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Drosophila: a polyvalent model to decipher host-pathogen interactions

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In the past few years, several genetically amenable organisms have been used as models for the analysis of host-pathogen interactions. Among them, the fruit fly Drosophila melanogaster has been useful in elucidating signaling pathways and mechanisms that are used by the host to prevent and combat microbial infections. In addition, this model has also been used to identify virulence factors of opportunistic pathogens. Finally, the characterization of naturally infectious pathogens for Drosophila has illustrated its ability to activate immune responses that are adapted to its aggressors and highlights the potential of these flies to disseminate bacterial pathogens. Altogether, these approaches will allow the mechanisms involved in microbial infection and host defense responses to be dissected using genetic and genomic approaches.

Host-pathogen interactions are multi-step processes that involve several factors in both partners. In microbes, this complex process involves adhesion to the host surface and eventually leads to colonization and persistence, which provokes tissue damage and disease; these steps involve at least two sets of so-called virulence genes. The first subset of genes comprises those involved in the regulation of the adhesion process, which involves binding of microbial ligands to host receptors [1]. This molecular interaction is highly specific and has been shown to define the host range of several pathogens [2]. The target cells or organs are defined by the expression pattern of the receptors and their accessibility to the pathogen. The second subset of genes is mostly involved in adaptation and survival of the microbe within the host's hostile environment. In the host, defense mechanisms are activated following infection and eventually permit clearance of the microbe.

Modeling host-pathogen interactions

To study most processes involved in host-pathogen interactions a system must be set up that includes both the host and pathogen. Because many pathogens being investigated infect humans, surrogate model hosts that are amenable to biological analysis are required for their study. *In vitro* models devised on the basis of cultured-cell approaches are very useful, especially to identify host receptors or to dissect the intracellular fate of invasive bacteria. However, *in vivo* surrogate model hosts are preferred to *in vitro* models because they are more relevant

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to the natural disease process. In these in vivo models, the infectious agent should have the same tropism, and the interaction that takes place the same outcome, as occurs in the normal host. Most pathogens usually colonize a restricted number of hosts, thereby limiting the number of models that are available. To overcome this limitation, several strategies have been adopted; one possibility is to study genetically related pathogens that are naturally infectious in laboratory models (e.g. infection of mice by Salmonella enterica serotype Typhimurium). Another strategy is to use artificial routes of infection (e.g. intraperitoneous or intravenous injection) or to use a permissive developmental stage in 'non-host' models (e.g. suckling mouse model for Vibrio cholerae [3]). A third approach has made use of genetically modified animals that express a human receptor for a specific pathogen [2]. However, this latter approach relies on knowledge of the human receptor, which remains to be identified for several pathogens (e.g. Salmonella enterica serotypes).

Genetic analysis is the method chosen by most to dissect the mechanisms involved in complex host-pathogen interactions. It allows the identification of host and bacterial factors through genome-wide screening of genetic variants. Several methodologies have been set up to generate large numbers of bacterial variants. However, despite the availability of mammalian host models, the number of animals that can be used is limited. To overcome this problem, methodologies have been developed that involve screening a relatively large pool of bacterial variants for attenuated virulence phenotypes [4,5]. These methods allow the identification of genes that are required for in vivo virulence; these genes can then be further tested individually. Ideally, the host should also be amenable to genetic analysis. Although mammalian models are amenable to reverse genetics, identification of genes through forward genetics remains a challenge. Therefore, non-mammalian models, such as the mustard weed Arabidopsis thaliana [6,7], the nematode Caenorhabditis elegans [8,9], the social amoeba Dictyostelium discoideum [10,11] and the fruit fly Drosophila melanogaster [12,13], have been used as host models because they are easily amenable to both forward and reverse genetics. Their affordability and short generation time make them suitable models to carry out intensive analysis and therefore provide an appropriate alternative to mammalian models when characterizing the interplay between hosts and pathogens. Other insect models that lack tools for genetic analysis have also been developed, for example,

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the larger size of the greater wax moth *Galleria mellonella* facilitates a biochemical approach [14-16]. The use of these models has revealed a good correlation between pathogenesis in mammals and in lower organisms [17]. In this review, we describe in further detail different features of *Drosophila*-pathogen interactions that will probably improve our understanding of host-pathogen interactions.

