Gene flow and biological conflict systems in the origin and evolution of eukaryotes

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The endosymbiotic origin of eukaryotes brought together two disparate genomes in the cell. Additionally, eukaryotic natural history has included other endosymbiotic events, phagotrophic consumption of organisms, and intimate interactions with viruses and endoparasites. These phenomena facilitated large-scale lateral gene transfer and biological conflicts. We synthesize information from nearly two decades of genomics to illustrate how the interplay between lateral gene transfer and biological conflicts has impacted the emergence of new adaptations in eukaryotes. Using apicomplexans as example, we illustrate how lateral transfer from animals has contributed to unique parasite-host interfaces comprised of adhesion- and O-linked glycosylation-related domains. Adaptations, emerging due to intense selection for diversity in the molecular participants in organismal and genomic conflicts, being dispersed by lateral transfer, were subsequently exapted for eukaryote-specific innovations. We illustrate this using examples relating to eukaryotic chromatin, RNAi and RNA-processing systems, signaling pathways, apoptosis and immunity. We highlight the major contributions from catalytic domains of bacterial toxin systems to the origin of signaling enzymes (e.g., ADP-ribosylation and small molecule messenger synthesis), mutagenic enzymes for immune receptor diversification and RNA-processing. Similarly, we discuss contributions of bacterial antibiotic/siderophore synthesis systems and intra-genomic and intra-cellular selfish elements (e.g., restriction-modification, mobile elements and lysogenic phages) in the emergence of chromatin remodeling/modifying enzymes and RNA-based regulation. We develop the concept that biological conflict systems served as evolutionary “nurseries” for innovations in the protein world, which were delivered to eukaryotes via lateral gene flow to spur key evolutionary innovations all the way from nucleogenesis to lineage-specific adaptations.

Keywords: antibiotics, biological conflict, endosymbiosis, immunity proteins, restriction-modification, RNAi, selfish elements, toxins

INTRODUCTION

Ever since the emergence of the endosymbiotic hypothesis as the primary model for the origin of eukaryotes there has been considerable interest in two major issues which it brought forth, namely large-scale lateral gene flow and genetic conflicts. While the exact details of the nature of this endosymbiotic event are still debated, by its very nature the endosymbiotic hypothesis implies gene flow between the alphaproteobacterial mitochondrial progenitor and the nucleo-cytoplasmic progenitor of archaean ancestry (Martin and Muller, 1998; Esser et al., 2004; Rivera and Lake, 2004; Aravind et al., 2006; Gabaldon and Huynen, 2007; Pisani et al., 2007; Sapp, 2007). This phenomenon is not just relevant to the origin of eukaryotes, but also several other symbiogenic events that shaped the subsequent evolution of eukaryotes, such as the origin of the primary photosynthetic eukaryotes, including the plants, and the numerous secondary or tertiary photosynthetic eukaryotes (Delwiche, 1999; Palmer, 2003; Bhattacharya et al., 2004; Keeling, 2004; Huang and Gogarten, 2007; Obornik et al., 2009). In the former event, not just the well-known gene flow from cyanobacteria, but also complementary contributions from a chlamydia-like endosymbiont have been postulated (Huang and Gogarten, 2007). Additionally, there are other inter-organismal interactions that have occurred throughout eukaryotic evolution, which have resulted in comparable gene flow, albeit in a more episodic fashion (Anantharaman et al., 2007). Eukaryotes are characterized by a wide-range of close organismal associations. Indeed, cytoplasmic symbiotic bacteria, comparable to the progenitors of the mitochondria and chloroplasts, and infection by several types of large DNA viruses are a common feature of many eukaryotes, including representatives of the metazoan and amoebobazoan lineages (Batut et al., 2004; Collingro et al., 2005; Ogata et al., 2006; Iyer et al., 2006b; Nikoh et al., 2008; Bertelli et al., 2010; Raoult and Boyer, 2010; Schmitz-Esser et al., 2010; Georgiades et al., 2011). There are also examples of some
rather dramatic inter-eukaryotic associations, like endoparasitism as exhibited by apicomplexans, karyoklepsy, or “theft” of chlorophyte nuclei (along with the chloroplasts) observed among ciliates, or karyoparasitism, involving injection of parasitic nuclei into host cells, which is observed in certain rhodophytes (Fields and Rhodes, 1991; Goff and Coleman, 1995; Johnson et al., 2007). Further, it has been noted that the phagotrophic nutrition of many eukaryotes can also result in a more general form of genetic chimerism, facilitated by the constant engulfment of genetic material of particular types of bacteria and eukaryotes (Doolittle, 1998). Yet other eukaryotes, such as the rotifers, appear to even actively engage in uptake and incorporation of genetic material from their environments—in addition to the proposed role in compensating for the lack of sexual reproduction, this phenomenon also serves as a conduit for notable “alien” gene flow (Gladyshev et al., 2008). Thus, it has become increasingly clear in the past two decades that gene flow between distant lineages and the consequent genomic chimerism might have a notable role in the evolution of eukaryotes.

Inter-organismal and intra-organismal genetic conflicts are a quotidian feature across all organizational levels of life (Smith and Price, 1973; Maynard Smith and Szathmáry, 1995; Hurst et al., 1996; Burt and Trivers, 2006; Werren, 2011). In their simplest form they include various trophic interactions between organisms, such as predation. Such conflicts might also arise between different cells of the same species cooperatively aggregating to form a multicellular assembly or developing as a multicellular organism due the emergence of “cheaters,” whose genetic interests do not align with the remaining cooperating cells (Dao et al., 2000). At the level of a single cell, as the interests of different genomes residing within it are not necessarily aligned with each other, there is potential for yet another level of genetic conflicts (Burt and Trivers, 2006). Such conflicts have a long evolutionary history in the prokaryotic superkingdoms in the form of the interactions between plasmids and the cellular genome. However, the endosymbiotic origin of eukaryotes made it one of their quintessential features because it brought together multiple distinct genomes (i.e., the nuclear and mitochondrials) in a single cell (Maynard Smith and Szathmáry, 1995; Werren, 2011). Such inter-genomic conflicts within the cell further expanded in course of eukaryotic evolution due to additional associations introducing interactions with genomes from plastids, nucleomorphs, and endosymbiotic/parasitic and intra-cellular bacterial predators of mitochondria (Sassera et al., 2006; Werren, 2011). In several cases symbiotic bacteria are involved in multi-level cooperation-conflict relationships: For instance, the bacterial symbiont Phototrophbus enables predatory nematodes to feed on insects by killing them with toxins (Bowen et al., 1998), whereas the endosymbiotic bacterium Hamiltonellonea defensa protects aphids against parasitoid wasps by deploying toxins against them (Degnan et al., 2009). Conflicts between the cellular genomes and viruses that exploit them for their own reproduction add yet another dimension to conflicts occurring within cells (Iyer et al., 2006b; Raoult and Boyer, 2010). Finally, there might be genetic conflicts within a single genome itself, arising from a wide variety of selfish elements trying to maximize their own fitness at the expense of the remaining genes (Burt and Trivers, 2006; Werren, 2011). These selfish elements are often characterized by a degree of intra- and/or inter-genomic mobility and assume a bewildering array of forms, including numerous distinct types of transposable elements, restriction-modification, and toxin-antitoxin systems (Kobayashi, 2001; Anantharaman and Aravind, 2003; Burt and Trivers, 2006; Ishikawa et al., 2010; Leplae et al., 2011). The former elements catalyze or facilitate their own proliferation, while the latter elements enforce cellular genomes to retain them by killing cells in which they are disrupted. Despite being primarily selfish elements, they might on occasions confer a fitness advantage to genomes, as this indirectly augments their own fitness (Burt and Trivers, 2006; Werren, 2011).

These conflicts are often directly mediated by particular molecules, either proteins or small molecules which act as “chemical armaments”; although in multicellular forms it might be reflected as morphological features that serve as weaponry (Smith and Price, 1973; Anantharaman and Aravind, 2003; Degnan et al., 2009; Ishikawa et al., 2010; Leplae et al., 2011; Werren, 2011; Zhang et al., 2011; Iyer et al., 2011b). Not surprisingly, each of the many levels of organismal conflict have sparked off intense “arms races” between the interacting organisms (Dawkins and Krebs, 1979), whose signatures are often seen in the form of extensive diversification of the proteins directly participating in, or synthesizing molecules deployed in conflict (Cascales et al., 2007; Zhang et al., 2011). Concomitantly, there is also a similar rapid diversification of proteins directly involved in defending or serving as antidotes against the chemical armaments deployed in the conflict (Anantharaman and Aravind, 2003; Leplae et al., 2011; Zhang et al., 2011; Iyer et al., 2011b). Importantly, both the offensive and defensive molecular adaptations involved in these conflicts can be transmitted between genomes by way of lateral transfer and is an important factor both in the spread of antibiotic production and resistance among prokaryotes (Walsh, 2003; Aminov and Mackie, 2007; Skippening and Ragan, 2011).

