

In-silico definition of the *Drosophila melanogaster* matrisome



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Abstract

The extracellular matrix (ECM) is an assembly of hundreds of proteins that structurally supports the cells it surrounds and biochemically regulates their functions. *Drosophila melanogaster* has emerged as a powerful model organism to study fundamental mechanisms underlying ECM protein secretion, ECM assembly, and ECM roles in pathophysiological processes. However, as of today, we do not possess a well-defined list of the components forming the ECM of this organism. We previously reported the development of computational pipelines to define the matrisome - the ensemble of genes encoding ECM and ECM-associated proteins - of humans, mice, zebrafish and *C. elegans*. Using a similar approach, we report here that our pipeline has identified 641 genes constituting the *Drosophila* matrisome. We further classify these genes into different structural and functional categories, including an expanded way to classify genes encoding proteins forming apical ECMs. We illustrate how having a comprehensive list of *Drosophila* matrisome proteins can be used to annotate large proteomic datasets and identify unsuspected roles for the ECM in pathophysiological processes. Last, to aid the dissemination and usage of the proposed definition and categorization of the *Drosophila* matrisome.org), The FlyBase (https://flybase.org/), and GLAD (https://www.flyrnai.org/tools/glad/web/).

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Introduction

The extracellular matrix (ECM) is an assembly of hundreds of proteins that structurally supports and biochemically regulates the cells it surrounds [1,2]. The ECM organizes the tissues of all metazoans [3]. It plays a role in a number of biological processes, from development and homeostasis [4–6] to pathological processes including fibrosis and cancer [4,7,8]. With a growing interest from the scientific community in the ECM and the emergence of high-throughput technologies generating large datasets came the realization that a robust definition of the proteins contributing to the formation of the ECM was needed. We thus defined the matrisome of human and mouse [9–11]. This was achieved by developing a computational approach based on protein sequence analysis using key structural features of ECM proteins, including the presence of a signal peptide and specific protein domains found predominantly in ECM and ECMassociated proteins [9,12]. We further proposed to classify the matrisome into the core matrisome, which is the compendium of genes encoding proteins forming the structure of the ECM (collagens, glycoproteins, and proteoglycans), and the matrisomeassociated ensemble comprising genes encoding accessory proteins and proteins involved in the remodeling of the ECM [9,10,13]. The adoption of these definitions by the scientific community has allowed the identification of ECM proteins previously unsuspected to play roles in physiological or pathological processes [14-16] and of ECM signatures in -

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Fig. 1. Bioinformatic workflow to define the *in-silico* matrisome of *Drosophila melanogaster*. The databases FlyBase, DIOPT, and Ensembl were interrogated with the full list of human and mouse matrisome and matrisome-associated gene symbols. Selected InterPro domains, including domains characteristic of collagens, proteoglycans, ECM-affiliated proteins, and cuticle-binding proteins (see Supplementary Table 4) were used to identify ECM-domain-containing proteins in the reference proteome. The Gene Ontology annotations related to the ECM were then used to identify previously-annotated ECM components. Finally, selected published literature using proteomic and/or bioinformatic methods, as well as reviews on the subject, were searched to identify ECM proteins not identified by the orthology-based or protein-sequencing methods. These data were combined and manually curated to generate the first complete *Drosophila* matrisome. *Hs, Homo sapiens; Mm, Mus musculus; Dm Drosophila melanogaster.*

omic datasets predictive, for example, of cancer patient outcome [7,17–19]. This prompted us and others to further define the matrisome of several model organisms: zebrafish [20], *Caenorhabditis elegans* [21], and planarians [22].

In recent years, there has been a surge of interest in using the genetic tractability of Drosophila melanogaster to identify fundamental mechanisms underlying ECM assembly, structure, and function, since several ECM proteins and processes contributing to the formation and assembly of the ECM are conserved between Drosophila and other organisms [23-25]. This surge is most evident in studies of basement membrane (BM) biology [26,27]. BM is an ancient and highly conserved ECM that lines the basal surface of epithelial and endothelial tissues and surrounds muscles, adipose tissue, and nerves [26,28,29]. Studies using Drosophila have made particularly strong contributions to our understanding of BM secretion and assembly [30-45], and the role BMs play in shaping

tissues during development [35,42,46–53]. They have also shown how BMs heal after injury [54,55] and how they regulate the immune response [56–60]. More recently, work in *Drosophila* has introduced a new role for BM proteins in intercellular adhesion [31].

