuclei, while in OGP 3 many oocytes acquire a brown colour due to yolk granules accumulation. More like a pre-ovation stage, in OGP 4 large and yolk-filled oocytes prevail, and yolk spheres are visible. In OGP 5, when in *A. hebraeum* oviposition begins, the ovary appearance is the same as in OGP 4, but ovulated oocytes are visible in the ovary lumen\textsuperscript{25}. It is important to note that *A. hebraeum* females take 10 days after detachment from the host to start oviposition. In contrast, in *R. microplus* females this period takes only 3 days. Nevertheless, the OGP system classification is useful to normalise ovary maturation period, thus allowing to compare protein levels among female ovaries.

**Tick ovary proteome overview.** In this work, ovaries from *R. microplus* partially engorged females (PEF) and fully engorged females (FEF), which were manually collected, were analysed to perform a temporal proteomic study. PEF were divided in eight groups according to average weight of individuals (as an indicative of blood meal acquisition and ovary maturation) ranging between 10 and 270 mg (groups PEF-10 to PEF-270) (Fig. 1). FEF were grouped according to days post-detachment (FEF-D1 to FEF-D4) (details in section “Ovary extracts and protein digestion” in Materials and Methods). These groups represent sequential stages of ovary maturation and oogenesis. The electrophoretic profiles of PEF and FEF ovary proteins reveal a clear difference in the amount of some proteins (Fig. 2). As ovaries mature, vitellin (Vt) accumulation (polypeptides ≥55 kDa) is visible, especially in late vitellogenesis (after PEF-53) and in FEF groups. Also, the profiles show that the number of polypeptides ≤55 kDa decreases as ovary develops. These electrophoretic patterns correlate with microscopy observations (Fig. 1), where ovaries go from off-white to a brownish colour, due to the amount of Vt-containing yolk granules (Vt has a typical colour due to its haeme content). To contemplate a whole panel of changes in the protein profile observed during ovary development until the end of oogenesis, experimental groups PEF-10, -24, -35, -53, -84, -189, and -270, and FEF-D1 and -D3 were subjected to proteomics analyses (Fig. 2).

Altogether, a total of 3,756 unique proteins were found in the proteome of *R. microplus* ovaries. Among them, 3,335 proteins were identified using normalised spectral abundance factor (NSAF) semi-quantitative approach, while 2,883 proteins were identified by dimethyl labelling quantitative method. From the proteins detected by dimethyl labelling, there are 421 unique proteins when compared to NSAF. Thus, 2,462 proteins were detected in common by NSAF and dimethyl labelling (Fig. 3; see Supplementary Tables S1, S2 and S3). Dimethyl-labelled proteins with a p value and ratio variance ≥0.06 were excluded from the analysis. Through NSAF semi-quantification, more than 2,000 proteins were identified in PEF-24, -35 and -270 groups; more than 1,000 in PEF-10, -53 and -189 and FEF-D1 groups; and more than 800 proteins in FEF-D3 group (Fig. 4 and Supplementary Table S2). Despite a distant phylogenetic relationship, the number of proteins described here in *R. microplus* is similar to what was reported in a comparative study of ovarian proteome performed with the shrimp.*
Penaeus monodon, which found a total of 1,638 proteins\textsuperscript{19}. Also similarly to the present study, Velentzas et al.\textsuperscript{26} reported the descriptive proteome of Drosophila melanogaster ovary, in which 2,103 proteins were found.

After mating, ovary engages in oogenesis to produce eggs that will provide embryo development; therefore, this organ is in constant physiological and biochemical modification. Broadly, in all ovary stages circa 80% of the proteins are involved in basal metabolism. These proteins are clustered as: synthesis and modification machinery, nuclear regulation, cytoskeletal proteins, proteasome machinery, transcription machinery, energetic metabolism, extracellular matrix/cell adhesion, immunity, oxidation/detoxification metabolism, signal transduction, and storage. Also, the NSAF overview highlights that protein amount in some clusters changes over time during the temporal analysis performed. Proteins related to transcriptional machinery, protein synthesis machinery,
and nuclear regulation show a decrease in abundance as ovaries develop, while storage proteins are upregulated (Fig. 4).

