Proteomics in evolutionary ecology

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Proteomics in Evolutionary Ecology

Running title: Evolutionary dynamics in the proteome

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Evolutionary dynamics in the proteome

Abstract

Evolutionary ecologists are traditionally gene-focused, as genes propagate phenotypic traits across generations and mutations and recombination in the DNA generate genetic diversity required for evolutionary processes. As a consequence, the inheritance of changed DNA provides a molecular explanation for the functional changes associated with natural selection. A direct focus on proteins on the other hand, the actual molecular agents responsible for the expression of a phenotypic trait, receives far less interest from ecologists and evolutionary biologists. This is partially due to the central dogma of molecular biology that appears to define proteins as the ‘dead-end of molecular information flow’ as well as technical limitations in identifying and studying proteins and their diversity in the field and in many of the more exotic genera often favored in ecological studies. Here we provide an overview of a newly forming field of research that we refer to as ‘Evolutionary Proteomics’. We point out that the origins of cellular function are related to the properties of polypeptide and RNA and their interactions with the environment, rather than DNA descent, and that the critical role of horizontal gene transfer in evolution is more about coopting new proteins to impact cellular processes than it is about modifying gene function. Furthermore, post-transcriptional and post-translational processes generate a remarkable diversity of mature proteins from a single gene, and the properties of these mature proteins can also influence inheritance through genetic and perhaps epigenetic mechanisms. The influence of post-transcriptional diversification on evolutionary processes could provide a novel mechanistic underpinning...
for elements of rapid, directed evolutionary changes and adaptations as observed for a
variety of evolutionary processes. Modern state-of the art technologies based on mass
spectrometry are now available to identify and quantify peptides, proteins, protein
modifications and protein interactions of interest with high accuracy and assess protein
diversity and function. Therefore, proteomic technologies can be viewed as providing
evolutionary biologist with exciting novel opportunities to understand very early events
in functional variation on cellular molecular machinery that are acting as part of
evolutionary processes.

Key words: Evolution, natural selection, sexual selection, peptide mass spectrometry,
protein diversity, post-translational modification, protein-protein interaction.

Introduction

Evolutionary theory as initially formulated by Charles Darwin [1] has become a
foundation for biological sciences and ranks among mankind’s most important scientific
discoveries. The empirical support for evolutionary theory shows that traits under natural
selection require two characteristics that make them evolvable: variation and inheritance.
For evolutionary processes such as for example host-parasite / predator-prey coevolution,
sexual selection or ecological adaptation to occur, phenotypic variation between
individuals needs to be generated and maintained for a trait so that selection can
differentially act upon them. Furthermore, traits need to be heritable so that individuals
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with advantageous characteristics pass these onto the next generation and thereby change
their frequency within a population, resulting in fundamental biological developments
such as adaptation and speciation. The discovery of DNA by Watson and Crick [2] as the
molecule responsible for biological information storage and inheritance offered biologists
the possibility to develop and use a range of molecular tools such as PCR, sequencing
and cloning to understand the implications of Darwinian thinking at the molecular level
and across evolutionary timescales. Crick also formulated the central dogma for
molecular biology [3] (Figure 1), where heritable information is coded as genes, typically
DNA but sometimes RNA, from which proteins are produced via transcription and
translation. The central dogma presents proteins as the endpoint of information flow
where any changes are not translated back to RNA and DNA and thus proteins are
typically not considered as drivers of evolutionary processes. As a consequence of this
history, evolutionary biologists are predominantly gene-focused and the technical
opportunities to aid their study of genes and genomes have developed at breathtaking
speed up to the present day. The gene mutation paradigm as the key to evolution has
dominated modern molecular biology, there have been prominent thinkers such as Woese
[4-6] however who proposed RNA and translation to protein as central drivers of
phylogenetic relationships in the tree of life. Woese also highlighted the role of horizontal
gene transfer between prokaryotic cells (i.e. the swapping of DNA encoding for a whole
new protein in bacteria and archea, Figure 1) as more critical to great swathes of
evolution than point mutation of the organism’s own genes [4, 5]. Horizontal gene
transfer allows for rapid evolution to occur at the level of the ecosystem rather than the
level of the organism and the introduction of an entirely new protein agent into a cellular
milieu and indeed into a protein network. The importance of horizontal gene transfer in fungal evolution [7] and even in very recent examples of grains in pathogenicity of fungi is well documented [8].

