Review Article

Novel Technologies in Studying Brain Immune Response

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Over the past few decades, the immune system, including both the adaptive and innate immune systems, proved to be essential and critical to brain damage and recovery in the pathogenesis of several diseases, opening a new avenue for developing new immunomodulatory therapies and novel treatments for many neurological diseases. However, due to the specificity and structural complexity of the central nervous system (CNS), and the limit of the related technologies, the biology of the immune response in the brain is still poorly understood. Here, we discuss the application of novel technologies in studying the brain immune response, including single-cell RNA analysis, cytometry by time-of-flight, and whole-genome transcriptomic and proteomic analysis. We believe that advancements in technology related to immune research will provide an optimistic future for brain repair.

1. Introduction

The role of the immune system in brain function and development has been highlighted by several studies [1-3]. Immunomodulation has even been considered a potential therapeutic strategy for neurodegenerative dysfunctions, including Alzheimer’s disease, stroke, and other neurodegenerative conditions [4].

Immune cells are complex and dynamic and characterized by diverse cell profiles. Besides, their activity largely relies on their interactions with each other, along with other pathological processes following brain injury [5, 6]. As a result, to obtain a comprehensive knowledge of the brain immune system, detailed profile information of each cell is required. The progress of human science and technology, especially the establishment and improvement of RNA sequencing, mass cytometry, and whole-genome transcriptomic and proteomic analysis, allows us to delve deeper and wider into how the brain immune system functions and responds to various pathophysiological conditions. In these techniques, RNA sequencing methods have assisted in depicting the complexity and diversity of the examined immune cells [7]. Cytometry by time-of-flight (CyTOF) overcomes the limitations of traditional flow cytometry approaches to analyze more than 20 parameters, allowing us to detect single-cell protein information, which is vital for revealing cell functions [8]. The focus of whole-genome transcriptomic and proteomic analysis is to chart and map cell information at DNA, RNA, and protein levels, plotting a much deeper, uncharted territory for future neurobiologists. In this paper, we will review the use of these technologies in the brain’s immune system. The literature was searched using Medline between 2006 and the present using search terms including “single-cell RNA”, “Cytometry by time-of-flight”, “whole-genome transcriptomic and proteomic analysis”, and “immune response”. The search terms were cross-referenced, and the search was limited to English...
language articles. All of the articles, found including those associated with the initial search results, were evaluated for methodology and results and were included if deemed applicable to this review.

2. Immune Response in the Brain

The brain was previously thought to be an “immunologically privileged” organ because peripheral immune cells have been considered unable to cross the blood–brain barrier. Now, mounting evidence suggests that glial cells, particularly microglia and astrocytes, constitute and modulate the complex neuroimmune system of the brain in response to infection and drug intervention [1, 2]. As research continues, macrophages, oligodendrocytes, and endothelial cells have also been found to play an important role in the process of neuroimmunity [9]. Activation of these neuroimmune cells may impair healthy neurons and result in brain damage by releasing pro-/anti-inflammatory cytokines and reactive oxygen species (ROS). Notably, this system is also important for brain metabolism and development by regulating synaptogenesis, neurotransmitter transmission, and trophic support for neurons [10, 11]. Therefore, brain immunity may play a pivotal role in the pathology of several neurological disorders and the development of the brain. However, the process and mechanism of the neuroimmune system are not completely understood for the structural complexity of the brain. The possibility to elucidate the current neuroimmune state has attracted a remarkable interest of a large number of neuroscientists. New insights in this field will provide novel methods to measure the neuroinflammatory process, facilitate the discovery of potential effective biomarkers, and eventually lead to the alleviation, treatment, and prevention of these nervous system diseases which may stimulate a neuroimmune response.

3. Novel Tools in Studying Brain Immune Response

3.1. Single-Cell RNA-seq Technology. In recent years, single-cell RNA-seq (scRNA-seq) has become widely used for transcriptome analysis in the field of the immune system. It provides quantitative measurements of RNA molecules within a single cell, leading to the present rapidly expanding world of single-cell transcriptomics. Single-cell genomics makes it possible to create a high-resolution characterization of cells and allows the profiling of numerous molecules from single cells. It is hoped that scRNA-seq will be used to create a comprehensive reference map of the types and properties of all human cells, as a basis for understanding, diagnosing, monitoring, and treating health and disease.