The ever-expanding genomic data from both eukaryotes and prokaryotes, along with genome-scale analysis, has considerably elucidated the major trends in the genomic chimerism arising from the bacterial and archaean progenitors of the eukaryotes (Martin and Muller, 1998; Esser et al., 2004; Rivera and Lake, 2004; Aravind et al., 2006; Gabaldon and Huynen, 2007; Pisani et al., 2007). These analyses have particularly helped differentiate the cellular systems which have a primarily archaean provenance (e.g., core DNA replication, core RNA metabolism, and translation) as against those with a primarily bacterial provenance (various aspects of energy, anabolic, and catabolic metabolism). However, uncovering the origins of specific systems, which appear to be eukaryotic synapomorphies (or shared derived characters), have required a somewhat distinct computational approach relying on in-depth analysis of protein sequences and structures (Aravind et al., 2006, 2011; Burroughs et al., 2011). Such analyses revealed glimpses of a collusion between gene flow through lateral transfer and the selective forces acting on molecular players in organismal and intra-genomic conflict in shaping the evolution of key components of systems such as eukaryotic chromatin, RNA-based gene regulation, and certain signaling pathways. However, this aspect of eukaryotic evolution is
considerably under-appreciated. Hence, in this article we present a synthetic overview of: (1) how large-scale lateral gene flow between interacting organisms has facilitated the emergence of new adaptations deployed in inter-organismal conflict. (2) How adaptations developed due to the intense selection for diversity in the molecular participants in organismal and genomic conflicts were dispersed by lateral transfer and subsequently exapted for various eukaryote-specific adaptations. Due to limitations of space, we do not provide a comprehensive survey of all known instances of the above processes. Instead, we attempt to highlight the importance of these processes in the emergence of key adaptations, not just in early eukaryotes, but also during their subsequent evolution, with diverse illustrations emerging from recent investigations. We must emphasize that in this article we mainly use published examples that have been reported in several individual studies on various biological systems or protein families. However, this is the first time they are being brought together to create a coherent picture. A detailed presentation of the methodological apparatus for sequence, structure and phylogenetic analysis of the presented examples is precluded due to limitations of space. However, we refer readers to the individual studies from which we draw our examples for details regarding the computational analysis of the proteins considered here. We use these to develop a conceptual framework for understanding the importance of the diversifying forces acting during biological conflicts in facilitating adaptations that played a role in the so-called “major transitions” of eukaryotic evolutions (Maynard Smith and Szathmáry, 1995).

MATERIALS AND METHODS

Sequence profile searches to establish the relationships between protein domains were performed using the PSI-BLAST (Altschul et al., 1997) and JACKHMMER (Eddy, 2009) programs that run against the non-redundant (NR) protein database of National Center for Biotechnology Information (NCBI). For most searches which were used to report the relationships presented in this work a cut-off e-value of 0.01 was used to assess significance. This was further confirmed with other aids such as secondary structure prediction and superposition on known structures, if available. Protein sequences were clustered using the BLASTCLUST program (ftp://ncbi.nih.gov/blast/documents/blastclust.html) to identify related sequences in gene neighborhoods. Multiple sequence alignments of all domains were built by the Kalign (Lassmann et al., 2009) and PCMA programs (Pei et al., 2003), followed by manual adjustments on the basis of profile-profile and structural alignments. Secondary structures were predicted using the JPred program (Cuff et al., 1998). A comprehensive database of profiles was then constructed using these multiple alignments and was used extensively in the annotation and analysis of protein domain architectures and gene neighborhoods. For other known domains, the Pfam database (Finn et al., 2010) was used as a guide, though the profiles were augmented in several cases by addition of newly detected divergent members that were not detected by the original Pfam models. Clustering with BLASTCLUST, followed by multiple sequence alignment, and further sequence profile searches were used to identify other domains that were not present in the Pfam database. Signal peptides and transmembrane segments were detected using the TMHMM and Phobius programs (Kall et al., 2007). The HHpred program was used for profile-profile comparisons to either unify poorly characterized families of proteins or find homologous structures in the PDB database (Soding et al., 2005). Structure similarity searches were performed using the DaliLite program (Holm et al., 2008). Preliminary phylogenetic analysis was conducted using a rapid but approximate-maximum-likelihood method implemented in the FastTree 2.1 program under default parameters (Price et al., 2010). In-house bench-marking suggested that these results are generally comparable to complete ML implemented in the Phylip (Felsenstein, 1989; Adachi and Hasegawa, 1992). Predicted lateral transfers to eukaryotes were further evaluated for false positives by ensuring they were embedded in contigs or complete chromosome sequences with other genes typical of eukaryotes, comparing exon-intron structure of the genes, studying their phyletic distribution within eukaryotes and comparing the protein distances of the predicted eukaryotic proteins (as measured by bit scores) with bacterial homologs. Structural visualization and manipulations were performed using the PyMol (http://www.pymol.org) program. Automatic aspects of large-scale analysis of sequences, structures and genome context were performed by using the in-house TASS package, which comprises a collection of Perl scripts.

RESULTS AND DISCUSSION

PARASITE-HOST CONFLICTS: EMERGENCE OF APICOMPLEXAN SURFACE PROTEINS FOR HOST INTERACTION DUE TO LATERAL TRANSFER

Apicomplexa are a remarkable clade of alveolate eukaryotes entirely comprised of highly specialized metazoan parasites (Levine, 1988; Vivier and Desportes, 1990). With other alveolates, such as ciliates, colpodellids, Perkinsids and dinoflagellates, they share organelles known as extrusomes, which allow delivery of a payload of proteins into target cells, such as their prey or hosts (Leander and Keeling, 2003). While basal apicomplexans, the archigregarines, are partial endoparasites that insert only the forepart of their cell into the host cells to suck nutrients, the derived apicomplexans are obligate endoparasites that reside entirely within the cells they invade (Leander et al., 2006). Basal apicomplexans typically have a single-host, but many of the derived apicomplexans like the malarial parasite *Plasmodium* and *Theileria* have evolved lifecycles with two distinct hosts (Levine, 1988; Vivier and Desportes, 1990). Genome analysis of multiple apicomplexans ranging from the relatively basal *Cryptosporidium* to the highly derived *Plasmodium* have shown that they have evolved a remarkable set of secreted or membrane-anchored (surface) proteins that interact with host molecules as a part of the invasion process or other cytoadherence events during their lifecycle (Kaslow et al., 1988; Kappe et al., 1998, 1999; Anantharaman et al., 2007; Arredondo et al., 2012). While surface proteins in each apicomplexan lineage show a wide range of lineage-specific domains (e.g., the Rifs and Dbl domain proteins in *P. falciparum*), they also contain a striking array of domains that are also found in surface proteins of animals (Pathy, 1999; Anantharaman et al., 2007) (Figure 1). Case by case phylogenetic analysis revealed that at least 18 types of non-catalytic
domains from apicomplexans are otherwise found only in the animal lineage, or alternatively are most closely related to versions found in the animal lineage (Anantharaman et al., 2007) (Figure 1). Functional studies in metazoans suggest that major- ity of these domains, such as the thrombospondin-1 (TSP1), sushi/CCP, MAM, fibronectin-type 2, scavenger receptor, kringle, and vWA domains are involved in adhesive interactions between proteins or proteins and carbohydrates on the cell-surface (Bork, 1993; Patthy, 1999). More recently structural analysis has revealed that the SRS and s48/45 domains, respectively, from coccidian and aconoidasidan apicomplexans, were probably derived through rapid sequence divergence from the ephrin-like domain found in metazoan signaling molecules (Arredondo et al., 2012). Genome analysis suggests that while some of these “animal-like” domains were acquired early in apicomplexan evolution, yet others were acquired only later by specific lineages (Figure 1) (Anantharaman et al., 2007). This suggests that the acquisition of a structurally diverse, but functionally comparable group of domains from their animal hosts has been a persistent feature of apicomplexan evolution. Although functional studies on apicomplexan surface proteins with animal domains are still in relatively early stages, two major themes are beginning to emerge: (1) Some of these
proteins appear to have a parasite-specific function in relation to their sexual development, such as in gamete fusion (Pradel et al., 2004; Arredondo et al., 2012). (2) Most others have been adapted for a diverse set of interactions pertaining to invasion of host cells or localization to particular tissues and are often secreted via specialized extrusomes of apicomplexans known as rhoptries (Bradley and Sibley, 2007; Santos and Soldati-Favre, 2011). Particularly striking is the recruitment of the TSP1-domain-containing adhesins early in apicomplexan evolution as part of the conserved invasion apparatus that depends on a cytoskeletal gliding motor unique to apicomplexans (Soldati-Favre, 2008).

Genome analysis has also revealed that apicomplexans possess an animal O-like glycosylation system with two separate arms performing the fucosylation and N-acetylgalactosaminylation of hydroxyl groups of serines or threonine on target proteins (Anantharaman et al., 2007) (Figure 1). The first of these has at its core two enzymes, the protein O-fucosyltransferase and a Drosophila fringe-like glycosyltransferase that elongates the initial fucose chain with N-acetylglucosamine (Varki et al., 1999; Luo et al., 2006). Also associated with this pathway is the fucose-GDP transporter that allows parasites to take up fucose (Luhn et al., 2001). Interestingly, this pathway modifies TSP1 and EGF domains, both of which appear to have been acquired by apicomplexans through lateral transfer from animals (Figure 1). The second pathway displays three distinct orthologous groups of proteins, which constitute the enzyme complex that transfers UDP-linked N-acetylgalactosamine to mucin-like target proteins typified by homopolymeric stretches of serines and threonines (Varki et al., 1999). Phyletic and phylogenetic analysis revealed that enzymes of both these arms of the O-linked glycosylation system and the fucose transporter are specifically related to their animal counterparts to the exclusion of homologs from any other lineage (Anantharaman et al., 2007). Furthermore, their phyletic patterns suggest that the glycosylation pathways were acquired in the common ancestor of endoparasitic apicomplexans, though they were either partially lost in haemosporidians or completely lost in piroplasms. Interestingly, in the more basal apicomplexans, like Cryptosporidium and the coccidians, there is a lineage-specific expansion of surface proteins with mucin-like S/T stretches, which are likely to be primary targets of the second arm of the glycosylation system (Stwora-Wojczyk et al., 2004; Anantharaman et al., 2007). Given the gut parasitism of these apicomplexans, it is possible that these glycosylated mucin-like proteins helped homotypic interactions with the gut mucosa, which is also enriched in surface mucus (McGuckin et al., 2011). However, emergence of vertebrate blood parasitism in haemosporidians and piroplasms probably rendered these useless, and perhaps even maladaptive due to the immune response directed against them, thereby favoring their loss.