The full genome sequences of thousands of species are now available allowing unprecedented base by base comparison of genes within and across families, genera and kingdoms and increasingly more sophisticated methodologies are also available to permanently or transiently manipulate gene sequence and expression to observed the effects. However, while this genomics has generated solid empirical evidence for evolutionary theory and provided detailed insights into evolutionary dynamics (e.g. [9]), a range of more fundamental questions still remained unresolved. For example, comparative genomics reveals that many genes often remain remarkably similar throughout evolutionary history, providing, for example, only preliminary answers to the question of why chimpanzees are chimpanzees and humans are humans based on DNA sequence alone [10, 11]. It is widely acknowledged that regulation of expression of genomes is the key differentiator between mammals but it remains unclear, how differences in gene expression of an identical gene pool can generate the tremendous phenotypic variation observed such as for example between humans and chimpanzees [10, 11]. Furthermore, dependence of molecular evolution of DNA on random mutations alone resulting in the eventual appearance of a gene with superior functionality [12-14] would relegate evolution to depend predominantly on chance events acted on by selective forces across generations. In our view while the focus on point mutation alone is weakening, the current framework pursued by many researchers still provides an
unsatisfying and insufficient explanation for fast co-evolving traits such as for example those under sexual selection or host-parasite co-evolution, where heritable changes in phenotype can often become visible within a handful of generations [15-19].

The predominance of evolutionary studies still focuses on genes and genomes through measures of mutation rates and genotype frequency changes in populations. There is no current evolutionary framework or substantial research literature to understand the importance of the role of translated agents – the proteins and their function - as drivers of adaptation. This can be very simplistically illustrated by the co-occurrence of ‘genome and evolution’ and ‘proteome and evolution’ in PubMed: a close to 50-fold difference in co-occurrence exists. Woese [4] pointed out a similar dilemma for RNA biology a decade ago where the importance of studying the evolution of translation of RNA to protein did not fit within the molecular biology paradigm. As he pointed out in his seminal contribution, “molecular biology has to bring evolution to the fore and integrate it fully – not hold it at arm’s length” [5]. Much has changed to resolve this as the explosion of data on numerous levels of RNA biology and the biological role of non-translated RNAs in influencing DNA [20] (Figure 1) has revealed a modern ‘RNA world’ in eukaryotes to mirror the ancestral RNA world at the time of archea and bacteria divergence [21, 22].

