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Taking insect immunity to the single-cell level

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Abstract (120 words)

For decades, insect immunology has contributed groundbreaking discoveries on the intricacies of innate immunity. It has profoundly impacted our understanding of innate immunology in mammalian systems and improved human therapeutic interventions, from the composition of vaccines to cell-based immune therapies. Our current knowledge of insect immunity mainly encompasses detailed molecular mechanisms and systemic responses to pathogen infection. However, the organs and specific cell populations involved in immune responses remain elusive. Here, we highlight the need for a better understanding of insect immune responses at the level of organs and cell populations. Not only will it improve our current understanding of tissue or cell-specific immune processes across species, but it will also pave the way for spatial modeling of within-host infection dynamics.
Over the last few decades, intense research efforts exploiting the power of the model organism *Drosophila melanogaster* have led to fruitful discoveries in the field of insect immunology and impacted the field of innate immunity at large [1]. Innate immune responses constitute the first line of defense against microbial agents and are defined by their rapid activation, efficiency, and lack of memory. Pathogen recognition is mediated by germline-encoded receptors recognizing pathogen-associated molecular patterns (PAMPs). Innate immunity is often distinguished from adaptive immunity, which emerged approximately 500 million years ago in vertebrates. Adaptive immune responses are based on pathogen recognition by highly specific antibodies generated by somatic gene rearrangements and long-term protective memory to counter pathogen re-exposures [2].

Detailed insights into innate antibacterial, antifungal, and antiviral immunity have been obtained in *Drosophila*, notably because of the vast genetic toolbox available in this organism. Early studies focused on the mechanisms of pathogen recognition and subsequent signal transduction. The discovery of an inducible antibacterial immune response in the fruit fly, first reported in 1972, led to the discovery of antimicrobial peptides (AMPs) that are not only conserved from insects to mammals, but are also controlled by highly conserved nuclear factor-κB (NF-κB) signaling cascades [3]. In 1997, the discovery of the Toll receptor as a key factor in *D. melanogaster* antifungal immunity [4] was a turning point for the field, as this signaling pathway turned out to be homologous to the myeloid differentiation primary response protein 88 (MYD88)-dependent Toll-like receptor (TLR) pathway in mammals [1].

Since those early days, critical molecular mechanisms for pathogen recognition have been elucidated, and the coordination of immune responses at the organismal level have been largely characterized [5]. Briefly, in insects (*Drosophila* sp.), humoral immunity is controlled by a set of highly conserved signaling pathways. The Toll and the Immune deficiency (IMD) pathways mainly control bacterial and fungal infections, and the Janus kinase-signal transducer and activator of transcription (Jak-Stat) and...
RNA interference (RNAi) pathways are primarily involved in antiviral defense [6]. In reality, these pathways are pleiotropic, and their roles against a specific class of pathogens are not always clearly distinguishable. Toll and IMD pathways also participate in antiviral immune responses [6], and the Jak-Stat pathway mediates the response to tissue damage that occurs upon bacterial, fungal, and parasitic infections [7].

While humoral immunity is relatively well-characterized, cellular immunity in insects (Drosophila sp.) remains more enigmatic. The best characterized immune cells are hemocytes, which circulate in the insect hemolymph and differentiate in four main subtypes: plasmatocytes, granulocytes, lamellocytes, and oenocytoids [8]. Our understanding of the immune pathways activated in those cells remains a black box. Moreover, little is known about their movement and communication within immune organs, somewhat due to the technical challenges linked to their study. In fact, harvesting hemocytes from an insect’s body cavity requires great technical skills [9] and their number ranges from ≈500 to 3,000, depending on the age, genetic strain, developmental stage, and infection status of the specific insect species [10, 11].

Beyond immune cells, immune organs are also poorly characterized. In adult insects, the fat body tissue has, in addition to its metabolic function, a central role in immunity [5]. It is a secretion site for antimicrobial peptides, but also a central signaling hub for the activation of cellular immunity [12]. In Drosophila, the fat body activates hemocytes, through a yet to be discovered mechanism that is Toll-dependent [13]. The exact cellular composition of the fat body remains unknown, as is the identity of the cells that can communicate with hemocytes, or coordinate immune processes.