Thus, apicomplexan genomics suggests that not just adhesion domains of surface proteins, but also entire modification pathways for them were acquired on account of lateral gene flow from their hosts. It appears likely that gene transfer from the host facilitated by the initial parasitic contact allowed the development of elaborate host interaction proteins that might have been central to the emergence of the intimate endoparasitism observed in apicomplexans.

COMMON MOLECULAR ADAPTATIONS OBSERVED IN INTER-ORGANISMAL, INTER-GENOMIC AND INTRA-GENOMIC CONFLICTS

In contrast to the above-discussed example, where a unique set of adaptations emerged due to lateral transfer in course of an evolving host-parasite conflict, several other molecular adaptations appear to be common across a wide-range of biological conflicts. These commonalities appear to be a consequence of two disparate forces: (1) Convergent evolution due to strong selection for particular types of molecular interactions in conflicts; (2) Rapid dispersion over wide phylogenetic distances of certain highly effective adaptations by lateral transfer. We briefly outline some of these adaptations below.

Deployment of proteinaceous toxins

Proteinaceous toxins are the mainstay across all major levels of biological conflict. Such toxins are seen in competition between multicellular eukaryotes (e.g., castor bean ricin, Aspergillus sarcin and various snake venom proteins) and between them and their pathogens (e.g., anti-microbial peptide toxins and defensive RNases such as RNase A and RNase L)(Rochat and Martin-Eauclaire, 2000; Rosenberg, 2008; Wiesner and Vilinski, 2010). Conversely, such toxins are also used by pathogenic and symbiotic bacteria directed against their hosts (e.g., the cholera toxin and the shiga toxin) (Aepfelbacher et al., 2000; Alouf and Popoff, 2006). Similarly, the importance of protein toxins is becoming apparent in inter-bacterial conflicts (Schwarz et al., 2010; Russell et al., 2011; Iyer et al., 2011b; Zhang et al., 2011). In this regard, an exciting recent discovery has been made of a highly prevalent system of secreted multi-domain toxins, primarily involved in intra-specific conflict between related strains of prokaryotes (Aoki et al., 2011; Iyer et al., 2011b; Zhang et al., 2011). These proteins are typified by the tendency to vary their C-terminal toxin domains through a process of recombination that replaces an existing toxin domain by a distinct one encoded by standalone cassettes, while retaining the rest of the protein’s architecture (i.e., N-terminal regions related to trafficking and presentation) intact (Zhang et al., 2011). Hence, these toxins are termed polymorphic toxins. They include contact-dependent versions, which have long N-terminal stalks comprised of RSH/TV or filamentous haemagglutinin repeats that present the C-terminal toxin domain at the tip, shorter diffusible versions, and versions injected or delivered via type VI and ESX/type VII secretory systems (Aoki et al., 2011; Iyer et al., 2011b; Zhang et al., 2011). Importantly, they share these delivery/presentation mechanisms with those toxins using conflicts with hosts (Schwarz et al., 2010). However, they are distinguished from them by the presence a specific immunity protein encoded by a gene downstream of the toxin gene (Aoki et al., 2011; Zhang et al., 2011). Given their role in intra-specific conflict, they are an important determinant of kin-recognition and thereby have an effect on the included fitness in prokaryotes. Inter-genomic conflicts between cellular genomes and selfish replicons residing in the same cell (e.g., classical bacteriocins and plasmid addiction toxins) and intra-genomic conflicts between selfish elements and the host genome (restriction-modification (R-M) systems and genomic toxin-antitoxin (TA) systems) also use protein toxins with related domains (Cascales et al., 2007;
Zhang et al., 2011). The protein toxins of TA systems enable them to act as selfish elements that favor their own retention or “addiction” by killing cells where they are lost or disrupted. However, they might also enhance the fitness of their prokaryotic host. Indeed, expression of chromosomally embedded TA systems has been observed in diverse pathogens such as Mycobacterium tuberculosis and Brucella abortus when they are replicating within human cells. Here, the action of the toxin actually helps the bacteria to persist effectively in the hosts (Korch et al., 2009; Heaton et al., 2012).

There are some frequently recurrent themes in these toxins deployed across different levels of biological conflict: Most prominent are enzymatic toxins that disrupt the flow of biological information—nucleases targeting genomic DNA, tRNAs and rRNAs, nucleic acid base glycosylases, nucleic acid-modifying enzymes such as deaminases, peptidases that cleave key protein targets, and protein-modifying enzymes such as ADP-ribosyltransferases and AMP/UMPylating enzymes that alter the properties of proteins, such as components of the signaling and translation apparatus (Anantharaman and Aravind, 2003; Cascales et al., 2007; Leplae et al., 2011; Zhang et al., 2011). For example, toxins with the restriction endonuclease (REase) or HNH/ENDOVI folds are seen in intra-specific, inter-specific, inter-genomic (i.e., plasmid-encoded colicins) and intra-genomic conflicts (Stoddard, 2005; Cascales et al., 2007; Zhao et al., 2007; Zhang et al., 2011). Alternatively they disrupt cellular integrity by forming pores in cellular membranes (Gilbert, 2002). The enzymatic domains deployed in these conflicts are characterized by rapid sequence and structure divergence due to selection arising from immunity proteins and resistance against them.

Use of small molecule toxins

Deployment of small molecule toxins or antibiotics, synthesized via dedicated secondary metabolism pathways, is another common strategy, primarily observed in inter-organismal conflicts (Walsh, 2003). They are particularly common in bacteria, and in certain eukaryotic clades, such as fungi and plants. Several distinct types of such molecules are synthesized, with aminoglycosidic, fatty-acid-based (polyketide) and peptide-based skeletons being prevalent (Walsh, 2003). These basic skeletons, which are often synthesized by large multi-domain or multi-protein complexes catalyzing one or more rounds of endoergic condensations of acyl moieties or amino acids, are typically subject to a wide variety of modifications enzymes such as 2-oxoglutarate-dependent hydroxylases, methylases and oxidoreductases (Walsh, 2003; Iyer et al., 2009). Related to antibiotics are siderophores that are secreted for chelation of essential environmental metals (Barry and Challis, 2009). While not being toxic, they are the center of inter-organismal conflict because several bacteria have evolved receptors for uptake of “non-self” siderophores that allow them to benefit from siderophores produced by other organisms in the environment (Lee et al., 2012). Organisms combat such siderophore-stealing by diversifying their siderophores through modifications similar to those of antibiotics (Samel et al., 2008). Similar pressures also apply to small molecule signals that are used, especially by bacteria, to communicate with each other, as they can also be potentially exploited by non-kin organisms (Brady et al., 2004). Thus, the related secondary metabolism pathways for antibiotic, signaling molecule and siderophore biosynthesis are under pressure for rapid diversification due to pressures from resistance and stealing. In most bacteria, components of these secondary metabolism pathways are encoded by multi-gene operons, which, as indicated by the large number of dioxygenases and oxidoreductases encoded by them, appear to have radiated concomitant with the first oxygenation event in Earth’s history (Iyer et al., 2010). Subsequently, they appear to have undergone diversification through recruitment of multiple non-ribosomal peptide ligases and acyl condensation enzymes, sequence divergence of individual enzymatic components, and recombination between distinct biosynthetic operons to synthesize new products (Walsh, 2003; Samel et al., 2008; Iyer et al., 2009, 2010).

Enzymes that facilitate mobility and replication of selfish elements

The fitness of intra-genomic and intra-cellular selfish elements depends on a variety of enzymes that allow their efficient propagation. One group of these enzymes is directly involved in the replication and transcription of the selfish DNA and provides autonomy from the host replication and transcription systems (Galun, 2003; Burt and Trivers, 2006). These enzymes include DNA polymerases, RNA polymerases, primases and reverse transcriptases, which in certain cases are distantly related to the cellular counterparts and in other cases, represent distinct, non-homologous enzymes with analogous activities. These enzymes often face selective pressures for diversification due to exploitation by defective or satellite element which lack their own replication or due to host defensive mechanisms (Galun, 2003; Burt and Trivers, 2006). Another widely used group of enzymes that do not directly catalyze nucleic acid synthesis are transposases/integrases, which often display nuclease domains related to the nuclease domain of toxins (see above) (Lilley and White, 2000; Galun, 2003; Stoddard, 2005; Burt and Trivers, 2006; Zhao et al., 2007; Mak et al., 2010; Zhang et al., 2011). One frequently encountered catalytic domain across a wide-range of transposons is a transposase/integrate domain of the RNaseH fold which is related to the nuclease domain found in the archaeal NurA and the argonaute nucleases (Aravind et al., 2000; Nowotny, 2009). This suggests that several of these mobile elements share an ultimate common ancestry in the form of an ancient RNaseH integrase. Additionally, these enzymes from selfish elements are characterized by a mélangé of structurally distinct DNA-binding domains (DBDs), which diversify considerably due to pressures for specific recognition of sequences in the selfish elements (Babu et al., 2006).