Drosophila as a model to characterize the complex host response

Drosophila immunity

Drosophila has developed effective defense mechanisms to protect itself from overwhelming infection in the microorganism-enriched environments in which it spends its entire life. First, the entire body of *Drosophila* is encompassed by a cuticle that prevents microbial penetration. Surface epithelia that are found in areas including the trachea and gut, which are major routes of infection, are also lined with chitinous membrane to prevent direct contact between cells and microorganisms. To avoid colonization by ingested microbes, a hostile environment is maintained in the gut through low pH and constitutive secretion of antimicrobial agents (e.g. lysozyme) [12,18].

When physical barriers are accidentally or experimentally breached, the introduction of microbes within the body cavity activates a strong inducible immune response. Because *Drosophila* has been the favorite model over the past 90 years for deciphering developmental and cellular processes, numerous genetic and genomic tools are available for this system (Box 1). The availability of such tools and various infection methods has facilitated a detailed description of these inducible defense mechanisms.

Box 1. Overview of Drosophila melanogaster genetics

In *Drosophila*, as in many other organisms, mutations can be generated using chemical or physical agents. If chemical agents generate point mutations, ionizing radiations usually generate deletions of various sizes that constitute a good tool to be used to define a physical area in which the mutation is located. Transposon-mediated mutagenesis has also been greatly emphasized because it allows the generation of different types of mutants (disruption or deregulation of generated small deletions by imprecise excision. Directed mutations can also be generated using homologous recombination [50]. Interestingly, mutations are easily tractable through generations using modified chromosomes, known as balancer chromosomes, which prevent meiotic recombination. These chromosomes are associated with phenotypic markers that allow genotypes of interest to be identified.

Synthetic *P*elements are extensively used as a vector for transgenesis [51] and constitute a powerful tool for gene expression management. The yeast UAS-GAL4 system is fully functional in *Drosophila* when carried as transgenes [52], and therefore it has been largely used as an inducible system to express the gene of interest in a spatio-temporal manner [53]. It involves binding of the GAL4 transcriptional activator to UAS sequences, inducing gene expression downstream of these sequences. The generation of such individuals can be easily achieved by crossing transgenic individuals carrying a UAS-transgene with transgenic individuals expression of the GAL4 driver. Depending on the promoter sequence driving the expression of the GAL4 gene, the transgene can therefore be expressed in any tissue or cell type and at any time.

Similarly, the UAS–GAL4 system can be used to drive expression of dsRNA that induces degradation of the mRNA of the target gene, which The *Drosophila* immune response consists of both cellular and humoral responses. Cellular responses mainly involve professional macrophages termed plasmatocytes, which engulf incoming bacteria through phagocytosis [12,18]. Microbes also induce a strong systemic humoral response. Cuticle breakage leads to the rapid activation of proteolytic cascades that provoke coagulation and melanization in an attempt to restrict microbial spreading within the organism. Moreover, numerous immune effectors are secreted by the fat-body into the blood. Expression of the gene that encodes these factors is under the control of two parallel nuclear factor (NF)-KB signaling pathways designated Toll and IMD (immune deficiency), which share strong similarities with those involved in mammalian innate immune responses (Figure 1) [12,18]. A repertoire of effectors, the expression of which is regulated by the Toll and IMD pathways, has been analyzed using microarray experiments [19-22]; for further information, see the Drosophila Immune Regulated Genes website (http://www. cnrs-gif.fr/cgm/immunity/drosophila_immunity_genes.html). Among these effectors are antimicrobial peptides, the expression of which is specified by the class of the infecting microbe. The IMD pathway is activated predominantly by Gram-negative bacteria, and in turn activates the expression of antibacterial peptide-encoding genes (e.g. dipter*icin*), whereas the Toll pathway is predominantly activated by Gram-positive bacteria and fungi, and regulates the expression of genes that encode antifungal peptides (e.g. drosomycin) and also a subset of antibacterial peptides [12,18]. The expression of these two pathways can be monitored in vivo using reporter constructs [e.g. effector – GFP (green fluorescent protein) fusions]