In addition to Drosophila, non-model insect species are more challenging to study but also medically more relevant. This review mainly highlights recent studies in mosquitoes (Anopheles and Aedes genera) that contributed seminal discoveries to the field. The completion of genome sequencing and assembly in important vector species, as well as the advent of RNAi-mediated knock-down and CRISPR-Cas9 technologies, have been instrumental in the study of mosquito immunity [14-16]. Comparative genomics
reveal a high degree of conservation of immune signaling pathways, combined with specific adaptations to particular pathogens or lifestyles. Overall, the core signaling pathways are well conserved between Drosophila and mosquitoes, whereas upstream signaling factors and downstream effectors diverged during evolution [17, 18]. The opinions expressed in this paper are mostly based on work in Drosophila and mosquitoes but are widely applicable to the study of ticks (Ixodes genus), sandflies (Phelobotominae subfamily) and further arthropods that are the focus of remarkable research efforts (Box 1).

Single-cell technologies encompass a vast array of methods to study genomics, epigenomics, proteomics and metabolomics at the single-cell level [19-21]. Here, we focus on single-cell RNA sequencing (scRNA-seq) and single-cell transcriptomics, which are on their own, revolutionary and paradigm-shifting. In 2009, the first study using single-cell transcriptome analysis based on a high-throughput sequencing platform led to the characterization of cells from early-stage murine embryos [22]. Several protocols have been developed over the last decade, leading to an impressive increase in throughput and data quality [21]. Historically, for many technical reasons, transcriptomic studies have typically been performed on large groups of cells originating from multiple individuals, multiple tissues, or pools of individual or tissues, therefore referred to as bulk RNA-seq. Hence, several levels of biological variation, including organismal, tissular and cellular differences, could be “hidden” in one transcriptome data set (Figure 1). Single-cell transcriptomics allow for the first time, to study cell-to-cell transcriptional heterogeneity in insects, including the gene expression stochasticity that was first reported over 15 years ago [23].

How can single-cell technologies help to accomplish a leap forward for studies of insect immune systems? Could these technologies enable, in the near future, the acquisition of a more comprehensive picture of insect immunity? Can they integrate into the same model, our knowledge of molecular mechanisms and systemic responses by bridging those components with a better knowledge of cellular and tissular immunity? In this opinion article, we argue that the increasing availability and applicability of scRNA-seq
present a new opportunity to deepen our knowledge of insect immunity and impact our global knowledge of innate immunology across species.

How can single-cell transcriptomics be instrumental to the study of insect immunity?

Deciphering the cellular composition of organs

The cellular composition of organs, including immune sites, remains to be investigated in insects. Early studies based on cellular morphology helped decipher the composition of tissues and led to the classification of hemocytes [24] and digestive tract cells [25]. However, a recent study revealed that Anopheles gambiae mosquito hemocytes display a high level of complexity, given that two highly distinct cellular “populations” -- as defined by their transcriptional profile -- actually share morphological and functional features, such as their phagocytic capacity [26]. Therefore, the classification of cell types historically based on morphology or function might need to be revisited. In addition, little is known about the cellular re-composition of organs after an infectious episode. In Drosophila, infection of the intestinal tissue with the Gram-negative bacterium Erwinia caratovora has resulted in dramatically increased stem cell proliferation and epithelial renewal, as shown by the increase of mitotic recombination in the intestinal tissue, accompanied by a higher number of stem cells (expressing the escargot marker gene) upon infection [27]. However, it remains to be determined whether the cellular content of the tissue pre and post pathogen exposure is comparable, or if infection-induced changes of immune site composition could impact subsequent immune responses. In fact, Plasmodium berghei (rodent malaria)-challenged mosquitoes constitutively secrete a hemocyte differentiation factor into their hemolymph that induces immune priming when experimentally transferred into naive mosquitoes [28]. The hemocyte differentiation factor was first discovered in mosquito hemolymph by mass spectrometry, and is a complex consisting of a protein (called Evokin) that is a lipid carrier of lipoxins [28]. Immune priming is characterized by an increased number of granulocytes (a subpopulation of hemocytes) and can be measured by extracting hemolymph from mosquitoes and counting the number of cells
present while assessing their morphology and binding properties [29]. P. berghei-challenged mosquitoes mount a strong immune response when the parasite breaks through the gut and, by doing so, brings the gut bacteria in contact with injured epithelial gut cell. This stimulation creates a state of systemic immune surveillance upon which the immune system is primed to respond again to a new exposure. Upon re-infection, the immune system mounts a stronger antibacterial response that is also detrimental to the survival of Plasmodium sp. parasites [29]. It will be interesting in the future to analyze the transcriptome of these cell populations at the single-cell level and understand the depth of immune priming processes. In fact, this could yield new insights into the mechanisms of immune memory that are so far poorly understood in insects.