In a similar way, we contend in this review that proteins are crucial molecules to study directly when addressing the scientific questions typically investigated by evolutionary biologists for a variety of reasons. First they normally represent the functional units (“the agents”) at the molecular level that are directly responsible for a phenotype seen on the
macroscopic scale. Secondly, most environmental factors, that are not direct mutagens, act firstly on proteins and only secondarily on the genome. Thirdly, proteins are responsible for determining transcriptional competency of significant portions of a given genome by controlling eu/heterochromatin modulation and thus access of the transcription machinery (Figure 1). This becomes increasingly more important given that work over the last two decades has revealed that genes can produce a substantial variety of proteins with fundamentally differed functions through both post-transcriptional processing and post-translational modifications [23]. The idea of post-translational marking as a driver of protein-protein interaction and of this marking as a ‘protein interaction code’ has been proposed based on a range of examples in yeast [24]. Thirdly, the state and function of a specific protein is influenced by both the proteome as a whole and the influence of the prevailing environment. Fourthly, proteins are the agents for epigenetic marking of genomes through histone modification and DNA methylation [25-27] (Figure 1), and hence the proteome has the potential for trans-generational influence both as the end product of the genome and also as epigenome modifying agents. Fifthly, and as pointed out by several recent reviews [28-31], advances in protein biochemistry now allow the assessment of the abundance, location, modification and function of proteins, from isolated single proteins to complex mixtures from whole tissue extracts [32, 33]. Proteomic technologies therefore provide capacity for the study of biological mixtures of proteins and the use of a range of separation techniques from gel electrophoresis to liquid chromatography coupled to different types of mass spectrometry to analyze, quantify and identify differences in the proteome [28, 34, 35]. Separations in gel, in liquid media and on solid surfaces provide physical arrays of proteins for
comparison of differences samples. Peptide mass spectrometry allows the pattern
matching based identification of peptides to track them back to all the specific genome
loci that encode them [36]. Increasing, the techniques of proteomics also allow the
assessment of smaller samples, faster and more accurately, and population level analysis
of individuals is already a reality [37]. Mass spectrometry can identify and quantify not
only the abundance of proteins, but also many modifications to proteins induced by
ecological stimuli and through genetic susceptibility to modification [32]. Changes in the
partnering and strength of protein-protein interactions will also soon be able to be
predicted, detected and quantified [38].

If we consequently consider protein synthesis and maturation as quantifiable mechanisms
to produce natural variation in gene products that selection can act upon and that can be
inherited, we need to integrate the analysis of proteins and their functions into a larger
framework for evolutionary biology using readily available molecular systems biology
approaches (see Table 1 for some examples). In our view this generates very exciting
opportunities for future research. There are a number of textbook examples where
proteins are involved in dynamics of evolutionary importance. For example trans-
generational movement of antibodies initiating immune competence from mother to child
[39], or Darwinian evolution of prion proteins in response to host competition [40]. But
what we will discuss in more detail here is that proteins more generally have an
underlying and fundamental role in phenotypic variation that is acted upon by evolution
across all species and considerable work is required to build and understand the
mechanisms that underpin it.
To build a strong framework of connection between proteomics and evolutionary theory we start with a general and understandable introduction into protein dynamics and protein networks and then consider the proteomic technologies and their possibilities and limitations for use by evolutionary biologists (Table 1). To exemplify some of the theoretical considerations we then provide our own experience in attempting to bridge these worlds through work on reproductive processes and pathogen susceptibility in two model insects – the honeybee *Apis mellifera* and the leaf cutter ant *Atta colombica*. In the later paragraphs we point out the major questions and challenges that remain to be studied. Our main aim in writing this paper is to encourage biologists of all fields to consider recent advances in understanding of proteins and the broader field of proteomics for their future work on evolutionary questions.

**Proteins are more diverse than the genes that encode them**

The primary structure of a protein is a string with each position occupied by one of 23 different amino acids. Each is decoded from defined sets of triplet bases containing one, two or three of the four bases of DNA. As macromolecules, proteins have a huge range of size, complexity and a much wider range of physical properties than the DNA that encodes them. Individual proteins can interact with substrates and products in catalysis, or with structural partner proteins through surface residue interactions. Different proteins can operate in aqueous or highly hydrophobic environments and in a temperature range of
over 120° C, from less than -20° C in arctic and alpine species to more than 100° C in species within volcanic vents [41, 42]. It is widely presumed by evolutionary biologists that proteins have little plasticity of function, no memory and very limited ability to adapt to changing molecular environments. However, protein biochemistry shows this is not really the case [43]. The functioning of proteins can generate alterations of their native conformation that can impact their future function. Proteins can also have their kinetic characteristics altered by external stimuli, thereby changing the way they interact with substrates and products. This can occur through covalent modification of amino acids by processes including phosphorylation, acetylation or glycosylation [24], or reversibly by allosteric activation or inhibition through the binding to proteins of small molecules other than substrates or products [44].