The usage of scRNA-seq has grown exponentially during these years (Figure 1). However, to analyze scRNA-seq data, novel methods are required. There are many scRNA-seq protocols, including CEL-seq2/C1, Drop-seq, MARS-seq [12, 13], SCRB-seq, Smart-seq/C1, Smart-seq2, STRT-seq, inDrop, Semi-seq [14], SPLIT-seq by using SMART-seq2 and plates [15–17], massively parallel single-cell RNA-seq [18, 19], droplets [20], microfluidics [21–23], 10x chromium system [24], sci-RAN-seq [25], Quartz-Seq2 [26], and the recently developed MARS-seq2.0 [27]. Each protocol has individual advantages and limitations, so their respective usage is often dictated by multiple features, such as biological context, cost, objective, and experimental scale [28, 29]. Ziegenhain et al. and Svensson et al. tested and compared the sensitivity and specificity between protocols that are commonly used today, providing a guide for immunologists before choosing an appropriate approach for addressing a particular research problem [29, 30], followed by Haque et al., who give researchers a further practical guide for their first scRNA-seq practice [31] (Table 1).

3.1.1. Characterizing the Diversity within a Population of Brain Immune Cells. The diversity of gene expression may be the key for the immune system to maintain homeostasis and against infections [32]. scRNA-seq provides a new and effective method to uncover the transcriptional patterns within a population of cells. It has been developed and optimized for diversity analysis in many kinds of cells, which is the main reason why immunologists choose scRNA-seq for study. Recently, deep single-cell RNA sequencing has been applied to study the developmental heterogeneity of microglia [33]. The results displayed an obvious difference in the gene expression of microglia between the adult and early postnatal brain by collecting and analyzing samples from different brain regions. These findings are important to show that the heterogeneity may have potential functions in immune cell development and differentiation. Furthermore, scRNA-seq has been used to target the diversity of glial cells and drug-induced neuroinflammation in cerebral organoids [34]. Previous cerebral organoid studies using scRNA-seq primarily focused on neural populations. Dang and his team turned their attention to glial cells; they confirmed the diversity of glial cells at the organoid level and the neuroinflammatory response within drug-treated organoid astrocytes with single-cell resolution.

3.1.2. Future Directions for scRNA-Seq in Brain Immune System

(1) Revealing Cell Fate Branching Point. While cell differentiation, proliferation, and reprogramming are controlled by complicated gene regulatory networks, each cell determines its own destiny by the integration of a large number of signals, which makes the cell fate difficult to be dissected for technology limits. In addition to heterogeneity discrimination, the appearance of single-cell techniques that can simultaneously detect mountains of molecules in a single cell could also shed some light on this challenge. Chen used single-cell profiling strategy to identify the heterogeneity of CD8+ T cells during acute and chronic lymphocytic choriomeningitis virus infections [7]. In this study, CD8+ T cells showed a different fate trajectory responding to the same virus infection. Notably, the distinct transcriptional profiles of CD8+ T cells and the transcriptional bifurcation timing have been determined with scRNA-seq [7]. Except for immune cells, this novel tracing technology also shows promise in other cell types including basal and myoepithelial cell population in mapping the cell fate and branch points
scRNA-seq is expected to be used to reveal cell fate branch points of the brain immune system (Figure 2).

(2) Classification of the Cellular Composition. Notably, scRNA-seq could provide an unexpected opportunity to systematically identify the cellular specializations of brain cells. To obtain a deeper understanding of the brain, scientists have attempted to establish experimental brain organoid models outside of the context of embryogenesis [36]. Brain organoids originating from different stem cells have been reported from time to time [37, 38], but the significance lies in whether they can generate the similar and rich diversity of cell types appropriate for the brain. This is one of the most important characteristics of the brain that distinguishes it from the rest of the body. However, few studies have attempted to classify the cellular composition of these organoid models, partly due to technical limitations. However, the situation is improving as technology advances. Velasco and her stem cell research team have tested four individual dorsal forebrain organoid models established by four distinct protocols and observed that each model can stably generate a reproducible cellular diversity without the requirement of an embryo [39]. Their important discovery was published in Nature recently. This is a typical case where cutting-edge technologies are used to solve the challenges of scientific research. Another important work worth mentioning is the development of a new algorithm called LIGER, which is designed to delineate shared and dataset-specific cell identity features [40]. As a promising analytical tool, LIGER could combine the single-cell RNA-seq information with DNA methylation profile data and credibly clarify the cell type and single-cell gene expression in specific subsets of human and mouse brain cells.

Single-cell sequencing transcriptomics can also be used to trace the origin and distribution schema of some key molecules and cells of the brain, which comprises thousands of cell types [41]. Undoubtedly, it is extremely useful for understanding the development and diseases of the human brain.
where so many mysteries remain uncovered. Technological advances in scRNA-seq make it possible for researchers to achieve a detailed cell-type analysis of different cells and tissues. Therefore, we can use this advantage of scRNA-seq to reveal immune cell types involved in a specific brain immune response caused by different causes and finally find a treatment target.

