Multiple sclerosis: getting personal with induced pluripotent stem cells

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Human induced pluripotent stem (iPS) cells can be derived from lineage-restricted cells and represent an important tool to develop novel patient-specific cell therapies and research models for inherited and acquired diseases. Recently, patient-derived iPS cells, containing donor genetic background, have offered a breakthrough approach to study human genetics of neurodegenerative diseases. By offering an unlimited source of patient-specific disease-relevant cells, iPS cells hold great promise for understanding disease mechanisms, identifying molecular targets and developing phenotypic screens for drug discovery. This review will discuss the potential impact of using iPS cell-derived models in multiple sclerosis (MS) research and highlight some of the current challenges and prospective for generating novel therapeutic treatments for MS patients.

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Facts

- Genetic and environmental factors are believed to be the underlying causes of the majority of autoimmune diseases such as multiple sclerosis
- Animal models only partially recapitulate pathogenetic features of autoimmune diseases
- iPS cells have the capability to differentiate into all cell types of the human body
- iPS cells represent an early stage of disease development

Questions

- The pathogenetic events involved in multiple sclerosis development and progression are still not completely understood
- Which pathogenetic events are involved in multiple sclerosis development?
- Are epigenetic aberrancies crucial for autoimmune diseases onset?
- Do iPS cells have the capability to model autoimmune diseases?
- Can iPS cells provide novel pathogenetic insights in autoimmune diseases?

Multiple sclerosis (MS) is a progressive, inflammatory, demyelinating central nervous system (CNS) disease affecting mostly young adults.1 Despite the real cause(s) remains largely unknown, MS has been conventionally classified as an autoimmune inflammatory disease affecting the white matter and only recently demonstrated to affect the grey matter as well.1 MS development has been associated with a genetic predisposition, which in concert with environmental factor exposure2 such as viral infections,3,4 vitamin D deficiency,5 and other factors, is responsible for disease initiation.6 Initial lesions are frequently associated with a perivascular inflammation that is also considered the origin of the blood–brain barrier breakdown found in MS patients. Thus, MS is characterized by chronic leukocytes infiltration of CNS and by self-limiting attacks to glial cells, ultimately leading to a severe neuron demyelination. One of the early features of MS is the presence of neurons having few layers of myelin rather than the usual 30 layers of compact myelin with a consequent reduction of the action potential conduction along nerves. Furthermore, recovery from acute inflammation results often in ion channel damage, which in absence of defined Ranvier nodes become abnormally distributed along the axons, concurring to the failure of efficient signal conduction. Importantly, myelin destruction followed by neuronal injury is responsible for both long-term disability and cognitive impairment7 in MS patients and nowadays, all current treatments focus in reducing or blocking the autoimmune reaction.

Despite the considerable resources invested in MS research, a significant number of open questions regarding pathogenesis, disease subtypes and response to therapy are still in need to be elucidated. Animal models of autoimmune

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Abbreviations: iPS cells, induced pluripotent stem cells; MS, multiple sclerosis; CNS, central nervous system; ES cells, embryonic stem cells; EBs, embryoid bodies; DMRs, differentially methylated regions; IL, interleukin; TYK2, tyrosine kinase 2; ACCN1, amiloride-sensitive cation channel 1, neuronal; KIF1B, kinesin family member 1B; miRNAs, microRNAs; SMA, spinal muscular atrophy

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...demyelinating diseases, mimicking MS phenotype, have been so far utilized with the hope to find effective treatments for MS. However, these animal models have failed to produce further pathogenetic insights of the disease, likely owing to the profound differences between the animal models and the human disease. The recent discovery that somatic cells can be reprogrammed to a pluripotent stem cell-like state has provided an important tool to study neurodegenerative disease in a controlled environment, including MS. Induced pluripotent stem (iPS) cells represent an early stage of disease development, and their use has the potential to identify specific disease pathways prior, during and after disease development. In addition, the possibility to obtain neurons and leukocytes with the same genetic background of MS patients can provide a deeper understanding of the genetic and epigenetic alterations contributing to the disease establishment. Recapitulating the human MS phenotype in vitro by using iPS cells might represent the rationale for the development of a drug screening approach to identify novel patient-customized targeting treatments.