**Proteins are organized in networks and groups of networks**

Complex phenotypic traits are not caused by a single protein but are usually the result of an organized co-occurrence of a set of proteins (Figure 1). Collectively such sets of proteins constitute functional networks with further aspects of flexibility. A protein network can have nodes and hubs, multiple inputs and outputs, and can operate at different states of flux depending on the starting conditions. One new enzyme added to the network can even reorganize the node-wiring diagram, bridge between pathways and thus impact the network flux more dramatically than its individual role as a single protein. Horizontal gene transfer is a means of introduction of new protein players into networks from other organisms in an ecosystem (Figure 1) [4, 6], and a raft of insights into the post-translational evolutionary processes regulating protein abundance have
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recently been reviewed [45]. The complexity of such networks is further increased because proteins often have multiple functions and can therefore be found in multiple proteomic networks, providing links and possibilities for interactions between them.

The presence of complex proteomic networks underlying macroscopically observed phenomena has several potential consequences: The different subtasks of networks are at risk of interfering with each other, both at the level of the proteins themselves and at the level of the metabolic pools that interconnect them. For example, proteases involved in host-parasite interactions can damage other proteins both in the host and parasite [46, 47], while enzymes producing free radical species can cause oxidation of other proteins through their reactive products [48]. Consequently we might expect evolutionary trade-offs to be apparent within protein networks and the ways network components can evolve is likely to be more restricted for those components that fulfill core functions in each network. This means the evolutionary clock will tick at different speeds for the corresponding genes coding for these proteins within a given network, depending on the biological activity of proteins or subgroups of proteins and their degree of connectivity to functional networks. Interestingly the high frequency of selection of functional modifications in metabolic enzyme loci associated with the TCA cycle and glycolysis and their secondary roles in cells has already been highlighted [49].

The potential for selection acting on proteomic networks
A problem we face is how best to describe the functioning as well as the evolvability of protein networks in a way that can illustrate the action of selective processes. Mathematicians use matrices to define the condition of an N-dimensional structure. By the same analogy we can describe a proteomic network as an N-dimensional matrix where each individual protein or pathway represents an additional dimension (vector) within the matrix and values within the matrix define individual network characteristics such as metabolic flux. The matrix is initially defined by genotype, but after its formation, it’s phenotypic expression can be represented in several different ways and powerfully influenced the interaction of its components:

Firstly, a protein matrix can be seen as information processing machinery. Input variables (an input vector that consists of internal or environmental stimuli) are entered into the matrix and ultimately produce a response variable with a phenotypic expression on the cellular or individual level. The stimulus could be a protein or a metabolite (or several or a combination) that enters a biochemical network and finally generates a product that has biological activity; for example, a neurotransmitter to influence development, or a metabolic substrate or antioxidant to nurture or defend a cell or tissue.

Secondly, a protein matrix can be seen as an information storage device. In such a case the input variables can be seen as a matrix as well and the interaction of matrices produce a new but altered one. This change, dilution or modification will affect the possible output from what has been mentioned above. These modified functions can be “stored” within the matrix in a short time frame. For example proteins within networks could be
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degraded by proteases or post-translationally modified or allosterically activated or inhibited, and thereby biochemical pathways blocked or metabolic flows diverged that would normally have been expected as physiological responses. Multiple modifications at closely arranged sites can ultimately represent logic functions, building logic gates that can impact the protein network function [32].

Evolutionary dynamics of protein network matrices

Our discussion about protein networks and selection acting on them as presented in the last chapters also allows us to propose elements of a continuum, starting from instant physiological effects influencing a trait under selection right through to the transgenerational dynamics that evolutionary ecologists typically investigate in their macroscopic trait studies. However, for biochemical credibility, such a pathway needs definable and measurable components that underpin mechanisms. In our view, many of the elements needed are already known and simply need to be placed within a connected framework of evolutionary biology as outlined below.