The temporal variation among protein levels was also supported by heat map analysis, which demonstrates the different patterns among ovary protein clusters (Fig. 5). The heat map shows that there are variations in protein profile, which are dependent of female weight, and allowed to identify a relationship between protein functions and ovary development. Clearly, this analysis reveals some trends: (i) groups representing early stages of ovarian development are mostly clustered in one branch, including PEF-10, PEF-35, PEF-53, and PEF-84; (ii) groups representing late stages of ovarian development (and group PEF-24) are clustered in another branch, including PEF-189, PEF-270 and FEF-D1; (iii) group FEF-D3 forms a separate branch. The abundance of proteins involved in storage, extracellular matrix, secreted proteins, protease inhibitors, immunity, amino acid metabolism, protein modification, lipocalins, and proteases increased significantly during ovary development, being upregulated at the end of ovary maturation (FEF-D3) (Fig. 5). Conversely, the abundance of proteins related to signalling, protein synthesis, and metabolism decreased over time (Fig. 5). Altogether (Figs 4 and 5), these differences in protein profile during ovary maturation indicate that when ticks detach from the host, ovaries are almost fully mature, with reduced synthesis of new proteins; at this point, the storage of elements important for embryo development becomes more conspicuous.

For more accurate protein quantification, a dimethyl labelling relative quantification was performed in the groups PEF-24, -35, -53, -84, -189 and -270. To analyse changes in ovary protein profile, the data of each group was normalised relative to the PEF-24 group, the smallest individuals who are in the beginning of the blood feeding. This allows to observe variations in protein amount among the developmental stage categories, i.e. during ovary maturation (Fig. 6). Proteins related to cell growth, differentiation, and development (nucleotide...
metabolism, nuclear export, transcription factor, signal transduction for apoptosis, energy metabolism, lipid metabolism, and protein export machinery) are the most abundant throughout ovary development. Moreover, proteins related to intermediate metabolism, lipocalins, and transposable elements are overall much less abundant, being absent in the beginning of ovary development (Fig. 6).

Figure 4. Functional cluserisation of *Rhipicephalus microplus* ovary proteins. Proteomic analysis was performed in partially engorged females, groups PEF-10 (a), -24 (b), -35 (c), -53 (d), -84 (e), -189 (f), and -270 (g); and fully engorged females, groups FEF-D1 (h) and -D3 (i). Pie charts represent the percentage of proteins found in each group respective to normalised spectral counting (NSAF) for each sample. Numbers of ticks composing each group are: PEF-10 (n = 10), PEF-16 (n = 9), PEF-24 (n = 28), PEF-35 (n = 23), PEF-53 (n = 8), PEF-84 (n = 9), PEF-189 (n = 5), PEF-270 (n = 5). FEF groups D1, D2, D3, and D4 were composed by ovaries from 5 ticks.

Figure 5. Protein abundance in *Rhipicephalus microplus* ovaries. Heat map of NSAF data for each protein is expressed as a per cent of total NSAF per group within each category. Z-scores were calculated and used to generate heat maps as described in Material and Methods section. Red colour indicates proteins of high abundance and blue colour indicates proteins of low abundance. Dendrogram on the left shows protein clustering according to functional annotation.
Altogether, the data corroborates previous observations from studies focusing on individual tick ovary proteins. Because of their fundamental role in ovary maturation and embryo development, we selected proteins involved in vitellin processing and energetic metabolism to be analysed and discussed in detail.