Discovering new immune cell populations

In addition to discovering new cell populations that are induced by infection, single-cell sequencing can also enable identification of cells that hold an immune function based on their transcriptional similarity with existing immune cells, but have not yet been characterized as such. In fact, it is likely that most tissues contain cells that mediate their defense and viability upon infection but are rare, non-canonical, or difficult to characterize. For instance, the immune capacity of glial cells in the Drosophila brain was only discovered recently, with the observation that they were able to secrete AMPs [30]. Additionally, infection of nervous tissues such as the brain by Zika virus has recently be modeled in Drosophila and pointed to a role for the IMD pathway in glial cells [31]. In fact, mutant flies that have a deficient IMD signaling failed to induce autophagy, an antiviral process that helps reducing Zika virus infection [31]. Single-cell sequencing, which is virtually applicable to any tissue or organs, is likely to contribute exciting discoveries of new immune cell types in insects that were so far unreachable using traditional methods.

Assessing heterogeneity of immune responses to infection
Recent studies on mammalian cells such as the human lung epithelial carcinoma line A549, the human hepatoma line (Huh7), or the murine fibroblast L929 line, utilized scRNA-seq to assess the diversity of the transcriptional responses to infections by influenza, dengue, Zika and West Nile viruses [32-35]. Taken together, these studies revealed astonishing heterogeneity at the level of host transcriptomes and viral abundance from cell to cell. The measurement of both host and viral RNA transcripts (also called visc-RNAseq) in the same cells can reveal correlations between gene expression, and make possible the detection of strong pro- or antiviral candidate genes [33]; indeed, this was previously done in a bulk RNA-seq study on dengue virus-infected midguts of *Aedes aegypti* mosquitoes, that identified the sterol regulatory element-binding protein (SREBP) as a host factor with a proviral role in the mosquito midgut [36]. Moreover, A scRNA-seq study performed on peripheral blood mononuclear cells isolated from dengue virus-infected human patients identified a set of genes that might be used as biomarkers for disease progression to severe dengue fever [37]. For example, the expression of MX2 within naive B cells is strongly upregulated before the onset of severe dengue fever [37]. The predictive power of such markers discovered by visc-RNAseq remains to be tested with larger study cohorts. Finally, quantifying the heterogeneity of immune responses to infection in relation to pathogen presence might also help identify by-stander effects of cells near infection sites that are not infected themselves, but which deploy immune responses linked to the detection of infection-linked signals in their environment. For example, in *Drosophila*, reactive oxygen species are released from necrotic cells and act as an immediate danger signal which can induce the recruitment of hemocytes to the injured tissue [38]. Indeed, it will be interesting to compare immune profiles of infected cells versus uninfected cells that have been alerted of an ongoing infection by danger signals in their intra-host environment.

**Assessing heterogeneity in pathogen genetic diversity upon infection**

Pathogen-inclusive scRNA-seq opens possibilities to assessing pathogen genetic diversity such as viral quasispecies at the intra-cellular level, as has already been done for hepatitis C virus, human immunodeficiency virus and vesicular stomatitis virus.
In insect vectors of viruses, it will be possible to identify cellular subtypes that are responsible for increasing viral diversity, or on the contrary, act as population bottlenecks. In fact, the organs and cell types that are sites of viral replication and enable the virus to genetically diversify or homogenize, are not characterized despite being important targets for future vector-control strategies. Intra-host viral diversity can impact viral fitness, and perhaps the transmission efficacy of a given virus population, as well as potentially, the epidemiology of an outbreak [40, 41]. For example, it was shown that the fitness of West Nile virus decreased as it spread from one mosquito organ to another: the virus replicated poorly in avian cells after a passage through Culex quinquefasciatus mosquitoes [41]. Single-cell technologies might help identify cellular sites and molecular mechanisms of viral diversity within the insect host and potentially support genetic interventions aiming at reducing pathogen transmission.