Immunity systems

Antagonistic actions in biological conflicts are countered by a variety of dedicated immunity mechanisms, which act over and beyond the immunity gained via sequence divergence of targeted proteins. The polymorphic toxins, plasmid-borne bacteriocins, TA, and R-M systems are all characterized by the presence of an antidote or immunity protein that neutralizes the toxin produced by them (Kobayashi, 2001; Leplae et al., 2011; Russell et al.,
COMMONALITIES IN THE MULTIPRONGED APPROACH OF INTRA-CELLULAR BACTERIA AND VIRUSES IN MANIPULATING EUKARYOTIC HOSTS

Endosymbiotic/parasitic bacteria utilize a multipronged approach by often simultaneously deploying several toxins or effectors, each with its own mode of action to manipulate the behavior of the eukaryotic hosts in which they reside. Yet genomics of these bacteria suggests that there is a relatively small set of strategies that are exploited by intra-cellular bacteria from across the bacterial tree, including representatives of alphaproteobacteria, gammaproteobacteria, chlamydiae, and bacteroidetes (Collingro et al., 2005; Ogata et al., 2006; Penz et al., 2010; Schmitz-Esser et al., 2010; Georgiades et al., 2011). The most commonly used approach is the deployment of proteins that alter action of the ubiquitin system, including E3-ligases with RING, U-Box and F-Box domains, deubiquitinating and desumoylating peptidases, especially of the PROTAC/ITCH type (Pancer and Cooper, 2006). In some cases, LRR and other domains might be combined with the SCF-type ubiquitin E3-ligases to allow degradation of proteins encoded by invasive replicons or cells (Thomas, 2006). In the vertebrate lineage, on two independent occasions, elaborate mechanisms involving mutagenesis and recombination have evolved to enable diversification of pathogen receptors, which respectively, utilize the immunoglobulin domain and LRRs (Pancer and Cooper, 2006).

EUNATION OF MAJOR EUKARYOTIC SYSTEMS: CONTRIBUTION FROM PROTEINS DEPLOYED IN INTER-ORGANISMAL, INTER-GENOMIC AND INTRA-GENOMIC CONFLICTS

In this section of the article we discuss with examples as to how several of the above-discussed players deployed in biological conflicts have played a major role in the emergence and elaboration of various eukaryotic adaptations. In doing so we take examples both from early events close to eukaryogenesis and also systems that evolved in particular eukaryotic lineages, such as metazoans.
Emergence of key players in eukaryotic chromatin protein complexes

Eukaryotes are distinguished from the two prokaryotic superkingdoms by their dynamic chromatin organized by histones with low complexity tails, which provides a veritable "ecosystem" for several protein-modifying and ATP-dependent remodelers (Allis et al., 2006; Kouzarides, 2007; Aravind et al., 2011; Iyer et al., 2011a). The mysterious origins of several of the unique components of eukaryotic chromatin have begun to considerably clear up with recent genomic data. SWI2/SNF2 ATPases, which had at least six representatives by the time of the last eukaryotic common ancestor (LECA), had already diversified to perform several distinct chromatin remodeling activities, such as sliding/ejection of nucleosomes, exchange of canonical nucleosomes with those containing alternative histones, or altering nucleosomal spacing (Iyer et al., 2008b; Hauk and Bowman, 2011; Hota and Bartholomew, 2011). Phylogenetic, domain architecture, and gene neighborhood analysis revealed that SWI2/SNF2 ATPases are superfamily II DNA helicases, which had their most extensive diversification as part of R-M systems and related systems that are likely to function as a defensive mechanism against bacteriophages (related to the phage growth limitation or Pgl system) (Iyer et al., 2008b) (Figure 3). In phylogenetic trees, the eukaryotic versions are nested within the radiation of SWI2/SNF2 ATPases from prokaryotic selfish elements and were transferred on at least three independent occasions to eukaryotes (Figure 3A). The first of these transfers occurred prior to the LECA, and by the time of the LECA had proliferated to spawn at least six distinct lineages (Iyer et al., 2008b). The remaining two transfers occurred much later in eukaryotic evolution, and gave rise to the Strawberry Notch and HARP-like SWI2/SNF2 ATPases (Iyer et al., 2008b). Bacterial R-M systems contributed a second ATP-dependent chromatin remodeling enzyme to eukaryotes, the MORC ATPase, which contains a composite module comprised of gyrase, histidine kinase, and MutL (GHKL) and S5 domains (Iyer et al., 2008a). Analysis of R-M bacterial systems showed that they display a vast radiation of several different types of GHKL-S5 module ATPases, of which the MORCs form
Eukaryotic SWI2/SNF2

REase-fused

SWIM-fused/associated

N6A-Methylase fused

HKD phosphoesterase fused

HARP/SMARCAL1

Bacterial HARP-like

HepA-like

Eukaryotic Sno

Bacterial Sno

VRR-NUC associated

FIGURE 3 | Evolutionary relationships of various families of enzymes illustrating the origin of eukaryotic versions within radiations of systems involved in inter- and intra-genomic conflicts. Reconstructed phylogenetic trees are shown for (A) The bacterial radiation of the SWI2/SNF2 ATPases. (B) MORC-like ATPases and (C) The Double-psi beta barrel containing RNA polymerases. Certain clades with multiple families such as the eukaryotic SWI2/SNF2 ATPases, the Topoisomerase ATPase subunits, the cellular DDRP and eukaryotic RdRPs are collapsed into triangles for clarity. Illustrative domain architectures or gene neighborhoods are shown next to the leaf. Genes in gene neighborhoods are shown in block arrows with the arrow head pointing from the 5' to the 3' gene. Proteins and gene neighborhoods are labeled by the gene name and species name separated by underscores. The trees represent only the overall topology because they were obtained by a combination of conventional phylogenetic tree construction and structure-based determination of higher-order relationships.
one distinct clade (Figure 3B). Given that basal excavate lineages, such as parabasalids and diplomonads lack MORCs, they appear to have been acquired by eukaryotes post-LECA, prior to the radiation of the large eukaryotic clade uniting animals, fungi, amoebzoans, and plants (Iyer et al., 2008a) (Figure 4). Both the MORCs and the SWI2/SNF2 ATPases use ATP hydrolysis to catalyze DNA-unwinding or large-scale looping of DNA in aiding the restriction activity of the REases. This activity has been
reused in a biochemically comparable, but functionally distinct, context to remodel protein-DNA contacts or facilitate higher-order looping in eukaryotic chromatin. In a similar vein, R-M systems might also account for the origin of the eukaryotic phosphoesterase enzyme TDP1, which hydrolyzes 3’-phosphotyrosyl bonds between DNA and the active tyrosine of topoisomerase Iib to release DNA from topoisomerase adducts (Gajewski et al., 2012). Sequence relationships of TDP1 suggest that it is likely to have been derived from HKD phosphoesterase domains found fused to SWI2/SNF2 ATPases in bacterial R-M systems (Iyer et al., 2006a).

Similar studies have shown that the DNA methylases of eukaryotes, which play an important role as encoders of epigenetic information that goes over and beyond the basic genetic information, also largely owe their origin to R-M systems and related methylation systems that protect prokaryotic genomes against restriction attacks by selfish R-M systems (Bestor, 1990; Iyer et al., 2011a). Both DNA cytosine (C5) and adenine (N6) methylases of eukaryotes appear to have been derived from bacterial R-M system and dcm methylases on more than 10 independent occasions (Iyer et al., 2011a). As none of the conserved eukaryotic lineages of DNA methylases can be detected in the parabasalids and diplomonads, it appears that the classical epigenetic DNA modification of cytosine was absent in the LECA. The primary conserved cytosine DNA methylase of eukaryotes, DNMT1, appears to have emerged only just before the time the heterolobosean-kinetoplastid clade branched off from the remaining eukaryotes, and phylogenetic analysis strongly supports its origin from a bacterial R-M system methylase-related to M.NgoFVII (Iyer et al., 2011a). Most other DNA methylases of eukaryotes can be attributed to comparable later acquisitions, primarily from other types of R-M systems. Recent discoveries have indicated that the reversal of cytosine DNA methylation in several eukaryotic lineages occurs via the action of Tet-JBP family of 2-oxoglutarate and iron-dependent dioxygenases (2OGFeDOs), which remove the methyl group through oxidative conversion to hydroxymethylcytosine and further oxidized cytosine derivatives that are then cleared by base excision repair (He et al., 2011; Iyer et al., 2011a). Interestingly, related enzymes, JBP1/2, catalyze the hydroxylation of thymine in the synthesis of base J, an epigenetic modification observed in kinetoplastids (Vainio et al., 2009). Prior studies on the evolution of 2OGFeDOs revealed that the eukaryotic Tet-JBP enzymes were derived from precursors encoded by caudate bacteriophages (Iyer et al., 2011a). Bacteriophages have been known to display a rich variety of DNA modifications, including hydroxymethylated pyrimidines, which enable them to evade restriction by different R-M systems in the host genome (Gommers-Ampt and Borst, 1995). Thus, the bacteriophage Tet-JBP enzymes appear to have first emerged as part of their counter-restriction strategy, and subsequently recruited to generating and erasing epigenetic marks on DNA upon being transferred to eukaryotes. Multiple studies have also revealed that not just enzymatic domains, but also specific DBDs found in eukaryotic chromatin proteins might have been acquired from bacterial R-M systems and replication apparatus of caudate bacteriophages. The SAD/SRA domain, which is a key player in eukaryotic chromatin as an epigenetic “reader” of hemimethylated cytosine marks, has been derived from the DNA-binding domain of REases from R-M systems that discriminate between hemimethylated and fully methylated sites (Iyer et al., 2011a). Likewise, the recently described HARE-HTH domain, which might have an important role in discriminating the DNA modification generated by the cytosine methylases, and the Tet/JBP enzymes has also evolved from bacterial R-M systems, where it is combined with several distinct REase domains (Aravind and Iyer, 2012). On the other hand, another DNA-binding domain, the HIRAN domain, which among other proteins is associated with the eukaryotic chromatin remodeling RAD5-type SWI2/SNF2 ATPases appears to have emerged from the replication apparatus of caudate bacteriophages (Iyer et al., 2006a).