A protein matrix is able to respond to its environment rapidly in seconds to minutes, such as for example the detection of the presence of a pathogen. This produces a matrix output that selection can operate on in minutes; for example, an oxygen radical is detoxified by antioxidant machinery, or alternatively it damages a protein altering the flux through a metabolic pathway, or a protein in phosphorylated changing its function or interaction partners. The manipulation of the protein matrix then moves it into a new equilibrium or
phenotype with altered attributes over a period of hours; for example changes in metabolic pools which can then feedback or feed forward on gene expression providing new components to the matrix. Prolonged changes in gene expression will alter the proteome, but can also influence both the mRNA pool and the smallRNA and miRNA pools. Through mechanisms such as RNA-directed DNA methylation or alterations of histone modification in regions of active transcription, over days to months, these altered gene expression changes can modify epigenetic marks on the DNA leading to altered epigenetic states of specific cells. Methylation patterning and corresponding changes in histone binding and modification can in some cases be heritable, providing a transgenerational flow of protein matrix attributes. There is also a further stage of this process, as there is now clear evidence that methylated regions of DNA have higher mutation rates [50] due to spontaneous deamination of methyl-cytosine and subsequent mismatching during DNA duplication to replace a CG couple with an AT couple, yielding a single nucleotide polymorphism. Hence changes to proteome matrices that can feedback to gene expression have the potential to influence the evolution of genes that influence them in a targeted fashion. Examples of the role of post-translational modification processes [51] and of DNA methylation [52, 53] in evolutionary processes have been reviewed.

Evolutionary biology from a protein perspective:
The ideas and hypotheses raised so far in this paper originate from the literature that has combined protein biochemistry with evolutionary ecology. Such empirical cross
disciplinary work conducted over the last decade provides a number of interesting examples of how questions derived from evolutionary biology have been approached using a combination of discovery based proteomic analyses and the typical, hypotheses driven approaches used in evolutionary ecology (see Table 1 for more details and examples). For example, proteins have been shown to be phenotypically plastic, if they lack a defined stable secondary or tertiary structure [54]. Regions within proteins that show such plasticity play a key role in protein-protein interaction networks, which might provide functional advantages. Because of their higher capacity to rewire with other protein partners, they also seem to evolve rapidly [54]. Proteomics has also been used in phylogenetic studies to understand the evolution of traits of interest, for example key physiological traits such as protein translation throughout the tree of life [55]. Proteomic approaches revealed how individual proteins such as Hsp90 can impact developmental networks and thereby influence morphological phenotypes and their evolution [56]. An interesting example of protein based inheritance has recently been reported in the honeybee *Apis mellifera*, where the egg yolk protein vitellogenin can bind to a bacterial bee pathogen and is used as a carrier of microbial fragments to eggs, resulting in immune priming of offspring [57].

Sexual reproduction typically involves behavioral interactions by two or more individuals on the macroscopic level but numerous key processes are occurring at the level of proteins [58]. These interactions are typically characterized by the presence of strong selective forces acting on individual fitness and by fast evolutionary change. Males have to provide females with sufficient numbers of viable sperm but often compete against
each other within the females’ sexual tract for access to eggs. Consequently, evolutionary ecologists had a long standing interest to untangle those traits that support the cooperative aspects of sexual reproduction from those responsible for the conflicts over paternity and understand their impact and male and female reproductive success. Proteomics has already been used in a wide range of different organisms to identify male reproductive proteins, which provided key insights into the make up of ejaculates [59-70] and triggered studies to unravel their effects on females and reproductive success [69, 71-75].

The identification of these proteins and a more detailed understanding about their function has enabled comparative studies to investigate the evolutionary history and phylogenetic relationships for some of these proteins [76-78].