Beyond viral infections, a recent scRNA-seq study that focused on Plasmodium sp. parasites revealed discrete transcriptional signatures during the parasite’s life cycle, as well as unexpected sex-specific roles for contingency gene families (var in P. falciparum and pir in P. berghei) that are known for enabling parasite adaptation to new environments [42]. In addition, scRNA-seq enabled the Malaria Cell Atlas to characterize thousands of single-parasite transcriptomes across the entire Plasmodium sp. life cycle [43]. Those studies are relevant as they may open multiple new research avenues -- from the analysis of nonculturable life stages of Plasmodium species (such as P. vivax), to the analysis of cell-to-cell variability in response to drugs and vaccines, as well as the development of blocking strategies against infection in humans. Overall, pathogen-inclusive scRNA-seq offers exciting research avenues that will allow to assess the impact of the host cells on pathogen evolution and adaptation.

What are the main challenges of single-cell sequencing implementation in insects?

Dissociation and fixation
The first step of any single-cell sequencing experiment consists of the preparation of the cell suspension. With the exception of hemocytes that circulate in the insect’s hemolymph, cells are embedded in tissues and organs that have to be dissociated in order to obtain a suspension of single cells. Dissociation methods are varied, ranging from the use of a diverse set of enzymes to the use of mechanical force (Table 1). Most often, several methods need to be tested, as their efficiency is often dependent on the cell type, tissue, organism, and insect species. Dissociation is an aggressive process that causes cellular stress and therefore transcriptional changes related to stress responses. Accordingly, if feasible, it is preferable to fix cells and thus preserve the transcriptome before dissociation. Several fixation agents are available (paraformaldehyde, acetone, methanol, Lomant’s reagent, RNAlater™, vivoPHIX™); some have been tested in side-by-side studies with live cells and do not seem to bias or alter the quality of the transcriptome [44, 45]. In all cases, a set of quality indicators should be measured: (i) dissociation efficacy, (ii) cell viability and (iii) cell integrity (Box 2). Importantly, dissociation methods can potentially affect the transcriptome of cells or a subset of cells [46]. It was recently shown that a subset of cells detected in several single-cell studies, for example in mouse muscle cells or zebra fish osteoblasts, was over expressing genes associated with tissue injury (i.e. heat shock proteins such as Hspa1a and Hspa1b). However, those cells were not found as a functional subset in vivo, pointing to a likely experimental artifact [46]. Noteworthy, these ‘injured’ cells might also be present as potential contaminants in bulk RNA-seq studies. However, such transcriptional signals would be harder or likely impossible to identify, as they would be averaged at the level of an entire tissue or individual. By contrast, In a single-cell study, ‘injured’ cells could be identified and excluded from subsequent analyses.

Insect cells present additional challenges at the dissociation step due to their limited number and reduced size. Insect cells are often smaller in size than their mammalian counterparts [47, 48]. For instance, Drosophila neurons are ten times smaller than mammalian neurons [49]. Moreover, insect organs and tissues are small in size. The Drosophila brain contains approximately 100,000 cells [48], while the human brain contains 100 billion cells [50]. While it is possible to dissect high numbers of organs for single-cell experiments in insects, never will it be possible to reach the order of
magnitudes of larger organisms. This dramatically complicates the execution of single-cell experiments, not only because of the sheer number of cells available but also because cell recovery after dissociation and cell viability have to be extraordinary. Once dissociation and potential fixation are optimized, RNA sequencing can take place. Apart from cellular sizes that might not be adapted to all microfluidic platforms, which are generally designed for mammalian cells, no specific considerations need to be taken for the sequencing of insect cells.

Data analysis and interpretation

Amongst the significant applications of scRNA-seq is the identification of novel cell types and states, founded on the detection of co-expression patterns and clustering of cells based on their transcriptomic similarity. Cell clusters can be analyzed further by identifying gene sets characteristic of each cluster, also called markers. Whereas markers for mammalian immune cells are generally well known, they remain elusive for most insect cells. Cluster analysis and marker discovery is, therefore, more challenging for insect cells, as preliminary knowledge about sub-groups to be detected is scarce. Recently, an unbiased classification of An. gambiae hemocytes was performed using a combination of scRNA-seq, high-content imaging flow cytometry, and single-molecule RNA hybridization [26]. It revealed an unexpected degree of complexity in the hemocyte population together with the existence of cellular cross-talk via exchange of mRNA between cells [26]. This combination of required tools highlights the need for orthogonal validation methods, especially in non-model insect systems for which there is little preconceived knowledge about cell populations, and almost no cell markers available. Overall, scRNA-seq data analysis and interpretation is challenging for insect cells and, given the poor availability of markers for many cell populations, will require cross-validation with other experimental methods.