In stark contrast to chromatin remodeling and epigenetic DNA modifications, enzymes catalyzing epigenetic modifications of proteins in eukaryotic chromatin appear to have extensively drawn from very different types of prokaryotic systems involved in inter-organismal conflict. Two key epigenetic modifications are acetylation of lysines and methylation of both lysines and arginines in histones and other proteins in eukaryotic chromatin (Allis et al., 2006; Kouzarides, 2007). Sequence comparisons show that the eukaryotic arginine methylases (PRMT) have been derived from within a bacterial radiation of peptide methylases (Aravind et al., 2011). The closest bacterial sister groups of the eukaryotic PRMTs are encoded in antibiotic-like secondary metabolite biosynthesis operons that also contain genes for peptide dioxygenases, non-ribosomal peptide synthetases and other peptide-oxidizing enzymes such as LSD1-related amine oxidases (Aravind et al., 2011). Bacterial PRMT domains are also incorporated as domains of gigantic antibiotic biosynthesis enzymes, such as anabaenopeptidil synthetase that synthesizes a peptide toxin of the cyanobacterium *Anabaena* (Rouhiainen et al., 2000; Aravind et al., 2011). Interestingly, the LSD1-like amine oxidases observed in these and other peptide antibiotic/toxin biosynthesis operons are also the precursors of eukaryotic histone demethylases that catalyze oxidative removal of methyl groups from mono- and di-methylated histone H3K4 (Allis et al., 2006; Kouzarides, 2007). All the remaining histone demethylases in eukaryotes belong to one large superfamily of 2-oxoglutarate-dependent dioxygenases known as the Jumonji-related dioxygenases (Iyer et al., 2010). These, along with LSD1, are absent in the earliest-branching eukaryotes such as parabasalids and diplomonads, and first appear as multiple paralogous copies just prior to the divergence of the heterolobosean-kinetoplastid clade from the other eukaryotes (Iyer et al., 2010). However, each of these multiple eukaryotic paralogous lineages have their own bacterial counterparts suggesting that they had already diverged in bacteria before being acquired. In bacteria, like LSD1, they appear in one or more copies in peptide antibiotic/toxin biosynthesis operons (Iyer et al., 2010), where they are likely to catalyze multiple oxidative modifications of peptides as previously observed in the biosynthesis of penicillin and its derivatives (Liras and Demain, 2009). Thus, it is plausible that eukaryotes have acquired multiple paralogous jumonji-related dioxygenases via the transfer of a single secondary metabolism gene-cluster with multiple versions of these enzymes. In eukaryotes, other than
histone demethylation, they also radiated to give rise to enzymes catalyzing the last step in the generation of the eukaryote-specific rRNA\(^{\Phi}\) modification, hydroxyxynutosine, and protein asparagine hydroxylation (Iyer et al., 2010). In contrast to these, the histone H3K79 methylase Dot1 appears to have emerged from a methylase effector delivered by intra-cellular symbionts and is seen in diverse bacterial endo-symbionts/pathogens of amoeboid protozoans and metazoans, like Parachlamydia and Legionella (Aravind et al., 2011).

Thus, components from R-M and virus-restriction systems, viral replication apparatus, peptide antibiotic/siderophore biosynthesis systems and effectors of intra-cellular bacteria, which are exemplars of intra-genomic, inter-genomic and inter-organismal conflict systems, have been harnessed as progenitors of distinguishing components of eukaryotic chromatin.

**Conflict systems and eukaryotic RNA metabolism**

Eukaryotes are characterized by the unique RNAi system, which is typified by small RNAs (usually 23–35 nt in length) that perform a number of roles ranging from post-transcription gene regulation to regulation of chromatin structure (Allis et al., 2006; Grewal, 2010). Of these small RNAs, the siRNA-type RNAs are particularly important in gene-silencing, and might be amplified by a distinctive enzyme of this system, the RNA-dependent RNA-polymerase (RdRP), which can be traced back to the LECA (Salgado et al., 2006; Ruprich-Robert and Thuriaux, 2010; Iyer and Aravind, 2011). Sequence-structure analysis of the RdRPs revealed that its two catalytic double-\(\beta\)-\(\beta\)-barrel (DPBB) domains are related to the catalytic domain found in the two largest subunits of the cellular RNA polymerases from all life forms (Salgado et al., 2006; Ruprich-Robert and Thuriaux, 2010; Iyer and Aravind, 2011). The search for RdRP cognates outside eukaryotes showed that they are prevalent in certain bacteriophages of firmicutes and also a variety of recently identified novel selfish elements in bacterial genomes (Figure 3C) (Iyer and Aravind, 2011). In these potential selfish elements they are often encoded alongside genes for different DNase domains such as those belonging to the RNase and URI endonuclease fold, which might aid in the mobility of the elements (Figure 3C). The RdRPs might also be combined with RNase H domain in the cyanobacterial versions suggesting that might function in the context of RNA-DNA hybrids (Iyer and Aravind, 2011).

Furthermore, structural analysis of the RNA-polymerases with DPBB catalytic domains showed that the RdRP-like enzymes belonged to a radiation of single-subunit RNA polymerases encoded by variety of selfish elements, from within which the cellular multi-subunit versions emerged via fission of the two catalytic domain-containing segments of the single-subunit enzyme (Figure 3C). It appears plausible that these RdRP-like enzymes of intra-genomic selfish elements and bacteriophages primarily arose as enzymes that aided their mobility by potentially acting as primases enabling their replication (Iyer and Aravind, 2011). Upon acquisition by the eukaryotic lineage, prior to the LECA, the enzyme appears to have been recruited as a part of the RNAi systems for amplification of small RNAs. Interestingly, the RdRP is not the only nucleic acid polymerase that has been recruited to RNA metabolism from a prokaryotic selfish element. Recent studies on the domain architectures and sequence relationships of the most conserved splicing factor of eukaryotes Prp8, which is part of the spliceosomal catalytic center, has revealed that it has been derived from the polyprotein of a retroelement replete with the reverse transcriptase, “thumb” and RNaseH domains (Dlakic and Mushegian, 2011). However, in Prp8 the active site of the reverse transcriptase domain is disrupted, suggesting that it merely functions in a nucleic acid-binding capacity rather than as an active enzyme (Dlakic and Mushegian, 2011). It is conceivable that this retroelement was associated with the ancestral group-II introns that invaded the genome in the pre-LUCA period to give rise to the spliceosomal introns of eukaryotes.

On several occasions, components of yet another prokaryotic inter-organismal conflict system, namely the recently characterized polymorphic toxin systems, appear to have contributed to eukaryotic RNA-processing and modification systems (Zhang et al., 2011). In eukaryotes, small nucleolar RNAs (snRNAs) are required for modification and maturation of rRNA in the nucleolus. In several eukaryotes certain snRNA, like U16 and U86, are directly released from the introns encoding them by the endonucleolytic action of the endoU RNase (Laneve et al., 2003). Sequence and structure analysis revealed that the endoU RNase of eukaryotes is nested within a vast radiation of RNase domains that function as toxins in bacterial polymorphic toxin and related secreted toxin systems (Zhang et al., 2011). Thus, acquisition of the endoU domain appears to have enabled eukaryotes to bypass splicing to directly release snRNAs from introns. RNA-editing via deamination of cytosine and adenine has considerably expanded in eukaryotes and is observed not just in tRNAs but also in mRNAs and as part of a counter-viral strategy (Iyer et al., 2011b). The origins of certain divergent metal-dependent nucleic acid deaminase domains, such as those of the AID-APOBEC clade and the DYW clade, which catalyzes massive RNA-editing in plant chloroplasts and mitochondria, were rather unclear until recently (Zehrmann et al., 2011). Analysis of the polymorphic toxins revealed that one of the widely used toxin domains was the nucleic acid deaminase that had greatly diversified in such and related secreted toxins (Iyer et al., 2011b). Importantly, the origin of the both the DYW and AID-APOBEC-like deaminases could be placed within specific prokaryotic toxin groups (see below for details).