In the next section we will further elaborate on such reproductive proteomics by summarizing our own findings, which offers us an opportunity to discuss our strategy and intent behind our scientific progress. Furthermore, commentary on the timeline of discovery and our unpublished and pilot data allows us to provide a more general overview of the benefits of proteomics for evolutionary research. Proteomics was without doubt the key tool we used over the last 10 years to gain major insights into highly complex biological processes and to begin to hone in on proximate mechanisms that underlie reproductive traits such as high quality sperm and long-term sperm storage. We started with the identification of lists of proteins in samples of interest. While this is often flagged as a limitation of proteomics, because it cannot provide causal relationships, it provided our protein landscape for the following years. As we illustrate below in detail, these parts lists guided consequently phenotypic studies providing not only answers to
our initial questions but a substantially broader and systemic insight into the molecular function of social insect reproduction.

**An example from our own research on insect reproduction**

We studied sexual reproduction in Hymenopteran social insects, being the eusocial ants, bees and wasps. Their societies are characterized by the presence of a single or very few reproductive animals, typically referred to as queens, and a non-reproducing cast known as workers [79-82]. Workers benefit from helping if they are related to the helped individual, known as inclusive fitness benefits [83, 84]. Because helping incentive increases with helper relatedness, social insects queens perform only a single round of mate choice and sperm acquisition and never replenish sperm once they have started to lay eggs. This has resulted in the evolution of spectacular reproductive adaptations in species with large and long lived societies, where males produce exceptionally large ejaculates of high quality [85], and queens store them for decades to sire millions of offspring [86, 87]. Although these reproductive traits must have been key during social evolution and will have contributed towards the remarkable ecological success of these animals [88], the proximate mechanisms to achieve such high levels of fertility or their evolution had remained completely unknown.

During mating males transfer ejaculates to females that consist of sperm and glandular secretions known as seminal fluid, which has increasingly been recognized to contain key molecules determining male and female reproductive success [89-92]. Seminal fluid can also produce mating plugs or mating signs [81, 93-96], which influence the mating
behavior of bumblebee queens [97, 98], providing the first evidence that seminal fluid components are important in social insects.

With this background we started our collaborative work together by a discovery inspired proteomic characterization of seminal fluid in honeybees [99], revealing a first insight and basic parts map of its molecular landscape. This list of proteins proved to be of significant scientific value because functional analyses predicted biological functions and generated a wealth of hypothesis and ideas that consequently guided our further experimental work. We found that seminal fluid proteins consisted of three major groups of proteins, that (1) keep sperm alive (2) defend sperm and queens from pathogens and (3) are molecular agents of sexual conflict. Follow up experimental work confirmed that seminal fluid is indeed exceptionally potent in keeping sperm alive and that proteins are the key molecules responsible for this effect [100, 101]. The detection of antimicrobial proteins in the seminal fluid triggered a search for pathogens in honeybee ejaculates, and resulted in the identification of two widespread bee pathogens in ejaculates, *Nosema apis* and *Nosema ceranae* [102, 103]. Our finding of antifungal proteins in seminal fluid implied that males may be able to combat these infections in their ejaculates. Experimental follow up showed that seminal fluid is indeed remarkably efficient in killing *N. apis* spores (Peng et al., submitted). Moreover, our data revealed that the pathogen is killed in at least two distinctly different ways, implying that there is redundancy in the defense system of seminal fluid as well as some specificity, because the biologically active molecules show no antimicrobial activity against a series of non-pathogenic microbes. Both redundancy and specificity of honeybee antimicrobial proteins were novel findings that had not been reported previously for insects. Thirdly, we found
molecules that we predicted to be involved in sexual conflict. The presence of ejaculates from multiple males within a female’s sexual tract can result in postcopulatory sexual selection, operating either as sperm competition [104] or cryptic female choice [105]. We hypothesized that the battlegrounds of these events are extracellular spaces dominated by secreted proteomes and the effective role and variability of these protein sets would be defining paternity success. We indeed found that seminal fluid proteins of polyandrous honeybees and leaf cutter ants are capable to kill sperm of rival males, known as sperm incapacitation [106].