(3) Revealing the Interaction between Cells. Apart from the cell-type analysis, scRNA-seq could be used to reveal an interaction between cells. scRNA-seq of the subventricular zone (SVZ) has been separately performed in young (3 months old) and old (28-29 months old) mouse brains to explore how the neurogenic niche, which comprises neural stem cells, as well as many other cell types, changed with age [42]. An expansion of T cells and a decreased proliferation of neural stem cells in old mouse brains were detected in this study. scRNA-seq results indicated that the presence of brain T cells correlated with an inhibition of neural stem cell proliferation, partly because of interferon-γ secretion. This gives us a new way to study the interaction between the peripheral and central immune systems.

scRNA-seq could also be used in conjunction with other popular techniques. It was performed on GFP-labeled inhibitory interneurons (INs) to explore how neocortical projection neurons instruct IN circuit development [43]. The limits of scRNA-seq include the limited cell number and the involvement of single-cell dissociation procedures, during which protease digestion or heating is required and may lead to cell death or other gene expression changes [44–46]. Recently, another promising transcriptome analysis method, single-nucleus RNA sequencing, has been reported without these deficiencies, which can be used not only in fresh tissue but also in frozen tissue [44] (Table 2).

### 3.2. Cytometry by Time-of-Flight

Cytometry by time-of-flight (CyTOF) or mass cytometry is a variation of flow cytometry in which antibodies are labeled with heavy metal ion tags, rather than fluorochromes. It provides researchers, who seek to explore the mysteries of the immune system, a deeper insight than ever before into the complexity of immune cells. Before the appearance of CyTOF in 2009 at Yale, scientists labeled cellular components with flow cytometry or fluorescence-activated cell sorting (FACS) to collect key cell feature data from a variety of cell phenotypes. The commonly available instruments and reagents determine the limitations of flow cytometry and FACS to analyze at most 20 parameters per cell [8]. In contrast to FACS using fluorescent reporters, CyTOF applies metal isotopes, which are detected via the time-of-flight mass spectrometry technique. It has successfully overcome those limitations and expanded the number of parameters that can be detected simultaneously due to the absence of signal overlap between metal isotopes. Besides, it is translated into four to five times more information, making it particularly suitable for valuable patient samples [47]. Most notably, CyTOF allows an investigator to measure an enormous number of extracellular and intracellular targets simultaneously. This advantage also enables the use of clustering techniques to perform the immunophenotypic classification [48]. However, CyTOF is not as perfect as we expect. Poor resolution for some positive and negative stains, harmful effects on cell division, the large expense associated with the machinery and reagents, lower cell numbers, and longer processing time than flow cytometry [54] are all drawbacks of CyTOF, indicating the areas to be improved upon in the future (Table 2).

Since this technology has been applied to explore the role of natural killer (NK) cells in the immune response to West Nile virus, researchers have started expanding the scope of research from West Nile to multiple sclerosis [55, 56], cancer

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Table 1: Comparison of six prominent scRNA-seq protocols. Smart-seq2 is found to be the most sensitive method, and the number of genes detected per cell is the most. Smart-seq is slightly less sensitive than Smart-seq2 and detects almost the same number of genes as Smart-seq2, whereas Drop-seq and MARS-seq have a substantially lower sensitivity and detect considerably fewer genes than others. Interestingly, there are small differences among these six methods in accuracy and dropout rate. CEL-seq2 is the most powerful method at a sequencing depth of 250,000 reads, but SCRIB-seq is the most powerful method with more reads. Drop-seq is the most cost-effective method; SCRIB-seq, CEL-seq2, and MARS-seq are less cost-effective, closely followed by Smart-seq and Smart-seq2. Due to its low library cost, Drop-seq may be desirable when analyzing large numbers of cells with low coverage. On the other hand, Drop-seq requires a relatively large number of cells in its current setting. Thus, if few cells are isolated by FACS, SCRIB-seq, MARS-seq, or Smart-seq2 protocols may be desirable.