has a role in promoting gene silencing [54]. In contrast to mammals, dsRNA does not provoke general protein synthesis blockage in *Drosophila*. It is therefore possible to clone large fragments of coding sequence as inverted repeats in a UAS expression vector to silence the expression of a specific target gene [54]. Even though RNA interference (RNAi) only generates hypomorphic mutants, it allows the study of genes that, when mutated, are associated with lethal phenotypes. It also permits the contribution of gene mutations in the pleiotropic phenotype to be identified in restricted cell types. The study of early lethal mutations can also be achieved by generating clones of cells that are homozygous for a mutation in an overall heterozygous individual. These clones can be obtained by forcing mitotic crossing-over that is mediated by expressing a site-specific recombinase (e.g. FLP–FRT system [55]).

Finally, the *Drosophila* genome sequence was completed a few years ago [56] and its annotation is under constant refinement [57]. The Genome Annotation Database (http://flybase.net/annot/) contains information about all the genes annotated, including their description, expression profile and mutant availability. In particular, this website gives access to large collections of mutants and transgenic lines that are available in *Drosophila* genetic centers. To date, the *Drosophila* genome is probably one of the most fully annotated eukaryotic genomes to be found in a database. The availability of these data has allowed the construction of DNA chips, allowing genome-wide analysis of different processes and a protein interaction map of the *Drosophila* proteome to be produced [58]; further information can be found at the *Drosophila* Interaction Database (http://portal.curagen.com/cgi-bin/interaction/ flyHome.pl).



Figure 1. Drosophila IMD (immune deficiency) and Toll pathways are predominantly activated by Gram-negative bacteria and by Gram-positive bacteria or fungi, respectively. These two pathways allow Drosophila to adapt its immune response to the class of the invading microbe. The IMD and Toll pathways display strong similarities with mammalian TNFR (tumor necrosis factor receptor) and TLR-IL-1R (Toll-like receptor-interleukin type 1 receptor) pathways, respectively. Shown in brackets are known mammalian homologs to Drosophila components of these pathways. Steps that are mediated by unidentified factors are indicated using question marks. Abbreviations: ANK, ankyrin domain; DD, death domain; DED, death effector domain; FADD, Fas-associated death-domain-containing protein; IKK, I-kB kinase; IRAK, IL-1R associated kinase; PGRP, peptidoglycan recognition protein; Rel, REL homolog domain, RIP, receptor-interacting protein; TAK1, transforming growth factor β (TGF- β)-activated-kinase; TIR, Toll-IL-1R domain.

that constitute a powerful readout to follow the course of the infection [23,24]. Recently, the combination of this latter approach with natural methods of infection has revealed that antimicrobial peptide synthesis is not restricted to the fat-body but can also be induced locally by surface epithelia after contact with incoming microbes [23-25].

Two different methods of infection have been developed to exploit the power of *Drosophila* genetics in the analysis www.sciencedirect.com

of host-pathogen interactions. The first method is injection, which involves either pricking the body cavity of the insect with a sharp needle that has been dipped in bacteria or microinjection of a precise dose of microbes directly into the body cavity (Figure 2a). Because the direct introduction of any microbe into the body allows the invading microorganism to be detected, this method of infection always leads to the activation of an immune response that is specific for the class of microbe injected. This method