**Vitellogenesis during ovary development.** Vt, a lipoglicoprotein, is the main source of nutrients for embryo development. Its degradation depends on a developmentally regulated acidification of the yolk granules, starting immediately after oviposition. Until the time of egg hatching, 40% of Vt content is consumed, implicating a very controlled enzymatic digestion mechanism. Therefore, to improve the understanding of the processes occurring during ovary maturation, critical proteases involved in yolk hydrolysis (BYC, THAP, VTDCE), and a receptor responsible for Vg uptake by the oocytes, *Rhipicephalus microplus* vitellogenin receptor, RmVgR, were selected for dimethyl labelling quantification and further analysis. These vitellin-degrading enzymes and RmVgR are present at high abundance during the rapid engorgement phase of PEF. Additionally, seven Vt polypeptides were identified, and most of them increase in abundance towards the end of the feeding phase. These polypeptides are products of the five different *R. microplus* Vgs, as observed by tBLASTn analyses, confirming the kinetic of Vt processing observed in previous studies. Accordingly, these results are consistent with the current knowledge on tick female physiology, with vitellogenesis being induced by blood meal and mediated by ecdysteroids. In *R. microplus*, adult blood feeding occurs for 7–8 days, and the last 48 hours are characterised by a rapid engorgement, when most of the blood is ingested. After completing the blood meal,
the fully engorged female weight exceeds by more than 100 times that of the respective larva\(^3\). As the rapid blood ingestion occurs, it leads to an acceleration of the metabolism to process the meal, providing nutrients for ovary maturation and, consequently, embryogenesis\(^35,36\).

RmVgR (\(R.\) microplus vitellogenin receptor, GenBank: KY781176)\(^32\), belonging to the low density lipoprotein receptor (LDLR)-like family, is responsible for Vg uptake from haemolymph into the oocytes. RmVgR knock-down in \(R.\) microplus female weighing between 25 and 35 mg results in delayed ovary development and reduced fecundity, suggesting a role in tick reproduction. RmVgR transcripts were specifically identified in the ovaries from PEF and FEF. As transcript levels decrease when females detach from the host (PEF-270), an increase in

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**Figure 8.** Vitellin (Vt) and vitellin degrading cysteine endopeptidase (VTDCE) in \(R.\)hipicephalus microplus ovaries analysed by western blot. Protein extracts from partially (PEF) and fully engorged females (FEF) ovaries were probed with rabbit antibodies (a) anti-Vt and (b) anti-VTDCE. PEF groups: -10, -16, -24, -35, -53, -84, -189, -270; FEF groups: -D1, -D2, -D3 and -D4. Gaps were created in the images to delineate the stages of ovary maturation, but both parts are from the same gel. This image is representative of two (for VTDCE) and four (for Vt) independent western blot. Numbers of ticks composing each group are: PEF-10 (n = 10), PEF-16 (n = 9), PEF-24 (n = 28), PEF-35 (n = 23), PEF-53 (n = 8), PEF-84 (n = 9), PEF-189 (n = 5), PEF-270 (n = 5). FEF groups D1, D2, D3, and D4 were composed by ovaries from 5 ticks. This image is representative of two (for VTDCE) and four (for Vt) independent western blot.

**Table 1.** Similarity match of vitellin polypeptides identified in \(R.\)hipicephalus microplus ovary proteome, compared to \(R.\)hipicephalus microplus vitellogenins deposited in GenBank. *Best match considering e-value 0.0. **Transcribed RNA sequence. ***mRNA sequence, partial CDS.
protein abundance is observed\(^2\). Relative quantification using dimethyl labelling corroborates RmVgR identification in all PEF groups, with a larger amount at the end of the rapid engorgement phase (PEF-189 and PEF-270) (Fig. 7a). However, receptor presence in FEF could not be detected by NSAF, probably due to a high abundance of storage proteins (e.g. Vt)\(^2\).

**Boophilus** yolk pro-cathepsin (BYC, GenBank: AY966003) is an acidic aspartic endopeptidase activated by autoproteolysis, that hydrolyses Vt and also haemoglobin\(^2\). This enzyme is synthesised in fat body and midgut and then secreted into haemolymph to be incorporated by oocytes, being allocated in the periphery of laid eggs\(^2\). BYC has a high specificity and low activity upon Vt, suggesting a role in continuously providing amino acids and energy for embryo development until larvae find a host\(^2\). Through dimethyl labelling relative quantification, BYC was identified in ovary from all groups between PEF-53 and PEF-270 (Fig. 7a). Also, as expected for an enzyme with a role in yolk degradation, BYC is present at higher levels in ovaries at late stages of oogenesis (PEF-270 group).