Although the core BM proteins (type IV collagens, laminins, heparin sulfate proteoglycans, and nidogens) are well known, proteomic studies have revealed that BMs can harbor numerous accessory proteins that vary by tissue [14,61,62]. A comprehensive list of these proteins will provide an important tool for *Drosophila* researchers as they continue to probe the diverse roles BMs play in animal development and physiology.

Drosophila also have ECMs that are unique to arthropods and are therefore not found in any other organism for which the matrisome has been defined. These include: the chitin-based cuticle that forms the animal's exoskeleton and lines the lumens of the foregut and hindgut [63–65]; non-cuticular, chitinbased ECMs that line the lumens of the trachea, salivary glands, and midgut [64,66]; the eggshell that protects the developing embryo [67,68]; and the salivary glue that is produced by the larva to affix the pupa to a surface [69]. Defining the list of proteins that comprise these ECMs will provide a reference dataset for the arthropod clade and aid with the annotation of large proteomic datasets, including the developmental proteome of *Drosophila* [70]. Moreover, because insects can be both disease vectors and agricultural pests, these data could provide an important source of information to combat these threats to human welfare.

Here, we define the in-silico matrisome for Drosophila melanogaster. To this end, we developed a computational pipeline that combines orthology comparison, protein sequence analysis, interrogation of experimental proteomic data, and literature search (Fig. 1) and identified 641 genes that we propose to comprise the Drosophila matrisome. We further classified these 641 genes into different structural and functional categories based on the model we have proposed for the matrisomes of other organisms [9,20,21]. We then describe the deployment of our list and terminology in the Matrisome Project website (http://matrisome.org) and in two databases, FlyBase [71,72] and GLAD [73], broadly used by the Drosophila community. Last, we illustrate how this new resource can be used to annotate -omic datasets.

In-silico definition of the *Drosophila melanogaster* matrisome

Identification of Drosophila orthologs of human and mouse matrisome genes

We first set out to identify the Drosophila orthologs of human and mouse matrisome genes. The sequence alignment tools built into Flybase (FB2017 03, released June 2017) [71], the Drosophila RNAi Screening Center's Integrative Ortholog Prediction Tool (DIOPT, Version 6.0.2, released June 2017) [73,74], and Ensembl (Ensembl 89, released May 2017) [75] were interrogated with the full list of human and mouse core matrisome and matrisome-associated gene symbols from each of the six categories of ECM components defined previously (Fig. 1) [11]. Although these sequencealignment meta-algorithms return a confidence score based on the number of algorithms returning a hit, we did not eliminate genes with low-confidence scores at this stage to maximize our potential to identify relevant genes.

The genes retrieved by each of the three databases (Supplementary Table 1A and 1B) were compiled to obtain a list of all predicted

Drosophila orthologs of human and/or mouse matrisome genes (Supplementary Table 1C). The results of this approach led to the identification of 834 putative *Drosophila* matrisome orthologs. Of these genes, 114 were orthologous to a human gene but not a mouse gene, whereas 51 were orthologous to a mouse gene but not a human gene. There were 296 human genes with no *Drosophila* ortholog (Supplementary Table 2A) and 340 mouse genes with no *Drosophila* ortholog (Supplementary Table 2B).