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Figure 2. Methods developed to infect *Drosophila*. (a) Bacterial injection is achieved by pricking adult flies in the thorax with a sharp needle that has been dipped into a concentrated bacterial solution or by microinjecting a precise dose of microbes into the body cavity. Through this type of infection, all microbes induce a strong immune response that is specific for that type of microbe as they are in direct contact with immune system sensors. (b) Natural infection of *Drosophila* larvae. To mimic the larval natural environment consisting of decaying fruit that is colonized by microbes, larvae are mixed with a solution of crushed banana containing ~ 10^{11} bacteria per ml. To date, only one strain has been shown to induce an immune response using this method, revealing a specific interaction between the two protagonists. Adapted, with permission, from [59], Academic Press.

could also serve as a possible screen for examining pathogenicity because Drosophila recovery from systemic infection depends on both the microbe and the genetic background of the flies. For example, following injection into Drosophila, Escherichia coli, Micrococcus luteus or Aspergillus fumigatus are not pathogenic for wild-type flies; in contrast to Pseudomonas aeruginosa, Enterococcus faecalis or Fusarium oxysporum. The second method of infection, known as natural infection, consists of either feeding Drosophila larvae or adults with a concentrated bacterial solution that has been mixed with their food [26,27] (Figure 2b) or spraying fungal spores directly onto the fly exoskeleton [28]. To date, only a few microbes have been shown to induce an immune response following natural infection [26,28] suggesting that additional specific interactions are required that are not revealed using the injection method.

The post-genomic era

Understanding the molecular dialogue between a host and its pathogen necessitates the identification of host factors that modulate the fate of the microbe. This can be achieved by the application of heritable RNA interference (RNAi) technology to Drosophila, which allows the generation of directed mutants [29]. Drosophila lines that conditionally express dsRNAs of interest greatly facilitate the silencing of host gene expression. In particular, the role of host factors that have previously been identified as immune responsive genes after bacterial challenge using microarray analyses [19-22] could be assessed using RNAi methodology. Because the Drosophila genome is fully annotated, in vitro dsRNA has been synthesized for every one of its genes [30]; more details can be found at the Drosophila RNAi Screening Center (http://www.flyrnai. org). Coupling RNAi technology with cell-culture-based models enables a functional genome-wide analysis of cellular processes using high-throughput screening to be carried out [30]. This methodology could be applied to analyze the influence of host factors that modulate the fate of bacteria using appropriate readouts, such as cytoskeleton rearrangement, intracellular localization, or bacterial or cellular survival.

Drosophila as a host to identify bacterial virulence factors

Human opportunistic pathogens: Pseudomonas aeruginosa and Serratia marcescens

Opportunistic pathogens are microbes that infect humans afflicted by specific genetic disorders (e.g. cystic fibrosis) or humans that have impaired defense mechanisms or physical barriers that have been breached (e.g. third-degree burns). They are ubiquitous, versatile bacteria that have developed the ability to adapt to a large number of different environmental conditions. Among them, P. aeruginosa and Serratia marcescens are opportunistic pathogens of major impact, and have been shown to be responsible for severe nosocomial infections and, in the case of *P. aeruginosa*, for life-threatening chronic lung infection in cystic fibrosis patients. Their inherent resistance to many antibiotic classes [31,32] constitutes a major clinical problem. This highlights the importance of characterizing their arsenal of virulence factors that are required for pathogenesis with the aim of defining new antibiotic targets. The study of these microbes in mammalian host models requires the development of conditions that mimic the clinical status of infected patients. Several mouse models have been developed but they are expensive and time-consuming. Recent studies have shown that these bacteria have a broad host range and are able to infect invertebrate hosts [11,13,16,17,33,34]. These studies have revealed that P. aeruginosa uses many of the same virulence mechanisms and effectors in both vertebrate and invertebrate hosts. Different studies have shown that *P. aeruginosa* virulence determinants that are known to be important for mammalian pathogenesis, such as global transcriptional regulators, type III secreted effectors or genes involved in quorum-sensing [27,34-36], were involved in fly death (Table 1). In these studies,

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Table 1. Pseudomonas aeruginosa virulence genes of known function required for full virulence in Drosophila model^a