Another aspartic proteinase of which the presence in *R. microplus* ovary was confirmed is tick haeme-binding aspartic proteinase (THAP, GenBank: AF288685). This enzyme is also involved in Vt and haemoglobin hydrolysis. It is distributed in the oocyte cytoplasm, surrounding the yolk granules, and its activity is regulated by haeme\(^2\). THAP is synthesised in ovaries and also in fat body and midgut, being transported via haemolymph to the ovary\(^2\). Considering that THAP is observed as two polypeptides in PEF ovaries and in laid eggs, that the most part of the polypeptides is converted to active enzyme after oviposition, it is conceivable that enzyme activation occurs during acidification of the yolk granules\(^2\). Yolk degradation is a long process, since *R. microplus* vitelligenesis takes 21 days, and a way to control THAP activity is through the haeme released by Vt proteolysis. Haeme competes with Vt to bind THAP, a negative feedback that slows down Vt degradation and prevents the oxidative damage caused by excess of haeme\(^2\). THAP is present in ovaries from PEF-35 to PEF-270 groups, and its abundance increases once females reach 84 mg (PEF-84 group); THAP levels keep rising until females reach 270 mg, when the feeding period is almost finished (PEF-270 group) (Fig. 7a).

**Vitellin-degrading cysteine endopeptidase** (VTDCE, GenBank: JQ080269) is a cathepsin L-like that hydrolyses Vt and haemoglobin\(^1\), and is found tightly bound to Vt in *R. microplus* eggs\(^9\). VTDCE is localised in midgut basal lamina and basophilic cells indicating that, just like BYC and THAP, it is synthesised in the midgut and then transported through the haemolymph to the ovaries, where it is internalised into pedicel cells and oocyte cytosol. A VTDCE inhibitor is present in female haemolymph which acts in a dose-dependent manner and is believed to control Vt hydrolysis before it is internalised into oocytes\(^9\). In this work, VTDCE was observed at low abundance in ovaries during the earliest phase of blood feeding, and at a higher abundance during rapid engorgement (PEF-189 and PEF-270), similarly to BYC and THAP (Fig. 7a). Also, this enzyme was detected in FEF-D2, -D3 and -D4 groups, when ovary protein extracts were probed with anti-VTDCE polyclonal antibodies (Fig. 8b).

A second cathepsin L-like that hydrolyses Vt\(^1\) and haemoglobin\(^1\), *Boophilus microplus* cathepsin-L like (BmCL1, GenBank: AF227957)\(^1\), was detected in the present analysis. This enzyme, also known as RmLCE (*Rhipicephalus microplus* larval cysteine endopeptidase), is synthesised in *R. microplus* PEF\(^2\) and FEF\(^3\) midgut, and its transcription was observed in total extract of PEF, FEF and larvae\(^2\). BmCL1 is localised in secretory vesicles of *R. microplus* PEF midgut, and its presence was identified also in PEF and larvae (5 to 10 days) total protein extracts\(^2\). This is the first report of BmCL1 presence in *R. microplus* ovary, being identified during oogenesis in PEF and FEF. The highest abundance of BmCL1 was observed during rapid engorgement (PEF-189 and PEF-270) and in the first day after detachment (FEF-D1) (Fig. S1), a profile similar to what is observed for other enzymes.

### Table 2. Similarity match of vitellin polypeptides identified in *Rhipicephalus microplus* ovary proteome, compared to tick vitellogenins deposited in GenBank. *Best match considering e-value 0.0. **mRNA sequence, complete CDS. ***mRNA sequence.