How can single-cell sequencing data be exploited to model immune responses in insects?
Single-cell sequencing data sets are comparatively more complex and multidimensional than classical bulk RNA-seq data sets. Beyond the standard analysis methods, consisting of reducing dimensional space (by PCA and t-SNE), functional clustering or trajectory analysis [21], even more sophisticated data analysis methods could help informing insect immunity.

Subjecting multiple tissue samples over an infection’s time course to scRNA-seq could help inform infection dynamics in a deeper manner than has ever been possible. Each data set might describe degree of infection, correlating it with transcriptional responses at the cellular level in different tissues over time. Given the high dimensionality of such data, integration into a mathematical modeling framework is highly desirable. Such information might help identify key immune cells and processes, as well as model the progression of the pathogen through the host’s body. Once built, such a modeling framework could ideally be used to predict infection outcomes, severity, or further pathogen transmission potential.

For instance, when an Aedes mosquito bites a human infected with an arbovirus such as Zika virus, the bloodmeal containing the virus is first ingested in the mosquito midgut. The virus initially replicates in midgut cells before disseminating throughout the mosquito’s body. Eventually, the virus reaches the salivary glands and can be transmitted again through a new bite [51]. The exact progression of the virus throughout the mosquito’s body remains elusive. How does the virus circulate in the hemolymph? Is it carried by hemocytes or does it circulate freely? Which organs does the virus replicate in, and when? Are similar immune pathways triggered in the midgut, fat body, and salivary gland? Can we predict the transmission potential of a certain virus based on the host immune response and viral levels in certain organs? Such fundamental questions might be answered, at least in part, by the use of single-cell technologies and modeling of infection dynamics.

Concluding remarks
For decades, the study of innate immune processes in insects has greatly contributed to our overall knowledge of innate immunology. Here, we propose that the recent developments of single-cell transcriptomics represent an opportunity to make a significant leap forward in this exciting research field. Improving our knowledge of immunity at the cellular and tissular level has the potential to be paradigm shifting at the fundamental level (see outstanding questions). Indeed, it may also have a wider impact on novel strategies deployed to control vector-borne diseases.

Nowadays, performing single-cell transcriptomics remains challenging technically, but also financially. Although the protocols and technologies evolve rapidly, the costs of the technology remain high. Hopefully, the constant decrease of sequencing costs and commercialization of scRNA-seq by more industrial actors will make it more accessible.

Spatial transcriptomics are a recent and exciting technological development [52]. In addition to obtaining single-cell transcriptomes, this technology allows to map the transcriptional activity to its tissular location and place it in a morphological context. In fact, it relies on the deposition of tissue slices on arrays covered with probes that capture RNA. After sequencing of the array, the transcriptomes are reconstituted in space and mapped to their location of origin on the tissue slice [53]. This will help identification of immune cells, infected cells, by-stander cells, rare cell types on a 3D map, and may contribute to a more holistic understanding of immune responses at a cellular, tissular and organismal level.

Here, we focused on the promise of single-cell transcriptomics to improve our knowledge of insect immunology, without mentioning the development of other single-cell based technologies. Performing DNA sequencing at the level of single cells can also help identify cell-to-cell heterogeneity in single-nucleotide polymorphisms and copy number variations. Moreover, DNA mosaicism remains an unexplored territory in insects although it has been extensively investigated in the context of human health [54]. Single-cell proteomics and epigenomics have also been developed more recently, offering multiple and diverse research possibilities in the near future [19, 20].
Finally, we posit that single-cell transcriptomics will need integration at the systems level, i.e. using a holistic approach to capture emerging properties of cells and tissues functioning as a system. This will be possible with the complementary single-cell based technologies aforementioned, and more generally, with various approaches of genomics, proteomics, epigenomics, metabolomics and lipidomics [55]. For now, it remains challenging to perform experiments and analyze samples in such a multi-dimensional manner for technical and financial reasons. However, we predict that within the next decade, the development of sophisticated computational methods will enable seamless data integration and analysis at a systems level. Alexander von Humboldt was the first scientist to suggest that, while each element in nature demanded specialized analytical attention in order to establish its own specific rules of order and organization, the most important challenge was that of integrating all this knowledge. With his 250\textsuperscript{th} birthday, his vision remains as relevant and applicable to systems biology in the 21\textsuperscript{st} century.