Gene name	Function	Virulence in animal models ^b	Refs
Type III secretion system			
exsA ^c	Transcriptional activator	n.d./n.d./n.d.	[34]
exsD ^c	Negative regulator of type III secretion regulon	n.d. / n.d. / n.d.	[34]
dsbA ^c	Cytoplasmic disulfide bond oxydoreductase	62 / + / 60	[34]
toxA ^d	Exotoxin A – inhibits mammalian protein synthesis	40 / ++ / 2	[35]
Pilus type IV biogenesis a	nd chemotaxis		
pilH, I, J, K, L ^e	Type IV pilus biogenesis	n.d. / n.d. / n.d.	[13,36]
pilV ^{c,e}	Type IV pilus biogenesis	n.d. / n.d. / n.d.	[13,34]
fimV ^e	Peptidoglycan remodeling	n.d. / n.d. / n.d.	[13]
chpA ^e	Component of chemotactic transduction system	n.d. / n.d. / n.d.	[13]
Quorum-sensing and glob	al regulators of virulence		
gacA ^d	Global virulence regulator	0-50/+/10-100	[35]
pqsB ^d	Hydroxy-alkylquinoline synthesis	63 / + / 100	[35]
qscR ^e	Quorum-sensing – dependant genes repressor	n.d. / n.d. / n.d.	[27]
mvfR ^d	Quorum-sensing regulator	35 / n.d. / 3	[35]
Toxins or related			
plcS ^d	Phospholipase C (lyses eukaryotic cells)	40 / + + / 20	[35]
phzB ^d	Phenazine biosynthesis	18 / + / 10-60	[35]
PA5441 ^e	C.H.P., ^f Probably pyoverdine production	n.d. / n.d. / n.d.	[36]
Other transcriptional regu	lators		
mtrR ^d	Transcriptional regulator of multidrug transporter	53/+/n.d.	[35]
pstP ^d	Transcriptional regulator; RpoN-dependant operons	0 / + / 600	[35]
General metabolism and u	unknown		
pyrF ^e	Orotidine decarboxylase	n.d. / n.d. / n.d.	[36]
pgm ^e	Phosphoglycerate mutase	n.d. / n.d. / n.d.	[36]
pgk ^e	Phosphoglycerate kinase	n.d. / n.d. / n.d.	[36]
cca ^e	tRNA nucleotidyl transferase	n.d. / n.d. / n.d.	[36]
PA3001 ^e	Probable phosphate dehydrogenase	n.d. / n.d. / n.d.	[36]
PA4489 ^e	C.H.P. ^f	n.d. / n.d. / n.d.	[36]

^aMore information, including GenBank accession numbers, is available at the *Pseudomonas* website (http://www.pseudomonas.com)

^bBurnt mouse % lethality (wild-type strain 100%)/*Caenorhabditis elegans* killing (++, wild-type; +, reduced) [33]/*Galleria mellonella* LD50 (wild-type 1) [17]; n.d., data not available. Some gene names have been updated from the date of first publication [60].

^c*Pseudomonas aeruginosa* CHA strain.

^dPA14 strain. ^ePA01 strain.

^fC.H.P., conserved hypothetical protein.

Drosophila was used to test bacterial mutants after a primary screen had been carried out in other organisms. Another study has reported the use of *Drosophila* to screen a library of P. aeruginosa genetic variants [13], which led to the P. aeruginosa pil-chp transduction system being identified as playing an important role in virulence, but its role in a mammalian model remains to be confirmed. Similar studies have been carried out with S. marcescens using a multiple-host approach [37]. In this report, the authors first screened C. elegans for virulence factors before testing clones that were impaired in virulence in other invertebrate and mammalian models. In addition, it was shown that most of the genes found involved in S. marcescens virulence have counterparts in P. aeruginosa, in which they also play an important role in virulence. Altogether, these studies have revealed the existence of conserved virulence mechanisms among opportunistic pathogens that will eventually define the basic requirements for bacterial virulence.