| Vt ID | Best match against ticks Vgs* | Accession number |
|-------|------------------------------|------------------|
| Vt_1  | *Dermacentor variabilis* Vitellogenin | AY885250.2** |
|       | *Amblyomma hebraeum* Vitellogenin I | JX846593.1*** |
|       | *Isodes scapularis* Vitellogenin, putative | XM_002415179.1*** |
| Vt_2  | *Haemaphysalis longicornis* Vitellogenin-B | AB359900.1** |
|       | *Isodes scapularis* Apolipophorin, putative | XM_002401724.1*** |
| Vt_3  | *Haemaphysalis longicornis* Vitellogenin-B | AB359900.1** |
|       | *Isodes scapularis* Apolipophorin, putative | XM_002401724.1*** |
| Vt_4  | *Haemaphysalis longicornis* Vitellogenin-B | AB359900.1** |
|       | *Isodes scapularis* Apolipophorin, putative | XM_002401724.1*** |
| Vt_5  | *Dermacentor variabilis* Vitellogenin-2 | EU204907.2** |
|       | *Amblyomma hebraeum* Vitellogenin-2 | JX846594.1*** |
|       | *Ornithodoros moubata* Vitellogenin | AB440159.2** |
|       | *Isodes scapularis* Vitellogenin, putative | XM_002403922.1*** |
| Vt_6  | *Haemaphysalis longicornis* Vitellogenin-1 | AB359899.1** |
| Vt_7  | *Haemaphysalis longicornis* Vitellogenin-B | AB359900.1** |
|       | *Isodes scapularis* Conserved hypothetical protein | XM_002401721.1*** |
In this proteomic study, we identified a subset of proteins that have regulated abundance in ovary during ovary maturation, and their function and physiological significance are discussed. In summary, a variety of tick proteins studied as targets for the development of control methods, as well as new putative targets, were identified at different developmental stages.

**Figure 9.** Quantification of proteins in selected metabolic pathways related to vitellogenesis. Dimethyl labelling quantification was performed in ovary extracts of partially engorged female (PEF) groups -24, -35, -53, -84, 189, -270. The ratio is the intensity detected for each peptide, relative to the internal standard (a protein pool of all ovary groups used in this study). The average ratio of a protein is calculated as the mean ratio of its peptides. The average ratio of each metabolic pathway presented in this graph was calculated from the average quantification (with standard deviation) of enzymes involved in each pathway (Supplementary Fig. S1). Numbers of ticks composing each group are: PEF-10 (n = 10), PEF-16 (n = 9), PEF-24 (n = 28), PEF-35 (n = 23), PEF-53 (n = 8), PEF-84 (n = 9), PEF-189 (n = 5), PEF-270 (n = 5).

**Metabolic changes during tick ovary development.** Previous studies have identified a number of proteins that are involved in specific processes within oogenesis and embryogenesis in different arthropods\(^45-47\). Present results show that during *R. microplus* ovary development, vitellogenic proteins are upregulated from late-vitellogenic follicles to mature oocytes (i.e. from PEF-53 to more mature ovary). Meanwhile, throughout the vitellogenic process, RmVgR abundance is upregulated from early-vitellogenic oocytes (i.e., before Vt internalisation, more immature than PEF-53) to late-vitellogenic oocytes (Fig. 7a). Accordingly, different Vt peptides are observed in association with changes in abundance during the same period (Fig. 7b). These results indicate that ovary development and the consequent oocyte maturation processes occur in this tick similarly to other arthropods\(^48-50\). At the same time, a large variation was observed in the levels of proteins involved in carbohydrate metabolism during ovary maturation, especially enzymes involved in glycogen metabolism (e.g. glycogenin glucosyltransferase) (Fig. S2a). This is consistent with the dynamic of the final stages of oocyte maturation, which requires different compounds such as fatty acids, amino acids, electrolytes, purines and pyrimidines, and other metabolites\(^49\). Regarding metabolites, glucose, glycogen, lipids, and pyruvate were also observed by many authors in different arthropods\(^51-53\). Furthermore, carbohydrates facilitate oocyte maturation and developmental competence after fertilisation in mammals\(^48,54\), and are important for embryonic development in arthropods\(^55\), suggesting a highly conserved link between carbohydrate metabolism and oocyte development.

Large differences in protein levels were identified in early-vitellogenic ovaries, indicating that maternal sources sustain egg formation, which accumulates Vt (Fig. 2). The highest levels of the studied peptidases (BYC, VTDCE, THAP, and BmCL1) are observed at the end of the vitellogenic process, simultaneously to oocyte growth. In general, proteins at low concentrations during early vitellogenesis were linked to protein processing, sugar metabolism, amino acid metabolism, mitochondrial proteins, alcohol metabolism, and ATP synthesis (Figs 9 and S2). These endopeptidases accumulate in ovaries during oogenesis, and have a physiological importance at a later stage of embryo development, when they participate in yolk degradation, supplying amino acids for the synthesis of new proteins, and as a source of energy. A more gradual change was observed for the main yolk reserve protein, Vt (substrate of BYC, VTDCE, THAP, and BmCL1) (Fig. 7b), showing that during early oogenesis, Vt degradation provides the major substrate for embryo development.