This shows how our initial identification of proteins in the seminal fluid of honeybees generated a number of predictions about function, which were accurate because we were able to confirm the expected phenotypes through follow up experiments. The proteins identified were confirmed to be the biologically active molecules, and our functional analyses have already provided subsets of target proteins for further study [107]. Finally, our work indicated that seminal fluid proteins or protein networks interact with other proteomes, such as those of rival ejaculates, the queen or parasites, encouraging us to also identify these additional proteomes.

The honeybee sperm proteome revealed the presence of a very distinct subset of proteins [108], many of them being related to energy metabolism. Their high abundance in honeybee sperm implied that the survival of high quality sperm is closely associated with energy production. When we quantified the effect of some of these proteins on sperm metabolism we were able to confirm that these proteins are biologically active and are key for sperm survival. A second group of abundant proteins we detected in honeybee sperm are related to transcriptional or translational activities, which was surprising given
that sperm is often believed to be translationally and transcriptionally silent. Providing sperm with radiolabeled amino acids confirmed that sperm indeed produce proteins at a low rate [109].

As well as producing high quality sperm, social insects are also the ‘world record holders’ for storing sperm [81, 86, 87]. To achieve this, queens provide sperm with spermathecal fluid, a glandular secretion continuously added to sperm during storage. The proteomic profiles of spermathecal fluid was distinct from that of seminal fluid [110] indicating that sperm are able to survive in two very different biochemical environments. The seminal fluid proteome forms a loosely connected network of proteins, consistent with the expectation that these proteins are responsible for more individual tasks such as keeping sperm alive, killing rival sperm and parasites or manipulating female physiology [100, 101, 106, 110, 111]. The spermathecal proteome on the other hand keeps sperm alive for years, and its high connectivity seems to provide a biochemical environment that has been selected for maximized sperm survival [110]. We therefore expected the proteome of sperm to adapt to these changes in their biochemical environment. We have now confirmed this experimentally [107], and key enzymes with changed abundances were as expected related to energy production. Consequently, we were able to provide important molecular insights into the secrets of long-term sperm storage, which were facilitated by the presence of a relatively small number of enzymes that maximize ATP production and minimize oxidative stress.

Apart from the proteome differences between stored and ejaculated sperm we have also
found other proteomes to be remarkably plastic, for example between seminal fluid samples from males of different bee lineages [111] or between seminal fluid of infected and uninfected males (Grassl et al in preparation). These studies reveal that the proteomes of ejaculates provide a useful model system to understand proteomic networks and their functioning in an evolutionary framework, because the proteomes investigated are substantially smaller than those of entire cells, organs or organisms, and the strong selective forces or ecological stimuli that impact them result in fast changes in these secreted proteomes and functional changes in proteomic networks that can be measured in vitro. These studies merely mark a starting point and demonstrate the feasibility of fruitful collaborations between proteomics and evolutionary biology researchers – what we plan now are studies that quantify the phenotypic variation of proteomes and to quantify their hypothesized heritability. Furthermore, the identification of individual proteins or protein networks of interest now allow us to also conduct phylogenetic comparisons to understand the evolutionary history of proteins and the way underlying proteomic networks co-evolved with traits of interest, similar to what has been done in other species [77].

Our work on evolutionary proteomics revealed that proteomic investigations tend to take substantially longer to conduct and publish, as they generate larger datasets that require substantially more time to analyze than classical studies typically conducted in behavioral ecology. However, such detailed data mining resulted in the development of predictions and hypotheses and the accuracy of these predictions was found, in time, to be highly consistent with the mechanisms uncovered. Consequently, proteomics can also be used as a highly efficient and accurate tool for the development and formulation of testable
The future of evolutionary proteomics