Even though the requirement for some virulence factors depends on the host [37,38], invertebrate models provide a powerful tool to reveal new aspects of bacterial pathogenesis. These examples show that the use of *Drosophila* as a model host provides novel insights into the identification of both host and pathogen factors that either enhance or restrict bacterial pathogenesis.

Mycobacterium marinum: a macrophage-hidden bacterium

Mycobacterium tuberculosis causes severe respiratory infections in humans, however, it is difficult to work with established models of mycobacterial infection. Mycobacterium marinum is an appropriate mycobacterial model because it is closely related to *M. tuberculosis* and displays pathogenic properties in its natural hosts (fish and frogs) that are similar to those of M. tuberculosis in humans. In both cases, the bacteria escape from the host immune system by hiding in macrophages. Notably, mutants of *M. marinum* that are impaired in macrophage survival can be complemented using homologous genes from M. tuberculosis [39], suggesting that these two bacteria share similar virulence mechanisms. Recently, Dionne *et al.* [40] showed that injection of M. marinum expressing GFP into flies resulted in the accumulation of bacteria within hemocytes. This study showed that after injection the tagged bacteria localized in hemocytes, suggesting a macrophage tropism. In hemocytes that were recovered by bleeding, internalized M. marinum did 240

not co-localize with either acidified organelles or internalized dead E. coli, suggesting effective subversion of phagosome maturation similar to that described for vertebrates. Altogether, these results with M. marinum indicate that Drosophila is a promising in vivo model to study bacteria that use macrophages as host cells and has the potential to be applied to other bacteria that escape host defenses by hiding out in macrophages.

Use of Drosophila to identify virulence factor targets

Another promising approach is the conditional expression of microbial factors in Drosophila that are normally injected into host cells through type III secretion systems; this approach might permit the identification of host factors that are targeted by the type III effector proteins. Type III secretion of virulence factors is a common strategy used by different classes of pathogens to interfere with the cell physiology [41]. In vivo genetic studies have been performed where the type III secreted protein is expressed in the yeast Saccharomyces cerevisiae system to determine their function [42]. Owing to the recent developments in fly genetics, it is possible to express genes in a spatiotemporal-dependent manner in non-vital, easy to screen organs, such as the eye. As observed with the HIV accessory protein Vpu (S. Netter, pers. commun.), conditional expression of type III effectors in the eye might lead to morphological modifications. By crossing such transgenic lines with mutant collections, it might be possible to identify genes that suppress or reinforce these phenotypes, therefore revealing genetic interactions. Biochemical and genetic approaches have been successfully applied in Drosophila to identify and study targets of HIV accessory proteins [43,44].

Drosophila as a host in natural bacterial infections

The *Drosophila* injection model has been shown to be relevant for identifying pathogen virulence factors that are important for mammalian infection. However, this method bypasses the first steps of the infection process. Therefore, isolation of the bacteria that elicit a *Drosophila* immune response through natural routes of infection might reveal strategies that are used by microbes to persist in their natural host. Deciphering host-pathogen interactions in such a way will provide new insights into the initial stages of infection. Because virulence pathways appear to be conserved [45], this will help generate an integrated model for host-pathogen interactions.

Drosophila live in an environment full of microorganisms. However, only a few microbes are known either to trigger the immune response [26] or to be pathogenic [27,46] in *Drosophila* following ingestion. Among pathogenic strains, the *qscR* mutant of *P. aeruginosa* kills *Drosophila* adults faster than the parental strain, apparently by constitutively expressing several quorum-sensing regulated genes [27]. However, induction of the immune response has not been reported.