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Over the last few decades, RNA sequencing technologies have evolved from using pools of individual insects, to single individuals, to single organs. At every step, the transcriptome obtained is an “averaged” picture of gene expression. It reflects several individuals, several organs, or several cell types. Only using single-cell sequencing is it possible to get a unique cellular transcriptome.

Table 1. Summary of the published work applying scRNA-seq to insect cells.

<table>
<thead>
<tr>
<th>Organism</th>
<th>Cell type</th>
<th>Dissociation method</th>
<th>Fixation method</th>
<th>Sequencing method</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>D. melanogaster</em></td>
<td>Total embryonic cells</td>
<td>Douncer homogenizer (mechanical)</td>
<td>Methanol</td>
<td>Drop-Seq</td>
<td>[56]</td>
</tr>
<tr>
<td><em>D. melanogaster</em></td>
<td>Olfactory projection neurons</td>
<td>Papain Liberase™</td>
<td>/</td>
<td>Smart-Seq2</td>
<td>[57]</td>
</tr>
<tr>
<td><em>D. melanogaster</em></td>
<td>Eye disc cells</td>
<td>Collagenase Trypsin</td>
<td>/</td>
<td>Drop-Seq</td>
<td>[58]</td>
</tr>
<tr>
<td><em>D. melanogaster</em></td>
<td>Total brain cells</td>
<td>Dispase Collagenase I</td>
<td>/</td>
<td>Chromium Smart-Seq2</td>
<td>[48]</td>
</tr>
</tbody>
</table>
Trypsin

Cel-Seq 2
ATAC-seq

<table>
<thead>
<tr>
<th>D. melanogaster</th>
<th>Midbrain cells</th>
<th>Papain / Collagenase I</th>
<th>Drop-seq</th>
<th>[49]</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. gambiae</td>
<td>Blood cells</td>
<td>/</td>
<td>Smart-Seq2</td>
<td>[26]</td>
</tr>
</tbody>
</table>

**Box 1.** scRNA-seq for studying immunity in arthropod vectors beyond mosquitoes.

The majority of published work describing the use of scRNA-seq in insects has been performed in *Drosophila* sp. and mosquitoes (Table 1). However, scRNA-seq is a technology that is applicable to arthropod vectors beyond mosquitoes, such as fleas (*Siphonaptera* order), ticks (*Ixodes* genus), tsetse flies (*Glossina* genus) and sandflies (*Phlebotominae* subfamily). Most current knowledge on insect immune signaling pathways has been acquired in *Drosophila*, and served as a roadmap for studying immunity in other arthropods. However, recent studies have revealed that immune signaling pathways in…. are mechanistically more distinct than anticipated, emphasizing the need for in-depth studies in other arthropods [59, 60]. For example, ticks have a smaller repertoire of Toll receptors compared to *Drosophila* [61] and have also undergone gene losses in the IMD pathway, despite both pathways remaining functional [62, 63]. This suggests that unidentified immune pathway components might be replacing predicted canonical factors to mediate the recognition of microbial invaders in ticks, although this remains to be further investigated [64]. It might be possible in the future to isolate individual cells in which those pathways are active (e.g. using flow cytometry) and analyze their transcriptome by scRNAsseq to find the unknown components that keep the pathway functional. Another recent study [65] analyzed the role of a mutualist bacterial symbiont on the adult immune response of the tsetse fly, *Wigglesworthia* [65]. It revealed that the symbiont induced the upregulation of an odorant binding protein (Obp6) in the gut of intrauterine tsetse larvae relative to…. which was necessary and sufficient to induce the subsequent production of crystal cells mediating the melanotic immune response [65]. In the future, it might be possible to determine, using scRNA-seq, exactly which cell types secrete Obp6, and which tissues are targeted by this molecule. Beyond these examples, we argue that scRNA-seq will help elucidate fascinating biological processes, and enable in-depth
studies in a broad range of arthropods that have so far been understudied because of the limited toolbox available.