**Prokaryotic conflict systems and protein-modifying enzyme and second messenger in eukaryotic signaling systems**

Recent studies on the diversity of catalytic toxin domains deployed in bacterial polymorphic and related secreted toxins systems are also throwing light on the emergence of what were previously considered uniquely eukaryotic signaling systems (Figure 4). One such is the polyADP-ribosylation system, which modifies aspartate, glutamate and lysine side chains in both cytoplasmic and nuclear proteins including histones, with profound effects on DNA repair, chromatin organization, telomere dynamics, centrosomal and mitotic spindle organization, and endosomal trafficking (Ame et al., 2004). The enzymes catalyzing this modification, polyADP-ribosyl polymerases (PARPs), can be traced back to the LECA, but their emergence in eukaryotes remained a mystery (Citarella et al., 2010). The closest relatives of the PARPs are
found among toxin domains of a toxin used in inter-bacterial conflicts delivered via a distinctive phage-derived, injecting secretory system known as the Photorhabdus virulence cassette (Hurst et al., 2004; Zhang et al., 2012). Related PARP domains are also found as effectors of intra-cellular symbionts/parasites of amoebae and metazoa such as Legionella dancourtii. Recently, a novel family of ADP-ribosyltransferases (ARTs), distinct from the PARPs, was identified, and typified by the Neur14 protein of humans (De Souza and Aravind, 2012). These ARTs might have an important role in the organization of the eukaryotic centrosome among other processes. They also seem to have been derived from effectors delivered by endoparasitic bacteria, such as Waddlia (Hurst et al., 2004). The use of mono-ADP-ribosyltransferases by diverse bacteria as toxins in intra- and inter-specific conflicts (i.e., polymorphic toxins) and those directed at host proteins is well-known (Koch-Nolte et al., 2008; Laing et al., 2011; De Souza and Aravind, 2012). Indeed, other than the PARPs and Neur14-like ARTs, the eukaryotes also possess several mono-ARTs which are nested within the radiation of bacterial toxin ARTs. Thus, on more than three occasions eukaryotes appear to have recruited the toxin ART/PARP domains as protein-modifying enzymes, with the event giving rise to the PARPs probably happening before the LECA (Figure 4). While in bacteria these enzymes appear to largely function as toxins, in eukaryotes they appear to have been utilized to post-translationally modify proteins and provide an additional level of coding information (Koch-Nolte et al., 2008; Laing et al., 2011). Beyond the events spawning pathways that are widespread in eukaryotes, polymorphic and related toxin systems also appear to have contributed to the origin of signaling systems unique to certain lineages, such as metazoa. In addition to ARTs, other bacterial toxin domains utilizing NAD as a substrate have also been recruited to metazoan signaling. The ADP-ribosyl cyclase domain was previously observed only in animals (in the CD38 and CD157 proteins) and generates two messenger molecules, namely cyclic ADP ribose (cADPr) and nicotinic acid adenine dinucleotide phosphate (NAADP), respectively, from NAD and NADP (Guse and Lee, 2008). The former two nucleotides function as messenger molecules that induce calcium signaling pathways via the ryanodine receptors (Guse and Lee, 2008). The discovery of the ADP-ribosyl cyclase as a toxin domain in bacterial polymorphic toxins provides a potential explanation for the sudden origin of this signaling enzyme in animals (Zhang et al., 2012). Additionally, fungi too appear to have independently acquired this domain from bacteria, suggesting that it might have been recruited on more than one occasion in eukaryotic evolution (Zhang et al., 2012).

The Teneurin/Odd Oz proteins found in metazoa and choanoflagellates function as developmental regulators with a potential role in cell-surface adhesion in diverse processes such as cell migration, neuronal path finding and fasciculation, gonad development, and basement membrane integrity (Minet et al., 1999; Silva et al., 2011). These proteins appear to have been derived from a complete bacterial polymorphic toxin, with both the N-terminal RHS/YD repeats, which form a stalk and the C-terminal toxin domain that is a derived version of the HNH/EndoVII fold (Zhang et al., 2012). While the C-terminal toxin domain has lost its active site residues in the animal lineages, it is cleaved and secreted as a potential neuromodulator (Qian et al., 2004). On the other hand the N-terminal RHS repeats appear to play a role in adhesion between different Teneurin/Odd molecules, which is a key aspect of their cell-cell signaling function (Silva et al., 2011). Other than the toxin domains, certain other domains in eukaryotic signaling pathways have also been acquired from bacterial polymorphic toxin systems. The hedgehog signaling pathway is a eukaryotic signaling pathway initiated by the hedgehog proteins, which undergo autoproteolytic cleavage to release signaling messengers (Ingham et al., 2011). The HINT domain, which catalyzes this autoproteolytic cleavage in the eukaryotic hedgehog proteins, is likely to have been derived from the HINT domains commonly found in bacterial polymorphic toxins, where they apparently facilitate the autoproteolytic release of the C-terminal toxin domain into target cells (Zhang et al., 2011). In metazoa, hedgehog activates a down-stream signaling cascade in target cells to activate the transcription factor Gli (Ingham et al., 2011). The Suppressor of Fused (SuFu) protein tethers the Gli in the cytoplasm in the absence of the hedgehog signal to prevent constitutive activation. This SuFu protein of the animal hedgehog pathway also has its origin in bacterial polymorphic toxin systems, where members of the SuFu superfamily function as immunity proteins that neutralize a structurally diverse range of toxin domains (Zhang et al., 2011).

**The eukaryotic ubiquitin system: origin and elaboration**

One of the most remarkable features of eukaryotes is the ubiquitin system, which comprises of several parallel enzymatic cascades which ligate Ubiquitin or an Ubl protein to target proteins, typically on a lysine residue (Hochstrasser, 2009). These cascades are typified by an E1 enzyme, which activates the Ub/Ubl terminal COOH group by adenylation and trans-thiolation to transfer it to and E2 enzyme. The E2 enzyme may then either directly or via an E3 enzyme transfer the Ub/Ubl to the target protein. In eukaryotes, such modifications often target proteins for degradation via the proteosomal system, where the Ub/Ubl is first cleaved off and released by a JAB domain metallopeptidase (Kerscher et al., 2006). In addition to proteosomal degradation, Ub/Ubl modifications also alter the interactions, localization and biochemistry of the target proteins and are modulated by a series of peptidases (DUBs) that debiquitinate them (Burrows and Johnston, 2012). Until recently it was thought that the Ub-system was a purely eukaryotic innovation. However, multiple studies have shown that the antecedents of the Ub-system first emerged in prokaryotes as part of a dramatic radiation of UbIs and E1-like enzyme in operons for the biosynthesis of cofactors (e.g. thiamin and molybdopterin), cysteine, and peptide secondary metabolites such as siderophores, antibiotics/toxins and small molecule signals (Burroughs et al., 2011, 2012). A subset of these operons is highly mobile (i.e., widespread dispersal across distant lineages) and evolved features characteristic of the eukaryotic Ub-systems, namely the presence of E2 and sometimes E3 enzymes and the debubquitinating JAB peptidase (Burroughs et al., 2011). The fact that these operons are mobile, and usually tend to couple the ubiquitinating enzymes with debubquitinating JAB peptidases, presents parallels to the R-M systems (Iyer et al., 2006c). Like
them these systems combine opposing actions in the modifying and de-modifying enzymes, and have no links to the metabolic enzymes that are typical of the operons with E1-like enzymes and Ubls that synthesize small molecule. Hence, we posit that these are potential selfish elements that act like the R-Ms, but at the protein level, by possibly destabilizing proteins through transfer of the Ub and restoring the original protein by removal of the Ub by the JAB peptidase. Since these Ub-like systems are closest to the eukaryotic versions, it is very likely that they were derived from them. On account of their mobility they are seen in several bacteria and certain archaea (e.g., the Caldiarchaeum) (Iyer et al., 2006c; Burroughs et al., 2011; Nunoura et al., 2011); hence, it is possible that eukaryotes might have acquired the precursor of their Ub-system either from their archaeal precursor or from endosymbiotic bacteria (Figure 4).

The only DUB that is consistently observed in prokaryotic Ub-like systems is the JAB peptidase domain (Iyer et al., 2006c; Burroughs et al., 2011). Eukaryotes, however, possess several other DUBs, most of which belong to the papain-like peptidase fold and a few to the Zincin-like metallopeptidase fold (Iyer et al., 2004). Interestingly, papain-like peptidases (e.g., Otu-like peptidase domain) and Zincin-like metallopeptidases are frequently found among the toxin domains of effectors delivered by a range of intra-cellular bacteria (Loueiro and Ploegh, 2006). These were previously thought to be lateral transfers from hosts to their endo-symbionts/parasites, which are used to interfere with the host Ub-system (Lomma et al., 2010; Schmitz-Esser et al., 2010). However, recent studies on polymorphic toxin systems suggest that such peptidase domains are far more widely distributed in bacterial toxins and often among toxins of free-living bacteria deployed in inter-bacterial conflicts (Zhang et al., 2012).

Hence, it seems more likely that they first emerged in bacteria as part of the polymorphic and related secreted toxin systems and were acquired by eukaryotes and recruited as DUBs in course of the development of the mitochondrial endosymbiosis (Figure 4). Not surprisingly, these DUB-like peptidase domains are common among intra-cellular bacteria such as Wolbachia, Rickettsia and Odyssella, which are closely related to the mitochrondrial precursors (Figure 2). Indeed, these DUBs probably originally emerged as part of the strategy utilized by these bacterial endosymbionts/pathogens that countered the immunity mechanism based on ubiquitination of target proteins. Interestingly, several of these papain-like DUB domains are also related to polyprotein-processing peptidases of eukaryotic RNA viruses and retroviruses (Iyer et al., 2004). It is conceivable that the emergence of the Ub-system in eukaryotes also provided a means for RNA viruses to escape constraints placed by the eukaryotic mRNA cap on internal translation initiation, by simply enabling translation of polypeptides that are then processed by the DUB peptidases. In course of viral evolution many of the DUB domains were probably incorporated into their own polyproteins to allow autoproteolytic processing.