| Protocol      | Sensitivity     | Gene number detected | Accuracy       | Dropout rates | Power           | Cost efficiency | Cell volume |
|---------------|-----------------|----------------------|----------------|---------------|-----------------|-----------------|-------------|
| CEL-seq2      | More sensitive  | More genes           | Small differences | Small differences | Most powerful   | More cost-effective | Small cells |
| Drop-seq      | Less sensitive  | Less genes           | Small differences | Small differences | Less powerful   | Most cost-effective | Large cells |
| MARS-seq      | Less sensitive  | Less genes           | Small differences | Small differences | Most powerful   | More cost-effective | Fewer cells |
| SCRIB-seq     | More sensitive  | More genes           | Small differences | Small differences | Most powerful   | More cost-effective | Fewer cells |
| Smart-seq     | Much more sensitive | Much more genes         | Small differences | Small differences | Less powerful   | Less cost-effective | Small cells |
| Smart-seq2    | More sensitive  | High number of genes | Small differences | Small differences | More powerful   | Less cost-effective | Fewer cells |
Finally, we use specialized software to analyze and present the data. PCR, and then, the cDNA sequencing library was prepared. Transcription. After this, cDNA amplification was conducted by droplet-based or plate-based approaches. Next, the transcription of primed RNA was converted into complementary DNA (cDNA) using reverse transcription. After this, cDNA amplification was conducted by PCR, and then, the cDNA sequencing library was prepared. Finally, we use specialized software to analyze and present the data.

3.3. Whole-Genome Transcriptomic and Proteomic Analysis. Whole-genome transcriptomics and proteomics are well-established approaches that can supply information at the DNA, RNA, and protein levels with outstanding coverage and depth of sequencing. They have played an essential role in genome model development and refinement for a broad spectrum of diseases. During this process, whole-genome transcriptomic and proteomic sequencing analysis improvement significantly promotes rapid progress.

3.3.1. Characterizing the Transcriptional Profile. Ciguatera-induced Chronic Inflammatory Response Syndrome (CIRS) can lead to a variety of neurologic deficits with few therapeutic options for the poor understanding of its pathophysiology. Recently, transcriptional profiles of seven important genes that can distinguish patients from controls have been identified by whole-genome microarrays in whole blood from ciguatera-induced CIRS patients [62]. Besides, this approach is appropriate in studies of vectors of rare infectious diseases. In a previous study, whole-genome sequencing has annotated the transcriptional profiles of Ixodes ricinus ticks and vastly extended the DNA and RNA databases for them, paving the way for further deep study of the vector and its hosts [63].

3.3.2. Clarifying the Genetic Phenotype Heterogeneity. Whole-genome transcriptomic technique was used to explore the deep phenotype of asthmatic patients with IL-17-high...
immunity by analyzing bronchial biopsies and blood samples [64]. Additionally, malignancies are aggressive with few therapeutic options and poor prognosis. Recently, whole-genome transcriptomic and proteomic sequencing analysis techniques exhibited the potential for clarifying the phenotype heterogeneity in tumors and the underlying mechanisms which promote cancer evolution. Whole-genome transcriptomic analysis has been carried out on surgically resected hepatocellular carcinoma (HCC) samples by a microarray technique for a novel understanding of the tumor microenvironment transcriptional networks [65]. This whole-genome-scale analysis provides a great potential for the actionable strategy design and precision necessary for targeted therapy in cancer treatment.

3.3.3. Identifying Novel Gene Targets and Developing New Antibiotics. Whole-genome transcriptomic and proteomic sequencing analysis was attempted in bacterial studies and the development of new antibiotics, since the first bacterial genome was sequenced twenty years ago. Using this technique, several potential genes and proteins related to glycopeptide resistance were identified by Alexander et al. [66]. Next, Jessica et al. found that the alterations of the bacteria cell surface genes encoded regulatory proteins, which were thought to be a crucial mechanism through which bacteria attained hypervirulence [67]. With these new genome sequencing analysis techniques that have become available, the similarities and differences regarding the genome sequences of Pasteurella multocida have been determined. Regarding the current data, the authors also found an interesting phenomenon, in which no obvious correlation was found between phylogenetic relatedness and host predilection or disease [68]. Hidekazu and colleagues used the whole-genome sequence to profile RS-1, the only strain of Desulfovibrio magneticus bacteria, and revealed the presence of three key genetic components for magnetotactic formation [69]. With respect to antibiotic discovery, approaches based on whole-genome transcriptomic and proteomic sequencing data have yet to identify genes that are important to bacterial survival and growth, thereby revealing potential targets for novel antimicrobial compound development [49].