**Conclusions**

This work describes 3,756 proteins involved on arthropod ovary development, and evaluates the modulation of several of them. Also, the profile of proteins involved in Vt degradation was corroborated by western blot analysis. In this proteomic study, we identified a subset of proteins that have regulated abundance in ovary during ovary maturation, and their function and physiological significance are discussed. In summary, a variety of tick proteins studied as targets for the development of control methods, as well as new putative targets, were identified at different developmental stages.
Materials and Methods

Ethics statement. Cattle used in this work were housed at Faculdade de Veterinária, Universidade Federal do Rio Grande do Sul (UFRGS), Porto Alegre, Brazil. This research was conducted according to the ethical and methodological aspects prescribed by the International and National Directives and Norms by the Animal Experimentation Ethics Committee of UFRGS. The protocols were approved by Comissão de Ética no Uso de Animais - CEUA–Universidade Federal do Rio Grande do Sul.

Ticks. Rhipicephalus microplus of Porto Alegre strain (Brazil) was reared on Hereford calves (Bos taurus taurus), which were brought from a naturally tick-free area and maintained in insulated pens. Calves were infested with 15-day-old tick larvae and after 22 days, partially and fully engorged adult female ticks were manually collected for ovary dissection.

Ovary extracts and protein digestion. Partially engorged females (PEF) were manually collected from two different calves, grouped by weight in order to represent different periods of development, and then the ovaries were dissected. The weight average, SD and number of ticks in each group was as follows: 10 ± 1.73 mg (PEF-10, n = 10), 16 ± 1.22 mg (PEF-16, n = 9), 24 ± 1.90 mg (PEF-24, n = 28), 35 ± 2.84 mg (PEF-35, n = 23), 53 ± 2.49 mg (PEF-53, n = 8), 84 ± 8.36 mg (PEF-84, n = 9), 189 ± 17.17 mg (PEF-189, n = 5) and 270 ± 16.71 mg (PEF-270, n = 5). Fully engorged females (FEF) were placed in four groups according to days post-detachment. All of them were maintained at 28 °C and 85% relative humidity. Ovaries from FEF one, two, three and four days after detachment (D1, D2, D3, and D4, n = 5 for each group) were obtained by dissection. The variation in number of ticks per group was consequence of ovary size, and each sample was normalised by total protein amount. Biological variability within the samples is guaranteed by the collection of ticks made in two different calves and the number of ticks composing each group. These are the same groups used in other experiments carried out concomitantly25, although data was analysed independently.

Proteins from samples of each experimental group were extracted using TRIzol® reagent (Thermo Fisher Scientific) following the manufacturer’s recommendations. Protein concentration was measured using BCA Protein Assay Reagent Kit, following the manufacturer’s recommendations. Ticks used for proteomic analysis were from groups PEF-10, -24, -35, -53, -84 and -270, and FEF-D1 and -D3.

Dimethyl labelling. Dimethyl labelling was performed according to Boersema et al.36. Samples (30 µg) dissolved in 2 M urea/100 mM Tris, pH 8.5. Proteins were reduced with 5 mM tris(2-carboxyethyl)phosphine hydrochloride (TCEP, Sigma-Aldrich) and alkylated with 25 mM iodoacetamide (Sigma-Aldrich). Proteins were hydrolysed without acidification. Columns and analytical columns were connected using a zero-dead volume union (Upchurch Scientific).