**Induced Pluripotent Stem Cells**

Among stem cells, human embryonic stem (ES) cells have been considered to hold greatest promises in biomedical science owing to their capability to differentiate into all the germ layer derivatives and given their potential as treatment in degenerative diseases. During the last few decades, a large amount of scientific efforts have been put in the development of functional equivalents hES-like cells for scientific and clinical purposes in order to overcome the ethical issues related to the hES use. Somatic cell nuclear transfer and cell fusion have been showed to induce reprogramming of differentiated cells to a pluripotent state; however, both techniques were highly inefficient for humans as well limited in number to be used on a large scale for disease modeling or regenerative medicine. Further, these methodologies did not solve the issues of immunological rejection of the transplanted allogeneic tissues derived from the pluripotent parental cells or the ethical issues relating to destruction of human embryos.

The landmark discovery that pluripotent stem cells can be directly derived through the ectopic expression of defined factors opened a new frontier for regenerative medicine especially for novel disease modeling and drug screening methodologies. Overexpression of OCT4 and SOX2 in combination with KLF4 and c-MYC, has been shown that differentiation of pluripotent stem cells is achievable in vivo by way of teratoma formation. This innovative approach allowed the isolation of several blood elements including hematopoietic stem/progenitor cells capable of multilineage reconstitution when transplanted in immunocompromised mice. This protocol represents an alternative route to derive specific cell types from pluripotent stem cells, whereas in vitro differentiation is not a feasible approach. Another aspect to consider when handling iPS cells is the impact of potential genetic and epigenetic alterations, occurring during the reprogramming process, on the differentiation potential. Although iPS cells show a normal

![Figure 1](Image)  
**Figure 1**  
Somatic cells can be reprogrammed to an ES-like state through the overexpression of four transcription factors. Pluripotent stem cells derived through this approach can differentiate to the three germ layer derivatives by the way of embryoid body formation (EBs).
karyotype after the reprogramming process, it has been reported that continuous passing of these cells is associated with the acquisition of chromosomal abnormalities (karyotype 46,XY,t(17;20) (p13;p11.2)) starting approximately at passage 13.30 Recent studies have demonstrated significant reprogramming variability among iPS cell lines, including somatic memory or aberrant reprogramming of DNA methylation.31,32 In particular, Lister and colleagues have identified differentially methylated regions (DMRs) occurring during reprogramming by comparing iPS with ES and somatic cells. Half of the identified DMRs were related to incomplete reprogramming while the other half was deemed as epigenetic errors as they were absent in either somatic cells or ES cells. These methylation errors can be potentially inherited by the iPS-derived differentiated cell types; however, the impact of such epigenetic heterogeneity on iPS cell-differentiation capabilities as well as an approach to reduce it have not yet been fully elucidated.

**iPS Cells Impact MS Genetics**

The possibility to generate iPS cells from individuals affected by several diseases with a genetic component33 has attracted a world-wide attention, initiating a driving force to create real patient-customized disease models, novel drug screening platforms and eventually regenerative medicine. Although there are high expectations for future applications of iPS cells to repair damaged organs including the CNS, the most compelling and important concerns/limitations remain the feasibility to produce ‘large-scale’ patient-derived cells carrying specific genetic susceptibility and to subsequently analyze these cells under controlled conditions in the laboratory. Recently, fibroblasts from MS patients have been reprogrammed to pluripotency34 making an important step toward MS research. Indeed, multiple sclerosis iPS cells are able to differentiate to all germ layer derivatives, including oligodendrocytes, astrocytes and functional neurons. Many iPS lines generated from several pathological conditions are characterized by specific genetic driving mutation.33 In contrast, MS is a multifactorial disease caused by a complex interaction between environment and genetic susceptibility thereby contributing to the pathological heterogeneity.35

A growing body of evidences suggests that the genetic component has a crucial role in the disease development36 (Table 1). Despite the MHC loci still represent the major dominant risk region for MS development,37,38 many other non-MHC genetic variants involved in MS pathogenesis have been recently identified.39 Interestingly, the majority of the immunological genes identified to have a role in the disease onset are known to be important for T-cell responses, including cytokine receptors for interleukin-7 (IL-7) and IL-2,40,41 and signal transducers such as tyrosine kinase 2 (TYK2) for type I interferons, IL-10 and IL-12.7 Genetic variants of adhesion molecules such as CD58 and CD226 are also demonstrated to be associated with MS development because of its involvement in abnormal stimulating signals.42 Interestingly, neurological gene variants have also been found associated with MS, including aminolide-sensitive cation channel 1, neuronal (ACCN1)43 and kinesin family member 1B (KIF1B).44 Although