Protein-domain-based approach to identify additional Drosophila matrisome proteins

Since it is well known that flies also have a large number of ECM proteins that do not have mammalian orthologs (see Introduction), we next used the UniProt Drosophila reference proteome (downloaded August 10, 2017) [76] to further expand our search for matrisome components (Supplementary Table 3A). Taking advantage of the conserved domain-based nature of ECM proteins [12], we selected InterPro domains [77] which were previously used to identify human and mouse matrisome proteins [9,10], including domains characteristic of collagens, proteoglycans, and ECM-affiliated proteins, to search for ECM-domain-containing proteins in the Drosophila proteome (Supplementary Table 4A). We also included in the search three domains characteristic of proteins involved in the production and maintenance of chitin-based ECMs: insect cuticle protein (IPR000618), chitin-binding domain (IPR002557), and chitin-binding type R&R consensus (IPR031311) [78]. Although three ECM domains were initially used to search the UniProt Drosophila reference proteome, the domain chitin-binding type R&R consensus (IPR031311), a conserved motif of 35-36 amino acids identified by Rebers and Riddiford (R&R) [79,80], was found to be redundant with the domain insect cuticle protein (IPR000618) for the identification of the 213 Drosophila proteins. (Supplementary Table 4B). To complete the list of proteins composing the Drosophila cuticle we further interrogated CuticleDB, a database of structural components of arthropods identified experimentally or through protein sequence analysis [81]. This allowed us to retrieve an additional 7 genes (CG13670, CG7548, CG8541, CG8543, Cpr65Ax1, Edg91, Lcp6) that were added to the class of cuticular proteins.

Using this method, we identified 353 *Drosophila* proteins with ECM domains: 140 using domains previously used to identify mammalian matrisome proteins and an additional 213 using domains characteristic of *Drosophila* proteins (Supplementary Table 4B). We compared the list of proteins identified with human matrisome domains to the proteins identified via gene orthology and found that

		/	Apical Matrix Classes and Sub-classes	# of Genes
		/	Cuticle	56
		/	Cuticle; Chitin-binding-domain-containing Proteins	2
		/	Cuticle; Chitin-binding-domain-containing Proteins; Chitinase	5
Matrisome Category	# of Genes	- /	Cuticle; Chitinase	2
		- /	Chitinase	2
Collagens	4	- /	Cuticle; Chitinase-like	4
ECM Glycoproteins	27	_ /	Chitinase-like	2
Proteoglycans	3		Cuticle; Tweedle	26
ECM-affiliated	106	_ /	Cuticle; R&R Chitin-binding-domain-containing Proteins	79
		- /	R&R Chitin-binding-domain-containing Proteins	2
ECM Regulators	98	-/	Chitin-binding-domain-containing Proteins	85
Secreted Factors	75		Chitin-binding-domain-containing Proteins; Chitin Deacetylase	5
Apical Matrix	328		Chitin Deacetylase	1
TOTAL	641		Chitin-binding-domain-containing Proteins; Mucin	9
	0.2	- \	Glue	11
			Zona Pellucida-domain-containing Proteins	11
			Eggshell; Vitelline Membrane	14
		\backslash	Eggshell; Chorion	12

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Fig. 2. The *Drosophila* matrisome. (A) The *Drosophila* matrisome comprises 641 genes. These genes are then divided into either categories which we have previously defined, or the newly proposed apical matrix category. (B) The genes that encode proteins that make up the apical matrix of *Drosophila* were further divided into classes and sub-classes.

49 of the 140 proteins discovered by domains characteristic of ECM proteins (35%) were not previously identified using the orthology approach (Supplementary Table 4C).

Gene-Ontology-based approach to identify additional Drosophila matrisome proteins

The Drosophila proteome retrieved from UniProt is also annotated with Gene Ontology (GO; http:// geneontology.org) - Cellular Component terms describing the intra- and extracellular localization of proteins [82,83]. The Gene Ontology terms extracellular matrix (GO:0031012), extracellular region (GO:0005576), extracellular space (GO:0005615), basement membrane (GO:0005604), and proteinaceous extracellular matrix (GO:0005578) were used to identify ECM components. The term proteinaceous extracellular matrix was found to be redundant with the term extracellular matrix, but the other four terms made significant contributions to the breadth of the search, which retrieved 1308 proteins from the Drosophila proteome (Supplemental Table 3B).