Using Drosophila lines expressing a diptericin-gfp reporter gene that reports a Gram-negative inducible response, Basset *et al.* [26] isolated bacterial strains of *Erwinia carotovora* spp. carotovora (also called *Pectobacterium carotovorum*) that are able to induce a strong immune response [26]. E. carotovora are phytopathogenic Gram-negative bacteria of agronomic interest as these bacteria are responsible for fruit soft rot. Among *Erwinia* strains, several induce a strong immune response in Drosophila larvae by natural infection but are not lethal to their host. It is now possible to apply genetic and genomic approaches on both partners to dissect this 'model' interaction. A genetic screen that was carried out with one of these E. carotovora strains, Ecc15, identified a single gene (evf) that is required for this interaction [47]. Importantly, transfer of evf to several Gram-negative bacteria increases bacterial persistence within the Drosophila gut and leads to the induction of immune responses. The *evf* gene is regulated by *hor*, a general regulator of virulence in several Gram-negative bacteria [48]. In Ecc15, *hor* is required for the infection of both *Drosophila* and plants, whereas evf was found only in E. corotovora strains that are able to infect Drosophila [47]. These results suggest that bacteria might acquire virulence factors and integrate them into pre-existing regulatory networks, which might constitute a potent evolutionary mechanism to broaden host range.

Finally, *Drosophila* is known to be a potential vector of bacterial pathogens [49]. The *Drosophila*-Ecc15 interaction revealed, for the first time, the genetic basis of bacterial persistence in its fly vector. By contrast, the *evf* gene was not found in other *Erwinia* strains that are infectious in *Drosophila*, suggesting that parallel strategies have been selected to allow longer persistence to occur in *Drosophila*.

Concluding remarks

The challenge of deciphering host-pathogen interactions is to understand both counterparts in this complex process. Identifying microbial factors required for *in vivo* virulence has been facilitated by the development of powerful genetic methodologies. Comparative genomics of bacterial pathogens will also give clues about conserved pathways and tropism specificity. Combined with genetic and functional data, it will allow the development of a general model of the role of various virulence genes during infection. The next step will be to build an integrated network of these genes and to understand their role and interplay in the complex process of infection. Such a network would allow the identification of general mechanisms that could be targets of new antimicrobial therapies. Invertebrate models constitute a promising alternative to mammalian models to carry out such analyses. Among these, Drosophila possesses induce defense mechanisms that are similar to those of mammals with respect to both the regulatory pathways involved and the spectrum of their effectors. Moreover, Drosophila is amenable to both forward and reverse genetics, making it a powerful genetic model to study host-pathogen interactions. The study of naturally infectious bacterial strains that are genetically related to human pathogens could highlight the role of virulence factors in the physiopathology of infection. Concerted efforts on the physiopathology of fly infection beyond the immune response would broaden the range of phenotypes and biological processes that can be analyzed.

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Microbiology websites

The Microbiology information portal

http://www.microbes.info/

A useful website providing a variety of microbiology links, including links to feature articles and microbiology-related news stories.

All the virology on the WWW

http://www.tulane.edu/~dmsander/garryfavweb.html

A comprehensive website on virology, including great pictures, virology jobs, a bookshop and links to other useful sites.

The Picornavirus homepage

http://www.iah.bbsrc.ac.uk/virus/Picornaviridae/

A website dedicated to the family of the *Picornaviridae*. Includes important up-to-date news features and articles, ongoing research at the Institute for Animal Health, comprehensive virus classification, links to other useful websites and details of future EUROPIC conferences.

The WWW Virtual Library: Mycology

http://biodiversity.bio.uno.edu/~fungi/ A useful website that provides extensive links to other related sites.

The World Wide Web Virtual Library: Parasitology

http://www.diplectanum.dsl.pipex.com/purls/ A website that provides extensive links to other related sites.

Parasitology links

http://www.galenica.cl/club/rec_parasitologia.html

The world of parasites

http://martin.parasitology.mcgill.ca/JIMSPAGE/WORLDOF.HTM Find out what parasites live with you in your country!

The aspergillus website

http://www.aspergillus.man.ac.uk/ This site includes laboratory protocols, treatment information, DNA sequence data, a comprehensive bibliographic database, an image library and discussion groups.

The E. coli index

http://web.bham.ac.uk/bcm4ght6/res.html A comprehensive guide to information relating to the model organism *Escherichia coli.*

Bacterial infections and mycoses

http://www.mic.ki.se/Diseases/c1.html

Lots of useful bacterial links.