| Table 1 Representative key genetic variants found in multiple sclerosis patients |
|------------------|-----------------|-----------------|
| **Gene** | **Proposed function** | **Reference** |
| **Immunological genes** | | |
| HLA-DR | Antigen presentation | Lincoln et al.37 |
| HLA-A | Antigen presentation | Fogdell-Hahn et al.38 |
| CXCR5 | Chemokine receptor | Sawcer et al.45 |
| IL2RA | Cytokine receptor | Hafler et al.46 |
| IL7R | Cytokine receptor | Sawcer et al.45 |
| IL12A | Cytokine | Sawcer et al.45 |
| IL12B | Cytokine | Sawcer et al.45 |
| STAT3 | Signal transduction | Ban et al.41 |
| TYK2 | Signal transduction | Sawcer et al.45 |
| MALT1 | Signal transduction | Sawcer et al.45 |
| CD40 | Co-stimulatory molecule | Sawcer et al.45 |
| CD86 | Co-stimulatory molecule | Sawcer et al.45 |
| VCAM | Adhesion molecule | Sawcer et al.45 |
| CD58 | Adhesion molecule | De Jager et al.42 |
| **Neurological genes** | | |
| ACCN1 | Neuronal pH-sensitive ion channel | Bernardinelli et al.43 |
| KIF1B | Axonal transport | Aulchenko et al.44 |
| ALK | Protein tyrosine kinase receptor; brain development | Hafler et al.46 |

the role of the ACCN1 channel is poorly understood, KIF1B encodes a protein involved in axonal transport of mitochondria and synaptic vesicle precursors. Although it is reasonable to hypothesize a contribution of these genes to the inflammatory process and neuronal degeneration observed in MS, the functional role of these factors during the disease development still requires to be specifically investigated. The advent of iPS cells makes it now possible to investigate the impact of genetic variants on specific cell population behavior and survival. In particular, generation of several neural population with a defined genetic MS background may allow functional evaluation of specific genetic variants and polymorphisms by unraveling their role in the disease establishment. A similar approach has the potential to shed light on the participation of several environmental insults, such as reactive oxygen species or abnormal cytokine levels, on apoptosis or cell death of specific neuronal population and to recapitulate at least partially the inflammatory environment of MS. Further, by enabling studies on the differentiation capability of MS-iPS-derived neural stem/precursors to terminally differentiated neurons and glial cells, this strategy will allow us to understand whether specific genetic variants interfere with self-renewal and regeneration of damaged CNS tissues in MS patients.

In summary, the use of iPS cells might unravel specific pathogenetic aspects of MS initiation and development that, up to date, remain largely unknown. Understanding the function of specific genetic variants, detected in neural and immunological population of MS patients, is critical to pave the way for the discovery of novel therapeutic targets to tackle this complex debilitating disease.

**iPS Cells Impact MS Epigenetics**

In the last few years, a large amount of studies showed that the interplay between environmental factors and individual genetic susceptibility is likely to produce a pathogenetic predisposition to MS.6 Currently, numerous studies point at
the role of epigenetic aberrancies to both disease development and progression.\cite{46}

Epigenetic modifications are heritable changes affecting gene expression without altering the DNA sequence. The molecular changes underlying epigenetics modifications include DNA methylation and histone modifications. The epigenetic modifications are important not only for tissue development but also for homeostasis and for the establishment of cellular identity. Transcriptional activation is associated with the presence of both lineage-specific transcription factors and of activation marks on lysine and arginine residues of histone tails. Gene repression instead is achieved by repressive marks on amino-acid residues of histone tails.\cite{47}

DNA methylation is linked to gene silencing and it is considered more likely an irreversible phenomenon.\cite{48}
Non-coding RNAs operate at both transcriptional\cite{59} and post-transcriptional\cite{56,60} level by finely regulating expression of specific target genes. All the epigenetic modifications are affected by environmental signals and their dysregulation can lead to many diseases including MS.\cite{51} Patients with MS have been found to share specific epigenetic modifications (Table 2), yet the functional role of such aberrancies in the disease development is not completely understood.

Recently, DNA methylation has been investigated in CD4+ peripheral blood lymphocytes from three twin pairs discordant for MS.\cite{52} Two genes TMEM and PEX14 were found to be differentially methylated. Furthermore, Liggett $et$ al.\cite{53} analyzed the DNA methylation pattern of 56 genes previously shown to be associated with cancer development using cell-free plasma DNA of MS patients. Interestingly, 15 out of 56 genes including PAX5, TP73 and FAS were found to be differentially methylated suggesting the potential use of different T-cell subpopulations to further investigate the role of DNA methylation in MS.