Manual curation of potential Drosophila matrisome genes

TOTAL

328

The three computational approaches described above identified 1585 genes encoding potential Drosophila matrisome proteins. As we reported for other organisms, a purely computational approach is not sufficient to identify ECM genes [9,10,21]. For example, examination of the list revealed the presence of proteins that share structural features with ECM proteins such as transmembrane receptors, anti-microbial peptides or accessory gland proteins, but that are clearly not ECM components. We thus undertook a knowledge-based approach to manually curate the list and either eliminate non-ECM genes or include ECM genes that had been missed by our computational approach. As GO annotations have been found previously to lack specificity to define ECM components [10], analysis of all proteins identified by GO annotation was performed using the Phobius signal peptide predictor [84]. Proteins that lack a signal peptide and did not exhibit other significant ECM characteristics were excluded along with proteins predicted to be

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cytoplasmic, proteins with multiple transmembrane domains, and proteins with contradictory GO annotation such as cytosolic (GO:0005829) or lysosomal (GO:0005764) localization. This step alone excluded 303 genes. We further consulted extensively the literature on the Drosophila ECM, and made direct queries of FlyBase [71] and UniProt [76] to examine closely the protein sequences and orthology relationships to mammalian ECM proteins of genes and decide whether there was sufficient evidence to classify these genes as part of the matrisome. This final curation step allowed us to identify some genes encoding ECM proteins that were missed by our computational screen and to eliminate some genes for which experimental evidence does not support their classification as matrisome components (see below).

The Drosophila matrisome is composed of 641 genes

Based on the combined result of these analyses, we propose that the Drosophila matrisome is composed of 641 genes (Supplemental Table 5). Interestingly, this number represents 4% of the 15,500 protein-coding genes in the Drosophila genome, which is comparable to the percentage of the genome encoding ECM proteins in humans, mice, zebrafish, and C. elegans [9,20,21] and is likely to be similar to the proportion of matrisome genes in the planarian genome [22,85]. Below, we describe how these 641 genes have been classified into matrisome categories based on their structure, localization, and/or function.

Classification of Drosophila genes orthologous or homologous to mammalian matrisome genes

Genes with orthology or homology to human genes were categorized based on the previously proposed mammalian matrisome divisions (core matrisome or matrisome-associated) and categories (collagens, glycoproteins and proteoglycans for the core matrisome, and ECM-affiliated proteins, ECM regulators and secreted factors for matrisomeassociated components) [9].

The Drosophila matrisome contains 34 core matrisome genes: 4 collagens, 27 glycoproteins, and 3 proteoglycans, the majority of which are orthologous to mammalian core matrisome genes (Supplementary Table 5A and 5B and Fig. 2A). Only 1 collagen (pericardin) [86,87] and 6 glycoproteins (artichoke [88], anachronism [89], Defense protein I (2)34Fc, glutactin [90], tiggrin [91], and tenectin [92,93]) were not homologous or orthologous to mammalian core matrisome genes. Since the expansion and complexification of the ECM emerged with the appearance of deuterostomes, a large number of ECM genes or families of genes were not found in Drosophila, including fibronectin, matricellular proteins such as thrombospondins and tenascins, and non-basement membrane collagens [23,94-96].

In addition to core matrisome genes, we predict that the Drosophila genome encodes 279 matrisomeassociated genes, including 219 that are orthologous or homologous to mammalian genes (Supplementary Tables 5A and B and Fig. 2A).

We previously defined ECM-affiliated proteins as proteins either somewhat structurally related to core ECM proteins or that have been found experimentally to be associated with the ECM in detergentinsoluble fractions of tissue lysates by proteomics [9,11]. Our computational approach predicts that 106 Drosophila genes encode ECM-affiliated proteins. Among these are galectins, C-type lectins (structurally characterized by three InterPro domains, IPR001304, IPR016186, and IPR016187), mucins, and semaphorins, some of which are orthologous or homologous to mammalian genes (Supplementary Table 5A). In addition, we classified under this category 6 collagen-triple-helix repeat-containing proteins and 9 fibrinogen-domain-containing proteins with no clear mammalian orthologs.