Peptide mixtures were analysed by LC-MS using quaternary HP 1100 series HPLC (high performance liquid chromatography) pump (Agilent Technology) connected to an LTQ (Linear Trap Quadrupole) XL or an LTQ Orbitrap Velos mass spectrometer (Thermo Scientific). Electrospray was performed directly from the tip of the analytical column. Mobile phases were as follows: solution A, 5% acetonitrile and 0.1% formic acid; solution B, 80% acetonitrile and 0.1% formic acid; solution C, 500 mM ammonium acetate, 5% acetonitrile and 0.1% formic acid. Flow rate was approximately 300 nL/min.
MuDPIT experiments were performed using a 45-minute transfer step followed by ten 2-hour cycles of salt pulse (solution C injection) and separation steps. Transfer step consisted of a linear gradient from 0 to 60% B over 25 minutes, followed by an increase to 100% B in 5 minutes. Column was re-equilibrated with 100% A for 10 minutes. Salt pulse consisted of a 4-minute solution C injection followed by 6 minutes of 100% A. Peptide separation gradient was from 0 to 10% B in 5 minutes, increase to 20% B over 35 minutes, increase to 50% B over 40 minutes, and increase to 100% B in 15 minutes. Column was flushed at 100% B for 5 minutes and re-equilibrated in 100% A for 20 minutes. At the end of the ten cycles, mobile phase during the salt pulse consisted of 10, 20, 30, 40, 50, 60, 70, 85, or 100% C, or 90% C/10% B. Due to a slight increase in peptide hydrophobicity, for dimethyl labelling experiments, the cycle with 40% C was omitted, and an extra cycle was performed in the end with 90% C/10% B.

The LTQ XL was operated in a data-dependent scan mode, with ESI voltage of 3 kV and inlet capillary temperature of 275°C. Full MS1 scans and CID MS2 scans of the 5 most abundant ions in each cycle were collected on the LTQ ion trap. AGC target was 3e6 and 1e4 for MS1 and MS2, respectively, and the mass range was 300 to 1200 m/z. Maximum injection times were 50 ms for MS1 and 100 ms for MS2 scans. Dynamic exclusion was enabled with repeat count of 1, repeat duration of 30 s, exclusion list size of 50 and exclusion duration of 30 s.

The LTQ Orbitrap Velos was also operated in a data-dependent mode, ESI voltage of 3.5 kV and inlet capillary temperature of 275°C. Full MS1 scans were collected in the Orbitrap, with mass range of 300 to 1200 m/z at 60 K resolution and an AGC target of 1e6. The 20 most abundant ions per MS1 scan were selected for CID MS2 in the LTQ Velos ion trap, with an AGC target of 1e4 and threshold intensity of 500. Maximum fill times were 250 ms and 100 ms for MS1 and MS2 scans, respectively, and dynamic exclusion was used with repeat count of 1, repeat duration of 150 s, exclusion list size of 500 and exclusion duration of 120 s.

**LC-MS/MS data analysis.** Label-free quantitative analysis was performed by comparing single MuDPIT technical replicates of each tick ovary groups (PEF-10, -24, -35, -84, and -270, and FEF-D1 and -D3). However, as the incomplete elution of peptides from ion exchange column allows peptide identification in more than one fraction, these fractions are analysed together, generating a unique result with redundant peptide data. The semi-quantitative method based on NSAF (normalised spectral abundance factor) was used. NSAF for a given protein is the number of spectral counts (SpC) identified for that protein, divided by the protein’s length (L), divided by the sum of SpC/L of all protein in the experiment.

Dimethyl labelling quantitative analysis was performed in ovary groups labelled as described above (section Dimethyl labelling). One MuDPIT technical replicate for each sample pair and internal standard (standard:PEF-24:PEF-35, standard:PEF-53:PEF-84 and standard:PEF-189:PEF-270) was used. Due to MuDPIT characteristics, one peptide can be identified and quantified in more than one fraction in the same run. This redundancy of measurements supports data reliability.