Nucleosomal histone tails are target of many enzymatic processes including acetylation, methylation, phosphorylation, sumoylation, citrullination and ubiquitination. Acetylation of lysine in position 9 and 14 and tri-methylation of lysine and arginine residues H3 histone tails in MS patients. This phenomenon also known as citrullination has been associated with protein instability and might be contributing to oligodendrocytes apoptosis\cite{54,55} and T-cell activation.\cite{55}

MicroRNAs have been also found differentially expressed between MS patients and healthy individuals. Using high-throughput miRNA profiling, key miRNAs involved in MS development have been identified in several tissues including whole blood, peripheral blood mononuclear cells and lymphocytes.\cite{56,57} Interestingly, few miRNAs involved in repression of genes associated with T-cell activation such as miR-17 and miR-20a\cite{57} were found significantly decreased (Table 2). Therefore, downregulation or upregulation of RNA molecules involved in the immune response could represent a key second hit during MS development.

Epigenetic changes have a pivotal role during nuclear reprogramming and are fundamental for pluripotency acquisition.\cite{58} From this prospective, nuclear reprogramming offers an ideal model to study the role of specific epigenetic abnormalities in MS development and progression. The generation of genetically matched but epigenetically distinct cells might help understanding the involvement of epigenetic alterations in the disease pathogenesis. Recently, it has been shown that the possibility to reset epigenetic abnormalities in cancer by nuclear reprogramming,\cite{55,59,60} leading the cancer epigenome to an ES-like ground state.

The same approach can be exploited in MS in order to reactivate immune response key factors poisoned by DNA methylation. This approach might allow the functional evaluation of epigenetic alteration role in MS development. Reactivation of critical genes of immune response regulation such as FAS and TP73 might lead to understand whether ‘apoptosis escaping’ driven by epigenetic mutations concurs to generate self-reactive clones after inflammation and whether those types of epigenetic abnormalities represent a precipitating pathogenetic event during autoimmune disease development (Figure 2).

### Table 2: Representative key epigenetic modifications found in multiple sclerosis patients

| Gene | Epigenetic abnormality | Source | Reference |
|------|------------------------|--------|-----------|
| TMEM | DNA methylation | CD4+ cells | Baranzini $et$ al.\cite{52} |
| PEX14 | DNA methylation | CD4+ cells | Baranzini $et$ al.\cite{52} |
| CDKN2B | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| FAS | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| MCI | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| MDS1 | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| PGK1 | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| TP73 | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| CDH1 | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| PAX5 | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| SYK | DNA methylation | Cell-free plasma DNA | Liggett $et$ al.\cite{53} |
| Not specified | H3 citrullination | PBMCs | Mastronardi $et$ al.\cite{54} |
| TNFalpha | H3 citrullination | Whole blood | Sharma $et$ al.\cite{56} |
| miR-17 | Downregulation | Whole blood | Cox $et$ al.\cite{56} |
| miR-20a | Downregulation | CD4+ cells | Jr Ode $et$ al.\cite{56} |
| miR-17-5p | Upregulation | CD4+ cells | Jr Ode $et$ al.\cite{56} |
| miR-155 | Upregulation | T-regulatory cells | Jr Ode $et$ al.\cite{56} |
| miR-25 | Downregulation | T-regulatory cells | Jr Ode $et$ al.\cite{56} |
| miR-106b | Downregulation | T-regulatory cells | Jr Ode $et$ al.\cite{56} |
Toward Pharmaceutical Applications of iPS Cells

The comprehension of how genetic variation contributes to disease pathogenesis is inevitable for the development of preventive strategies and new generation treatments. Historically, gain- and loss-of-function animal models have been used for this purpose; however, several disease phenotypes failed to be reproduced in these models. Furthermore, the limitation in obtaining human tissues, other than blood, has been so far an obstacle for conducting analysis and relevant research for specific diseases. Thus, the advent of iPS cells has represented a turning point to overcome the mentioned limitations. The possibility to use pluripotent cells derived from patients able to differentiate into cells or tissues not accessible otherwise (e.g., neurons, glial cells, etc) has initiated a new research field with the potential to unravel specific pathogenetic mechanisms, which are still largely unknown. In addition, owing to the unlimited self-renewal capabilities, iPS cells become a suitable tool for drug screening or toxicology studies. Furthermore, iPS cells can mimic disease models in so-called ‘disease in a dish’ system. In order to obtain a specific ‘disease in a dish’ phenotype, two aspects must be satisfied: (1) to derive pathological relevant cell populations and (2) to recapitulate key aspects of the disease onset. This approach might even more benefit from the application of three-dimensional (3D) cell culture model system. Traditionally, two-dimensional (2D) cell culture systems have been considered the chosen and most simplistic tools to reproduce ‘disease in a dish’ for drug discovery and screening purposes whereas ‘in vivo’ models have been preferred for the efficacy and safety assessment before proceeding to clinical trials. Unfortunately, both approaches display several limitations: 2D cultures are unable to mirror the physiological condition of the native tissue of study, thus impairing therapeutics screening as shown by the high drug failure rate (~95%).