The ECM regulators category groups enzymes participating in the synthesis or remodeling of the ECM together with the regulators of these enzymes (including inhibitors). We identified 98 ECM regulators (Supplementary Table 5), including matrix metalloproteinases [25], cathepsins, ADAMs, and two orthologs of the recently identified serine/ threonine kinase family Fam20 [97]. Our study also identified a total of 24 prolyl-4-hydroxylases (P4Hs). Prolyl-4-hydroxylases catalyze the formation of hydroxyprolines [24,98-100]. The most wellrecognized role of this post-translational modification is to stabilize collagen triple-helical structures. Interestingly, and as previously noted [24], the human genome encodes 44 collagen genes and 3 P4Hs, whereas the Drosophila genome encodes only 4 collagen genes, 6 collagen-triple-helix repeatcontaining proteins and yet 24 P4Hs. Both previous work [100] and interrogation of The National Human Genome Research Institute **mod**el organism **ENC**yclopedia Of DNA Element (modENCODE) database [101] indicate that the P4Hs are expressed in a tissue-specific manner and at different developmental stages. Whether P4Hs have additional substrates in Drosophila remains to be determined.

Last, we previously included secreted factors in our definition of the matrisome, since the ECM is recognized as a reservoir of growth factors and other soluble factors [102]. These 75 proteins (Supplementary Table 5) were defined using a combination of orthology or homology annotations, GO terms, literature references, and the presence of characteristic domains not previously used to define secreted factors but identified from the examination of the *Drosophila melanogaster* extracellular domain database (FlyXCDB http://prodata.swmed.edu/ FlyXCDB/info.list.new21_26.html, [103]). These domains are the PDGF/VEGF domain (IPR000072), the Spaetzle domain (IPR032104), the von Willebrand factor type C (IPR029277), the insulin-like domain (IPR016179), the eclosion hormone domain (IPR006825), and the interleukin-17 family domain (IPR010345).

Classification of Drosophila genes with no mammalian orthologs or homologs

Since the chitin-based ECMs, eggshell, and salivary glue are all secreted from the apical side of epithelial tissues, we classified both the structural and regulatory proteins associated with these ECMs under a category termed "Apical ECM" (Supplementary Table 5 and Fig. 2B). As a group, these proteins comprise nearly 50% of the Drosophila matrisome. To further subdivide this diverse group of proteins, an additional level of classification was created (Supplementary Table 5, column C) to reflect their respective proteins' domain structure, enzymatic function, or localization. The chitinbinding-domain-containing proteins and R&R chitinbinding-domain-containing proteins families refer to proteins containing InterPro domains IPR002557 and IPR031311, respectively. The Chitinase and Chitinaselike families also have a group of defining domains, the chitinase II domain, IPR011583, and three glycoside hydrolase domains, IPR029070, IPR001223, and IPR017853. The Tweedle family represents the only proteins with the domain DUF243 (IPR004145) [104]. Chitin deacetylases were identified based on the presence of a glycoside hydrolase/deacetylase domain (IPR011330). A group of 11 zona-pellucida-domaincontaining proteins was also identified [105]. These proteins have a shared structural attribute, the zona pellucida domain (IPR001507), which we originally used to identify core components of the mammalian matrisome. However, since zona-pellucida-domaincontaining proteins do not present clear orthology or homology with mammalian proteins, we classified them apart.

Groups without clear structural similarities were classified by other means. Proteins of the cuticle that did not meet the definitions above were classified by their shared GO term, chitin-based cuticle development (GO:0040003). Included in this class were also a number of genes reported to be cuticle proteins of low complexity [106,107]. The eggshell superfamily includes two protein classes, corresponding to the vitelline membrane and chorion layers of this ECM, respectively [68,108,109]. The vitelline membrane proteins were defined by GO term or literature search. The chorion proteins had all previously been assigned chorion-related GO terms and chorionrelated protein names, except Cp38 which has chorion in the name and is cited [108]. Finally, 11 proteins including new-glue and salivary glue secretion proteins, Eig71Ee [69,110], and the newly identified tandem paralog of Sgs5 (FBgn0038523) [111] were classified as glue proteins.