Protein and peptide identification was done with Integrated Proteomics Pipeline–IP2 (Integrated Proteomics Applications). Tandem mass spectra were extracted from raw files using RawExtract 1.9.9.2 and searched with ProLuCID against a local *R. microplus* protein database (Rm-INCT-EM) containing 22,010 sequences previously produced by our research group using Illumina Sequencing technology (BioProject ID PRJNA232001 at Transcriptome Shotgun Assembly (TSA) database–GenBank). The search space included all fully-tryptic and half-tryptic peptide candidates. Carbamidomethylation on cysteine residues (+57.02146) was used as a static modification. For dimethyl labelling, dimethylation (+28.0313) of N-terminus and lysines was considered as static modification, and the mass difference between intermediate and light labels (+4.0251), or between heavy and light labels (+8.0444), were considered as metabolic labelling. Data was searched with 50 ppm precursor ion tolerance and 600 ppm fragment ion tolerance. Identified proteins were filtered using DTASelect. Filtering required a minimum of 2 peptides per protein, at least one tryptic terminus for each peptide, and less than 1% FDR. Normalised spectral abundance factor (NSAF) was calculated according to Zybailov et al. Comparisons between runs were performed by identificationCOMPARE, part of the IP2 pipeline. Quantitative analysis was performed by Census integrated into IP2 pipeline. Protein ratios (medium/light and heavy/light) were calculated by quantitativeCOMPARE, using peptide ratios and including redundant proteins.

**Protein functional annotation.** The functional annotation and clustering of protein sequences were performed using Visual Basic programs developed by Dr. José Marcos Ribeiro. BLASTP tool was used to search proteins sequences against several databases. The final result presented here was manually curated and, in many cases, manually annotated. This catalogue was plotted on a hyperlinked Microsoft Excel spreadsheet (see Supplementary Table S1).

The abundance profile of dimethyl-labelled proteins was analysed by normalizing each PEF group relative to PEF-24 group. Proteins with *p* value and ratio variance ≥0.06 were excluded from the analysis (see Supplementary Table S3).

**Relative abundance and graphical visualisation.** Proteomic profiles were compared across samples as functional clusters of proteins. To determine the relative abundance of proteins, NSAF was used in a label-free relative quantification approach. NSAF as an index for relative protein abundance was input in Microsoft Excel as percentage of the total NSAF for respective samples, and visualised on pie charts according to protein clusters. To visualise relative protein patterns on a heat map, NSAF values were normalised using Z-score. Normalised NSAF values were used to generate heat maps using the heatmap2 function from the gplots library in R.

**SDS-PAGE and Western Blot.** Tick ovary protein extracts were analysed by 10% SDS-PAGE (sodium dodecyl sulphate–polyacrylamide gel electrophoresis). The gel was loaded with 50 µg of protein per sample lane (PEF-10, -16, -24, -35, -84, -189 and -270; and FEF-D1, -D2, -D3 and -D4) and stained with coomassie brilliant blue for protein visualisation.
The presence of vitellin and VTDCE in ovary extracts was evaluated by western blot. Fifty micrograms of each tissue extract were resolved by SDS-PAGE and eletroblotted on nitrocellulose membranes. Membranes were blocked with 5% non-fat dry milk in PBS (blocking solution), for 1 hour at room temperature. Membranes were incubated with anti-vitellin\(^{13}\) or anti-VTDCE\(^{14}\) rabbit sera (1:100 and 1:200), respectively, in blocking solution for 16 h at 4 °C. After three washes with blocking solution, membranes were incubated with anti-rabbit antibodies conjugated with alkaline phosphatase (diluted 1:5,000 in blocking solution) for 1 hour at room temperature. After washes with PBS and alkaline phosphatase buffer (Tris-HCl 100 mM, NaCl 100 mM and MgCl\(_2\) 5 mM), membranes were incubated with NBT (nitro blue tetrazolium, Thermo Scientific; 0.3 mg/mL) and BCIP (5-bromo-4-chloro-3-indolyl phosphate, Fermentas; 0.15 mg/mL) in alkaline phosphatase buffer.

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Conceived and designed the experiments: M.A.X., L.T., A.F.M.P., I.S.V.J., A.S. and C.T. Performed the experiments: M.A.X., L.T., A.F.M.P., J.K.D., and A.S. Contributed reagents/materials/analysis tools: J.R.Y., I.S.V.J., A.S., and C.T. Drafting the article: M.A.X., L.T., A.F.M.P., J.K.D., J.R.Y., A.M., C.L., I.S.V.J., A.S. and C.T. Critical revision of the article: M.A.X., L.T., A.F.M.P., J.K.D., J.R.Y., A.M., C.L., I.S.V.J., A.S. and C.T.

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