**Executers of apoptosis: multiple independent recruitments of domains from prokaryotic conflict systems**

One of the simplest counter-pathogen strategies is regulated cell death or apoptosis, in which a cell might sacrifice its own fitness and prevent the pathogen from replicating within it. This typically works in situations where the inclusive fitness accrued from saving kin from infection might contribute to fixation of altruistic behaviors such as apoptosis (Aravind et al., 2009). Such mechanisms are likely to be further enhanced with the emergence of colonial or multicellular organization. Some of the simplest programmed cell death systems seen in bacteria are constituted by intra-genomic selfish elements. For example, in Escherichia coli a defective prophage produces a toxin known as Lit with a zincin-like metallopeptidase domain to cleave the elongation factor Tu and kill the cell when infected by the phage T4, thereby preventing the further spread of T4 to remaining cells in the colony (Snyder, 1995). Likewise, under conditions of starvation, when resources are limiting, chromosomally encoded toxin-antitoxin systems, such as the entericidin locus, mediate cell death in bacteria like E. coli and allow certain cells to survive and grow at the expense of kin that have undergone cell death (Bishop et al., 1998). Thus, the principle of the use of toxins as mediators of programmed cell death appears to be an ancient one (Jensen and Gerdes, 1995; Bishop et al., 1998).

Although eukaryotic lack conventional toxin-antitoxin systems, the executioners of apoptosis resemble the prokaryotic toxins from these and other conflict systems in that they cleave or modify specific target proteins or permeabilize membranes in the cell committed to apoptosis. These effectors have been best studied in the animal lineage and include membrane-permeability regulators (the BCL2 superfamily), DNA-cleaving enzymes (e.g., the DNA fragmentation factor/CIDE), DNA-modifying enzymes (e.g., pierisin) and peptidases (e.g., the caspases) (Chou et al., 1999; Lugovskoy et al., 1999; Kanazawa et al., 2001; Riedl and Salvesen, 2007). Investigation into the provenance of these proteins has revealed multiple ancient connections to bacterial toxin systems. The core helical domain of the BCL2 superfamily (the first 6 helices) is specifically related to the translocation (T) domain of several host-directed toxins from distantly related bacteria such as diphtheria, botulinum, tetanus and Vibrio toxins (Chou et al., 1999). The T-domain undergoes a pH induced conformational change to assume a BCL2-like structure, inserts into the endosomal membrane and transfers the catalytic domain of the toxin into host cytoplasm. Given its sudden emergence in metazoans, it is likely that it was derived from a bacterial toxin and recruited as regulator of the permeability of the mitochondrial membrane. In metazoans these domains diversified into anti-apoptotic versions, which prevent the release of cytochrome C from mitochondria and pro-apoptotic versions which foster its release (Chou et al., 1999; Riedl and Salvesen, 2007). From animals, the BCL2 superfamily was secondarily acquired by large DNA viruses that infect animals, and include membrane-permeability regulators (the BCL2 superfamily), DNA-cleaving enzymes (e.g., the DNA fragmentation factor/CIDE), DNA-modifying enzymes (e.g., pierisin) and peptidases (e.g., the caspases) (Chou et al., 1999; Lugovskoy et al., 1999; Kanazawa et al., 2001; Riedl and Salvesen, 2007).
Salvesen, 2007). Structural studies had revealed that this domain contains an endonuclease domain of the HNH/EndoVII fold, but its origins remained unclear (Lugovskoy et al., 1999). Recent analysis of the bacterial polymorphic toxins revealed that a subset of them contains a toxin nuclease domain, which shares unique sequence signatures with the DFF/CIDE endonuclease domain to the exclusion of other representatives of HNH/EndoVII fold (Zhang et al., 2012). Here again, the relative abundance of the HNH/EndoVII fold among polymorphic and related toxin domains compared to its lone presence in DFF/CIDE, which is restricted to metazoans, points to an origin for the latter from a representative in the bacterial toxin systems. Pierisin-type ARTs are unusual enzymes that mediate apoptosis (thus far only known from lepidopterans) by ADP-ribosylating the N2 atom of guanine in DNA (Kanazawa et al., 2001). The lepidopteran pierisin-like ARTs are specifically related to the ART toxin domains found in certain bacterial polymorphic toxins and insecticidal toxins of insect pathogens, such as Bacillus sphaericus (Orth et al., 2011). This suggests that they were probably laterally transferred into lepidopterans from a bacterial symbiont or parasite, followed by their reuse as an apoptotic effector. In all the above examples the natural action of the bacterial toxins in disrupting or killing animal cells appears to have been harnessed as a mechanism to execute apoptosis.

Caspase-like peptidases are the central executers of apoptosis throughout eukaryotes and have been demonstrated to play a central role in cell death in animals, fungi, plants, and certain other eukaryotes (Aravind and Koonin, 2002; Riedl and Salvesen, 2007). Prior evolutionary analysis of the caspase-like superfamily revealed that they first diversified in bacteria into several clades such as the metacaspases, paracaspases and numerous other bacteria-specific lineages (Aravind and Koonin, 2002). Metacaspases were transferred to eukaryotes prior the LECA and are found in most eukaryotes (Figure 4). Subsequently, in the animal lineage and in dictyostelid slime molds metacaspases were displaced by a second acquisition from bacteria, the paracaspases, which then radiated in animal to give rise to the classical caspases (Aravind and Koonin, 2002). This phyletic pattern suggests that paracaspases are effectively functionally comparable to metacaspases, as they have displaced them on more than one occasion. Interestingly, several bacteria, particularly endosymbiotic/parasitic alphaproteobacteria (e.g., Agrobacterium, Labrenzia, Bradyrhizobium) encode metacaspases and paracaspases with N-terminal signal peptides that are likely to be secreted into their hosts (Aravind and Koonin, 2002). Hence, these peptidases were possibly first used in regulating endoparasite/symbiont-host conflicts to modulate the immune response and cell death in favor of the intra-cellular bacterium. Consistent with this, recent studies in humans have shown that the paracaspase modulates the T-cell-dependent immune response by cleaving A20, a deubiquitinating enzyme involved in the process, and is required for prevention of cell death in diffuse large B cell lymphoma (Coornaert et al., 2008; Ferch et al., 2009). This suggests that caspase-like peptidases might have been acquired on multiple occasions in eukaryotic evolution from endosymbiotic bacteria, which were probably utilizing them to regulate the survival of their host cells. On a similar note, the GIMAP/AIG1-like GTPases, which are deployed by certain endo-symbionts/parasites (Figure 2), could have given rise to the eukaryote representatives of this clade which are known to modulate both apoptosis and the immune response.

Thus, protein domains that originally diversified in prokaryotic conflict systems both as toxin and also as potential modulators of host defensive responses have had a notable effect on the evolution of apoptosis.

**Origin of antigen receptor diversification mechanisms and mutagenic immunity mechanisms**

Despite the enormous disparities in the immune systems of different eukaryotes, there are a few common strategies that are observed across most of them. These include the use of a relatively small number of families of protein domains as antigen receptors. Diversification of antigen receptors in most eukaryotes is a passive process of sequence divergence, probably under positive selection, within families of lineage-specifically expanded proteins (e.g., LRR proteins). However, in both jawed and jawless vertebrates two distinct and directed mechanisms for their diversification have been observed, namely recombination and active mutagenesis, which result in different populations of lymphocytes expressing different types of antigen receptors (Pancer and Cooper, 2006; Schatz and Swanson, 2011). In both jawed and jawless vertebrates the process of directed mutagenesis by DNA cytosine deaminases of the AID-APOBEC superfamily is utilized (Rogozin et al., 2007). Such mutagenesis is used either as a trigger for antigen gene-conversion, or for hypermutation or for antibody class-switching. Additionally, certain representatives of the AID-APOBEC family of cytosine deaminases are also major line of defense against retroviruses by mutagenizing their genomes by cytosine deamination (Chiu and Greene, 2006). Although AID-APOBEC-like deaminases were, until recently, thought to be restricted to vertebrates, sensitive sequence analysis showed that more divergent members exist in nematodes, cnidarians and several distantly related algal groups. Identification of these sequences helped establish that the fast-evolving AID-APOBEC deaminases have their ultimate origin in the toxin domains of polymorphic and related secreted bacterial toxins (Iyer et al., 2011b). Indeed, effectors with toxin domains most closely related to the AID-APOBEC deaminases are observed in the Wolbachia endosymbiont of the moth Cadre cautella and the plant pathogen Pseudomonas brassicacearum (Iyer et al., 2011b). Thus, these mutagenic deaminase domains, which were originally part of toxins deployed by bacteria, appear to have provided the basis for the unique mechanism for antigen receptor diversification in vertebrates. However, their role in anti-retroviral response suggests that they were probably initially recruited merely as mutagenic enzymes that targeted viruses (i.e., similar to the original toxin role but merely directed at viruses). Interestingly, several filamentous fungi show a lineage-specific expansion of related nucleic acid deaminases that also appear to have been derived from toxin domains of bacterial provenance (Iyer et al., 2011b). It is conceivable that these play a similar role as the counter-retroviral deaminases in potentially mutating cytoplasmic parasitic elements or preventing anastomosis by unrelated hyphae.
In jawed vertebrates, antibody and T-cell receptor diversity is generated by the action (V-D-J and V-J recombination) of a dedicated recombination apparatus comprised of two proteins Rag1 and Rag2, of which Rag1 is the catalytic subunit of the recombinase (Schatz and Swanson, 2011). The origin of the Rag1 recombinase in animals had remained mysterious until it was shown that their recombinase domain is related to the transposase domain of a distinct class of eukaryotic transposons known as the Transib elements (Kapitonov and Jurka, 2005; Panchin and Moroz, 2008). This transposase domain contains a distinctive version of the RNaseH fold and cleaves sites associated with the termini of these transposons, which show sequence relationships to V-D-J and V-J recombination sites. Thus, the Rag1 recombinase appears to have evolved from a “domesticated” selfish element whose recombinase domain and terminal recognition sites were reused as a mechanism to generate diversity. Indeed, domestication of selfish elements for generation of diversity in host-pathogen interfaces is a general phenomenon, which is not restricted to the animal immune system: In certain caudate bacteriophages, the mutagenic reverse transcriptase of an integrated retroelement has been shown to play a role in creating sequence diversity in a tail-fiber-associated protein (Medhekar and Miller, 2007). This allows the bacteriophages to recognize a changing landscape of cell-surface proteins on their hosts.