Box 2. Essential questions to ask during sample preparation for scRNA-seq.
- How many cells are recovered after dissociation? Is that number consistent?
- Is the tissue fully dissociated?
- What is the percentage of cell death?
- Are the live cells intact and healthy? Are there apoptotic or necrotic cell bodies in the cellular suspension?
- Are the cells metabolically active during dissociation potentially leading to changes in the transcriptome?
- Does the cell suspension contain cellular debris to be filtered?
- Are cells recovered after fixation?
- Are the cells potentially infectious?
- Do the cells need to be used immediately or can they be stored before use?

Glossary (450 words)

**Antimicrobial peptides**: In insects, AMPs are a class of small peptides with potent antimicrobial activity.

**Humoral immunity**: Immune responses mediated by factors present in extracellular fluids, mainly the hemolymph in insects.

**Toll pathway**: Immune pathway of insects mainly triggered by Gram-positive bacteria leading to antimicrobial peptide production.

**IMD pathway**: Immune pathway of insects mainly triggered by Gram-negative bacteria leading to antimicrobial peptide production.

**Jak-Stat pathway**: Immune pathway of insects triggered by the Unpaired cytokines mainly upon viral infection.
RNAi: RNA interference refers to a set of gene-silencing mechanisms induced by double-stranded RNA among which the small interfering RNA (siRNA) pathway is considered a major antiviral defense of insects.

Pleiotropic: A gene or pathway is called pleiotropic when it affects two or more seemingly different phenotypic traits.

Cellular immunity: Immune responses mediated mainly by hemocytes.

CRISPR-Cas9: Prokaryotic immune system based on Clustered Regularly Interspaced Short Palindromic Repeats and the endonuclease Cas9, which was converted into a powerful a gene-editing technique targeting specific sequences recognized by a complementary guide RNA.

Bulk RNA-seq: Conventional RNA sequencing technique that probes the transcriptome of a large number of pooled cells.

Immune priming: analogous to trained immunity, it describes the phenomenon by which a first pathogen exposure enhances host resistance to a subsequent challenge.

viscRNAseq: virus-inclusive RNA sequencing probes the host transcriptome together with intra-cellular viral RNA at the single-cell level.

Viral quasispecies: Cloud of diverse virus variants that are genetically linked through mutation and collectively contribute to the characteristics of the viral population.

Population bottleneck: In evolutionary biology, it is defined as an event that drastically reduces the size of a population.

Viral fitness: Usually equated to replicative fitness, it is defined as the capacity of a virus to produce infectious progeny in a given cellular environment.
PCA: Principal component analysis is a linear dimensionality reduction method commonly used to identify cell subpopulations by clustering their features in 2D or 3D.

t-SNE: t-distributed stochastic neighbor embedding is a dimensionality reduction method that helps identifying cell subpopulations and sometimes provides clearer visualization of clusters than PCA.

RNAi-mediated knock-down: Experimental gene-silencing technique using double-stranded RNA or siRNAs to neutralize target RNA molecules by RNAi.

Plasmatocytes: Macrophage-like cells involved in phagocytosis and encapsulation that are the major constituents of the insect hemocyte population.

Granulocytes: Adhesive cells mediating phagocytosis, encapsulation and degranulation in insects.

Lammelocytes: Large cells involved in encapsulation of invading pathogens in insects.

Oenocytoids: Large cells responsible for the production of prophenoloxidase in insects.

Crystal cells: Cells responsible for humoral melanization that accompanies some immune reaction in insects.

Pathogen-Associated Molecular Patterns (PAMPs): Diverse set of molecules that share a set of general “patterns,” or structures, that alert immune cells to destroy invading pathogens. One example is lipopolysaccharides present in the membrane of Gram-negative bacteria.

Toll-like receptor (TLR): A class of pattern recognition receptors that initiate the innate immune response by sensing pathogen-associated molecular patterns.
**Hemocytes:** A group of several cell types that are found in the hemolymph of insects.

**Copy number variation:** Phenomenon in which the number of copies of a particular gene varies from one individual to the next.

**DNA mosaicism:** Phenomenon that involves the presence of two or more populations of cells with different genotypes in one individual that is called a mosaic.

**Melanotic immune response:** A branch of the innate immune response that involves the synthesis of melanin, a dark pigment that can encapsulate invading pathogens or unhealthy cells.

**References**


Figure 1

Pool of multiple individuals

Single individual

Single organ

Single cells

Gene X expression

Individual 1
Individual 2
Individual 3
Individual 4

Organ 1
Organ 2
Organ 3
Organ 4

Cell 1
Cell 2
Cell 3
Cell 4