Although nuclear reprogramming appears to be a very inefficient process, to date many iPS lines from patients affected by genetic disease including Duchenne muscular dystrophy, Down’s syndrome, Parkinson’s and so on have been derived. Among genetic diseases from which iPS cells have been derived, spinal muscular atrophy (SMA) looks to be one of the most promising to generate a valuable disease modeling. SMA is a neurodegenerative genetic disease in which motor neurons degenerate bringing the young patients to paralysis and often death. iPS cells derived from SMA fibroblasts were capable to differentiate normally to neural tissues and motor neurons. However, motor neurons were selectively lost in the neural culture recapitulating the patient-specific SMA phenotype. Interestingly, valproic acid or tobramycin treatments were capable to increase the levels of SMN protective protein. These findings prove that generation of a valuable disease in a dish model might aid the identification of novel compounds and pathogenetic mechanisms in the disease initiation and development.

Heterogeneity of human diseases has never been recapitulated in animal models. Generation of iPS lines from patients’ samples with same disease but different outcome has the potential to shed light on the mechanisms driving the severity of a specific disease. For MS, unraveling the pathogenetic events behind the disease subtypes represents a major goal for understanding and generating the most appropriate and efficacious treatment for that disease phenotype. In particular, derivation of iPS cells from patients affected by different types of MS might facilitate identification of genetically distinct disease subtypes (Figure 3).

Understanding drug toxicity before starting a clinical trial is one of the greatest challenges in pharmaceutical research. A reliable and accessible platform to predict toxicity in vivo will offer two major advantages: (1) it will lead to more efficient generation of safer compounds and (2) it will decrease the costs for drug development. Animal models have been extensively used for toxicity purpose, however, differences between human and murine tissue entail a poor accuracy in toxicity prediction. iPS cells capable to differentiate to tissues often target of drug toxicity (e.g., cardiac, neural and hepatic...
tissues) represent now an important tool to dissect pre-clinical drug toxicity. Pluripotent stem cell libraries derived from skin of patients for whom that drug is being developed represent an important resource to predict accurately toxicity in a specific cohort of patients. Finally, the possibility to derive disease subtype iPS cells will help in developing patient-customized treatments. For MS case, this approach will allow for a timely identification of patients that would respond adversely or favorably to a specific compound improving and accelerating the drug development process.

**Concluding Remarks**

The major finding that somatic cells can artificially reach a pluripotent stem cell-like state holds new promises for discovery of novel therapeutic treatments. Recent studies warn some caution for the use of iPS cells for clinical applications due to both the use of exogenous DNA and for the presence of important epigenetic alterations found in several iPS lines. On the other hand, it is clear that the most promising application of iPS cells resides in disease modeling and drug screening. Establishment of the ‘disease in a dish’ using disease-specific iPS cells will lead to the identification of pathogenetic mechanisms that are still poorly understood. iPS cells promise to overcome all the limitations associated with animal models used to recapitulate specific disease phenotypes, offering an unlimited source of pathological tissues involved in specific disease development. Combination of exploratory genomic approaches such as new generation sequencing with functional experiments through iPS cells will elucidate key pathways in the initiation and development of many genetic disorders including neurodegenerative diseases.

Despite many advances in MS field have been achieved in the last few years, a number of open questions regarding disease initiation, disease subtypes and response to therapy still need to be answered. The generation of MS phenotype in a dish represents the biggest challenge to overcome all the limitations associated with the autoimmune-mediated demyelinating animal models used so far. iPS cells derived from MS patients will make accessible neural populations that so far have been poorly analyzed owing to inaccessibility of the brain tissues. Analyzing neural populations with the specific patient’s genetic background will undoubtedly shed light on disease-pathogenetic mechanisms promoting the generation of new effective treatments for this debilitating disease.

**Conflict of Interest**

The authors declare no conflict of interest.

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