Developing a framework for how proteins play an important role for evolutionary processes is critical for engagement of researchers with expertise in each area. Biochemical approaches to identify and study proteins and their abundance and functionality have been rapidly developing over the last decade and are now much more easily accessible for a wider range of scientists including evolutionary biologists [30, 31, 112]. Academic institutions as well as private companies offer collaborations and services to analyze protein samples of interest. With this technical feasibility, proteomes can be studied in an evolutionary framework that does not differ in any major way from any other phenotypic trait of interest studied by evolutionary biologists over the last 150 years. What is still needed are larger scale experimental studies quantifying natural variation in protein profiles within and between individuals of the same and different populations in order to understand how much of the theoretically achievable variation in proteomes is actually realized and selected for. Furthermore we need a better understanding of how the interactions between environmental factors and an individual’s proteome operate, both in the short as well as in the long term. Such studies will provide crucial insights into an individual’s opportunity to actively respond and adapt to changing environments and can test for the degree of heritability of modified proteomic networks characteristics. Finally, studies are needed that quantify the overall effect of proteins on evolutionary processes in order to understand whether they are only important for a
subset of processes such as sperm competition and immunity or whether they are in fact additional drivers of evolution more generally.

Summary

The traditional split between biological sciences focused on either proximate or ultimate questions is starting to diminish. One of the reasons for this is that molecular insights into how life works have exponentially grown over the last decade, due in significant measure to spectacular technological breakthroughs that now allow the study of molecular dynamics in cells and entire organisms and even their extrapolation to wider habitats and ecosystems. Genomics has provided evolutionary biologists with new and exciting opportunities to understand and investigate evolutionary concepts. The rapidly evolving field of proteomics now needs researchers with evolutionary questions to link protein networks and their functioning to complex organismic characteristics.

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Table 1: A focus on proteins as drivers of ecological and evolutionary processes offer researchers from various fields novel opportunities to explain biological phenomena such as rapid evolutionary and adaptive changes, especially in cases where the traditional focus on DNA mutation followed by natural selection provide unsatisfying proximate explanations for observed phenotypes. Furthermore, molecular biologists focusing on the use of -omics approaches to understand variation in their datasets can use evolutionary and ecological explanations in hypothesis building for followup experiments.

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<th>Literature</th>
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<td>(Proteome changes over generations)</td>
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<tr>
<td>Evolution of traits</td>
<td>Proteins modify genomic information flow through DNA modifications and structure</td>
<td>[50, 55, 56]</td>
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<td>Heritability</td>
<td>Proteins are all genome derived but can also act as non-genomic components of transgenerational inheritance</td>
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<td>Natural Selection</td>
<td>Proteins are the biologically active “agents” and proximate drivers of fitness</td>
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<td>Genotype x Environment Interactions</td>
<td>Protein networks are susceptible to environmental stimuli</td>
<td>[107]</td>
</tr>
</tbody>
</table>
Evolutionary dynamics in the proteome

<table>
<thead>
<tr>
<th>Host-Parasite Interactions</th>
<th>Protein diversity can be a driver of virulence and immunity</th>
<th>[8, 40, 47]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phenotypic Plasticity</td>
<td>Protein networks and matrixes are highly variable and change due to genomic and ecological factors</td>
<td>[23, 49], [54, 115]</td>
</tr>
<tr>
<td>Sexual Selection</td>
<td>Proteins can be molecular drivers of conflicts over paternity</td>
<td>[71-73, 116]</td>
</tr>
</tbody>
</table>
Figure Legends

Figure 1

The Central Dogma coupled to other regulatory steps and mechanisms of environment-dependent variation that influence the proteome. The central blue box presents the primary flow of molecular information as found in all living organisms, through which DNA encodes for genes that are transcribed into RNA which in many cases are translated into proteins. The latter are principally responsible for the expression of a specific phenotype. Research over the past decades has now shown that this central protein building system is augmented by a range of more dynamic protein and proteome-modifying factors which are influenced by environmental factors. This presentation highlights the role of proteins and their variation as additional drivers of physiological processes and their evolution rather than simply as end points of molecular information flow.