**Was the origin of the eukaryotic nucleus-related to inter-organismal and intra-genomic conflicts?**

As the endosymbiotic model for eukaryogenesis involves juxtaposition of two distinct genomes in the same cell, it implies an increased scope for genetic conflicts between the genomes and the intra-genomic selfish elements contained by them. Indeed, different scenarios exploiting such conflicts have been proposed. One of these argues that the mobile self-splicing group-II introns from the alphaproteobacterial mitochondrial progenitor invaded and proliferated in the progenitor of the nuclear genome (Koonin, 2006). As a consequence there was selection for the nuclear membrane as a physical barrier to protect unspliced intron-containing transcripts from the translation apparatus. This hypothesis posits that the pre-LECA eukaryotes were enormously enriched in introns (Koonin, 2006) as a consequence of reduced selection due to decreased effective population sizes (Lynch, 2007). However, direct evidence for highly intron-rich pre-LECA genomes is lacking based on available genomes and with the current data it is not possible to distinguish between: (1) the early proliferation of introns in eukaryotes being a consequence of the emergence of a protective barrier of the nucleus and (2) the nucleus being a consequence of the selective pressure imposed by intron proliferation. Moreover, there is little evidence for extensive proliferation of group-II introns in any prokaryotic lineage. In an alternative hypothesis, greater alignment of the genetic interests of the genomes of the pro-mitochondrion and the nucleus is likely to have happened with the transfer of genes, including those encoding ribosomal proteins, from the former genome to the latter (Jekely, 2008). This is likely to have resulted in chimeric ribosomes in the cytoplasm with potentially deleterious effects for both genomes. This hypothesis presents the nucleus as a physical barrier to prevent such chimerism and might also effectively explain the origin of the nucleolus, another defining feature of eukaryotes. It should be noted that nucleus-like structures have convergently evolved in certain representatives of the clade of bacteria uniting the planctomycetes, chlamydialae and verrucomicrobia (McInerney et al., 2011). In these cases there is no evidence for deleterious effects arising from intra-genomic selfish elements like group-II introns or ribosomal chimerism. Indeed alternative selective pressures could have facilitated nucleogenesis.

One key feature of bacterial endo-symbionts/parasites is their deployment of toxin/effector systems that contain nuclease and nucleic acid deaminase domains, both from poly-morphic and host-directed toxin systems (Iyer et al., 2011b; Zhang et al., 2011, 2012). These are observed in a variety of extant endo-symbionts/parasites such as Wolbachia, Ricketsia, Orientia, Odyssella, Legionella, Ameobophillus, and Protochloramidia (Figure 2). Indeed, such genome-targeting toxins are likely to play a role in the chromosomal disruptions produced by Wolbachia in the process of regulating sex-specific survival and killing of incompatible hosts (Duron, 2008). Interestingly, a key nuclear pore component, Nup96/98, has an autoproteolytic ZU5 domain (Mans et al., 2004). ZU5 domains appear to have originated in bacterial cell-surface proteins, such as polymorphic toxins, and play a role in the autoproteolytic processing of toxins on the cell-surface [ZU5 domains were also secondarily acquired again from bacterial sources to give rise to the animal apoptosis regulator PIDD (Riedl and Salvesen, 2007; Zhang et al., 2012)]. It is possible that this key nuclear pore component was derived from a toxin system of the ancestral endosymbiont. Thus, it is likely that nucleic-acid-targeting toxins were deployed by the mitochondrial progenitor, which could have threatened the integrity of the DNA of the nuclear genome precursor. Hence, the nucleus was probably selected for, as a physical barrier to minimize this threat. In this scenario, once the initial endosymbiotic association between the mitochondrial precursor and the archaean was underway, the selective pressure from the DNA-targeting toxins of the mitochondrial precursor favored the emergence of the nucleus very early in the development of the association. The early presence of the nucleus then favored the development of several characteristics of eukaryotes, including those that have been noted in the other hypotheses: (1) it would have allowed transfer of alphaproteobacterial ribosomal genes to the nuclear genome, as chimerism could be avoided due to presence of an additional compartment (Jekely, 2008), eventually leading the origin of the nucleolus. (2) It allowed retroelements associated with group-II introns to proliferate in nucleus (Koonin, 2006). This not only gave rise to introns but also the telomerase (Aravind et al., 2006). (3) The telomerase in turn facilitated the origin of multiple linear chromosomes, whose expression could now be coordinated as they were contained within the nuclear compartment. (4) Linear chromosomes, together with the nucleus, probably selected against the prokaryotic pumping mechanisms for chromosome segregation based on HerA-FtsK-like ATPases, and instead favored a cytoskeleton-based mechanism, which allowed for fixation of the microtubular apparatus. (5) The stabilization of multiple linear chromosomes contained with a nucleus also probably allowed for increased genome sizes in eukaryotes, as it removed the
constraints coming from containing the entire genome on a large circular chromosome segregated by the ATPase pumps.

In conclusion a number of mechanistically distinct scenarios support a role for organismal and genomic conflict systems in eukaryotic nucleogenesis. Further investigations of alternative scenario presented here might provide a new handle to understand key events in eukaryogenesis.

**GENERAL CONCLUSIONS**

In the above discussion, we provide a series of examples from across the eukaryotic phyletic spread for how the interplay between lateral gene flow inter-organismal, inter-genomic, and intra-genomic conflicts has shaped the evolution of numerous functional systems (Figure 4). These examples are by no means meant to be exhaustive—rather, they were chosen in order to provide a glimpse of the sheer variety of biological systems that are affected by the evolutionary contributions from such systems. One key theme that emerges from the above discussion is that domain families gained through lateral transfer in course of intimate inter-organismal associations, such as symbiosis and parasitism, can notably determine the very nature of these interactions. This is strikingly illustrated by the case of apicomplexan adhesion molecules implicated in host interaction: here manifold domains were acquired by the parasites via lateral transfer from their hosts, spawning unique “animal-like” interfaces for interacting with the host (Figure 1). The other recurrent theme, which transcends various biological systems, is how proteins/protein domains originally emerging in the context of various biological conflicts were recycled as regulatory molecules (Figure 4). Of these host-directed toxins, and the toxins, immunity proteins, structural modules and secretory components from bacterial polymorphic toxin systems have a distinct life beyond their locus of provenance in eukaryotic regulatory and defense systems (Iyer et al., 2011b; Zhang et al., 2011, 2012). We outline numerous occasions where these components were incorporated into regulatory systems of eukaryotes, and sometimes might have played a major role in the very origin of these systems. This process appears to be constantly on-going, all the way from the origin of eukaryotes to the terminal tips of the eukaryotic tree (Figure 4). The reason why proteins derived from biological conflict systems appear to be recruited for other functions might be attributed to the consequences of natural selection. Not surprisingly, toxin-immunity systems used in inter-organismal conflict have a large effect on the fitness of both the organisms producing toxins and those defending against them, thereby escalating an arms race situation. Many of the conflict systems deployed by bacteria might even function at the interface of symbiotic and parasitic interactions of bacteria and eukaryotes, thereby developing adaptations to effectively target components of eukaryotic systems. Toxins and immunity proteins of intra-genomic selfish elements are also under multiple levels of selection that foster their diversification. At one level they are under selection to evade host resistance to function effectively as addictive agents. At another level many of them might also be under selection to function as effective stress response mechanisms that allow their host genomes to survive adverse conditions. Consequently, there are strong selective pressures for constant diversification of toxins and the corresponding immunity proteins in various conflict systems. Hence, these biological conflicts could have functioned as evolutionary “nurseries” for innovations in both prokaryotic and eukaryotic proteins. Hence, lateral gene flow from symbionts, parasites and other modes of DNA uptake (Gladyshew et al., 2008; Nikoh et al., 2008) has enabled eukaryotes to have access to and import a “readymade” set of molecular innovations from such biological conflict systems. When recruited in non-conflict biological contexts, they can in turn spur the emergence of new interactions in eukaryotic systems. Thus, number of key eukaryotic innovations can be traced back to the above-described players in biological conflict systems, such as secondary metabolism operons, R-M, polymorphic and host-directed toxins systems, anti-phage systems, phage counter-restriction strategies, and mobile elements. These systems appear to have particularly expanded in bacteria on account of the presence of operons, extensive lateral transfer with several modes of DNA uptake and recombination, perhaps combined with high effective population sizes (Lynch, 2007). Thus, organismal and genomic conflicts as the basis for major molecular innovations, which in turn might facilitate major evolutionary transitions, can be considered a general evolutionary principle.

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