Accessing the *Drosophila* matrisome and utilizing it to annotate large datasets

The Drosophila matrisome is available from three sources

To facilitate the use of our definition and categorization of the Drosophila matrisome by the scientific community, the list devised here has been made available through three public platforms. Similar to the matrisome lists of human, mouse, zebrafish and C. elegans, the Drosophila matrisome list can be found on the Matrisome Project website (http://matrisome.org) [11]. Moreover, it has been implemented in two databases widely used by the Drosophila community. The Drosophila matrisome is available within the "Gene Groups" section of FlyBase (FB2019 04, accessible at: https:// flybase.org), which is the most comprehensive source of genetic information for this model organism [71,72,112]. In addition, as a result of the Matrisome analysis, two new terms were added to the Gene Ontology Cellular Component aspect: chitin-based extracellular matrix (GO:0062129) and adhesive extracellular matrix (GO:0062130), allowing more precise GO annotation of the constituents of these specific types of ECM. All Drosophila cuticle proteins and glue genes have now been annotated with these respective terms in FlyBase.

The *Drosophila* matrisome is also available in the Gene List Annotation for *Drosophila* (GLAD, accessible at: https://www.flyrnai.org/tools/glad/web/) database, which is maintained by the Perrimon laboratory to enhance the utility of the cell-based RNAi screening (DRSC) and in vivo fly RNAi (TRiP) collections for the community [73]. For consistency with the current GLAD nomenclature, the matrisome forms a new gene list/group; the matrisome divisions are listed as sub-groups, the categories as sub-sub-groups, and the families are listed under comments.

The Drosophila matrisome provides a powerful tool to annotate large datasets

One powerful application of the matrisome list for any species is in the annotation of large -omic datasets [11]. Thus, as a proof of principle, we used the newly defined *Drosophila* matrisome to re-

evaluate two recently published datasets that focus heavily on ECM-associated proteins. In the first study, Baycin-Hizal and colleagues identified 399 N-glycosylated proteins of the Drosophila head region using solid phase extraction of N-linked glycopeptides coupled to LC-MS/MS [113]. They reported that 4.5% of the proteins identified experimentally in their study were part of the ECM. We found, however, that 13% of the proteins they identified (which included 8 of the 26 glycoproteins and 2 of the 3 proteoglycans we have predicted) are in fact matrisome proteins, more than double the original number. In the second study, Sessions and colleagues reported changes in the abundance of ECM proteins in the Drosophila heart during aging [114]. 104 of the proteins detected were identified as ECM proteins using the Software Tool for Rapid Annotation and Differential Comparison of Protein Post-Translational Modifications (STRAP PTM) developed by Spender and colleagues. Of these 104 proteins, 27 are part of the matrisome, whereas 77 are not. Examination of these 77 proteins revealed that most are in fact localized intracellularly, with little evidence to support that they are ECM components. We retrieved the raw mass spectrometry data from the ProteomeXchange repository (PXD006120) and reannotated the data using the matrisome list. We identified a total of 46 matrisome proteins, finding 19 additional proteins not originally annotated as belonging to the ECM. Together, these two examples demonstrate the power of our matrisome list to comprehensively annotate large experimental datasets. We thus propose that the use of our annotations and nomenclature would assist in the comprehensive identification of ECM signatures contributing to cellular, physiological and pathological phenotypes.

Conclusion

We propose here an *in-silico* definition of *Dro-sophila melanogaster* matrisome that comprises 641 genes encoding ECM and ECM-associated proteins. We further propose their comprehensive classification according to structural and/or functional features. Of note, and as it has been the case with the human and mouse matrisomes, this list is meant to evolve as we gain knowledge in the functions of these genes. We hope that this list and nomenclature will aid with the annotations of large datasets, and thus further our understanding of the roles of the ECM in fundamental biological processes and pathophysiology.

Supplementary data to this article can be found online at https://doi.org/10.1016/j.mbplus.2019. 100015.

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