The advances in mass spectrometry and transcriptome sequencing technologies make the collection of genomic, transcriptomic, and proteomic information from the same tissue sample available. Integration of this data provides a comprehensive and unprecedented insight into the patient’s gene profile. In the past year or so, whole-genome transcriptomic and proteomic analysis has been applied in the CNS, combined with an exome sequencing approach to identify critical candidate genes contributing to abnormalities in spine development and cortical neuron dendritic and subsequent neurodevelopmental disorders [50]. With the help of an integrated transcriptomic and proteomic analysis, researchers have validated and improved the present existing

| Methods                        | Molecules analyzed | Parameters | Advantages                                                                 | Disadvantages                                                                 | Refs  |
|-------------------------------|-------------------|------------|-----------------------------------------------------------------------------|-------------------------------------------------------------------------------|-------|
| scRNA-seq                     | mRNA, surface protein (CITE-seq only) | <20        | (1) Incorporates surface protein and mRNA (2) Limited specialized equipment required | (1) Labor-intensive, lowest numbers (2) Protease digestion heating requirement | [43–45] |
| CyTOF                         | Protein, glycosylation epitopes, lipids, phosphorylation | >40        | (1) Higher simultaneous parameters (2) Permit using clustering to classify cell group (3) Identify cell populations readily without the need to stain with classical markers (4) No spectral overlap and other fluorescence-associated complications (5) Adjust the detector easily | (1) Lower throughput and longer run time than flow cytometry (2) Removal of photometric measurement precludes measurement of forward and side scatter (3) Poor resolution for some stains (4) Cannot isolate live cells (5) Expense of machines and reagents (6) Difficult to detect low abundance antigens (phycoerythrin or allophycocyanin) (7) Ion sensitivity decreases over time | [46–48] |
| Whole-genome transcriptomic and proteomic analysis | DNA, RNA, protein (high-quality proteomic data) | >8000 genes | Supply information at DNA, RNA, and protein levels | (1) Poor sensitivity of the spectrometers (2) High incorrect incidence for new peptide hits (3) Intrinsic biophysical features that cause some peptides undetectable (4) Largely dependent on computational technologies | [49–53] |
annotation of the zebrafish protein-coding genome [51]. Furthermore, several diseases have been investigated at whole genomic, transcriptomic, and proteomic levels, making the therapy strategies more promising. Using these advanced technical tools, multiple candidate genes and signaling pathways potentially involved in Dupuytren’s disease have been reported, validated, or proposed, which may help to elucidate the pathogenic mechanisms underlying Dupuytren’s disease [52].

Nevertheless, like all other popular areas of research, the current state of whole-genome transcriptomic and proteomic analysis is not without flaws, including the poor sensitivity of the spectrometers, the increased false positives for new peptide hits, and particularly the intrinsic biophysical features that cause some peptides to remain undetectable [53]. In addition, these techniques are extremely dependent on technologies that are used to enable sensitive and large-scale proteome profiling, like the most dominant technology, tandem mass spectrometry (MS/MS) [70]. Another promising technology that they use is ribosomal footprinting, which offers exceptionally valuable information at the translational product level and acts as a bridge between transcriptomics and proteomics data. However, it is technically challenging for many laboratories at present (Table 2).

Although whole-genome transcriptomic and proteomic analysis has been broadly applied in many areas, notably until now, there are few relevant studies found in brain immune research. The reason is likely due to the complexity of the brain structure, making these novel techniques promising in exploring the potential mechanisms of this pathophysiological condition.

4. Conclusions and Expectations

To adapt to the environment and fend off external threats, creatures from prokaryotes to humans have strong immune systems, and these immune systems have developed highly specific types of immune cells to prevent and eliminate diseases. The hottest single-cell technology is critical to understanding how the immune system generates potential responses to many different pathogens.

This review summarizes existing novel technologies and discusses their advantages and limitations and explores how these approaches can deepen our understanding of the immune response in neurological diseases. Today, major breakthroughs are scarce in the research of brain injury, including apoptosis [71, 72], inflammation [73, 74], and oxidative stress [75, 76]. Immunology is rapidly gaining attention recently [77]. Additionally, major strides have been made in human science and technology, shedding light on a deep understanding of the immune response after brain injury. However, few neuroscientists have applied these techniques to explore the impact of the immune system on brain repair. It is worth waiting to see how researchers will use these powerful tools to systematically address this issue. To achieve this, information from several fields, such as genomics, transcriptomics, and proteomics at the cellular and organism level, as well as database and computational technologies will be required. Genes and their products do not work independently. Without a doubt, further closer integration of genomics, transcriptomics, proteomics, and relevant promising approaches will be more important than ever before, which can influence the way in which researchers properly understand the brain immune response and, moreover, can sway the way in which other complex neurological problems are solved.

Disclosure

The funders had no role in the design of the study; in the collection, analyses, or interpretation of the data; in the writing of the manuscript; or in the decision to publish the results.

Conflicts of Interest

The authors declare no conflict of interest.

Authors’ Contributions

LL wrote the initial draft and prepared the figures and tables. CL, JK, XL, and ZL revised the draft; FX and JZ recommended the structure for the paper, which substantially advanced the draft.

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