INTRODUCTION

In late December 2019, the first cases of pneumonia with an unknown origin were reported in Wuhan, China. The pathogen was identified as a novel coronavirus that has a similar molecular structure to SARS-CoV-2. Since then, not only in China but throughout the world, the number of COVID-19 patients has risen dramatically. Coronaviruses (CoVs) are part of the Orthocoronavirinae subfamily of the Coronaviridae family of the Order Nidovirales. The Orthocoronavirinae subfamily is subdivided into four genera that include Alphacoronavirus (-CoV), Betacoronavirus (-CoV), Gammacoronavirus (-CoV), and Deltacoronavirus (-CoV). The World Health Organization reports that the outbreak of the deadly virus had been noted almost in all the countries worldwide. Newly no standard therapies are available to combat the situation and this remains the major challenge for healthcare professionals to provide effective treatment against the life-threatening condition. As a result, concerted efforts are required to develop safe and effective COVID-19 therapies, particularly for severe instances. A potential regenerative medicine method using the infusion of stem cells for the treatment of lung disorders has been reported. The recent study finds that infusing stem cells in the affected individuals showed promising results as the stem cells with their cytoprotective and pro-antigenic properties remain as an advanced treatment strategy in severe lung diseases to overcome this pandemic period. This review attempted to explore the immunomodulatory characteristics of Mesenchymal Stem Cells (MSCs) and how these properties make them beneficial for the treatment of SARS-CoV-2 patients.

1. METHODOLOGY

A literature search was conducted via online search platforms such as PubMed, Embase, and various resources with the help of keywords to identify recent research relating to the review's goal of analyzing the relevance of stem cells in battling SARS-CoV-2.

2. RESULTS AND DISCUSSION

NEW TREATMENT APPROACH INVOLVED IN COVID-19

2.1 Nanotechnology-Based Treatment Techniques

COVID-19 treatments rely heavily on nanotechnology. The effectiveness of nanotechnology in SARS-CoV-2 treatments is
COVID-19 patients. This is because the simplicity with which these sources may be extracted and the amount collected make them ideal for both experimental and therapeutic uses. Many MSCs have recently been generated from novel sources, including menstrual blood and endometrium. Bone marrow, adipose tissue, umbilical cord blood, and endothelial progenitor cells are the most popular types of MSCs used for lung disease treatment. Despite tremendous progress in the field of stem cell-based treatment, the primary constraints of this therapeutic strategy immunogenicity, restricted cell supply, and ethical issues, have yet to be resolved. MSCs have gotten a lot of interest because of their source potential, high proliferation rate, less invasive technique, and lack of ethical concerns when compared to other therapies they have a significant advantage. COVID-19 may trigger a destroying immune overreaction in the body and it produces large amounts of inflammatory factors, causing a cytokine storm including an overproduction of immune cells and cytokines.

2.4 Mechanism of MSCs

T-cells are activated during inflammation through several cell signaling processes. The initial signal that activates lymphocytes is mediated by antigen-specific T-cell receptors (TCR), whereas the second phase, known as co-stimulation, is independent of the TCR yet critical for allowing complete immune response activation and avoiding energy. MSCs can repress T-cells at both the primary and secondary activation phases by signalling through soluble substances such as cytokines and growth factors, as well as through mechanisms involving direct cell-to-cell interactions. IFN-promotes indole amine 2, 3-dioxygenase (IDO) activity in MSCs via the JAK/STAT signaling pathway, which converts tryptophan to kynurenine, which inhibits many cells and reduces the inflammatory response. Apoptosis of pro-inflammatory T-cells was found after increasing IDO activity by tryptophan deprivation due to a generalized decrease in cellular energetics, as evidenced by increased production and accumulation of released kynurenines.

Surprisingly, more than 30 soluble molecules have been found to promote MSC immunomodulation potential to control T-cell activation and proliferation. Human MSCs, in contrast to other species, generate IDO to inhibit T-cell proliferation and induce T-cell energy, which is characterized by a lack of proliferation and a reduction in cytokine output. These findings highlight the fact that MSCs have the greatest immune-inhibitory effect on T-cell proliferation. Cell-to-cell interactions are also important in MSC-mediated immunomodulation. MSCs do not express co-stimulatory molecules such as CD40, CD80, CD86, CD134, and CD252 even after being exposed to inflammatory signals. Furthermore, MSCs can inhibit T-cell proliferation by expressing CD39 and producing adenosine, which activates the adenosine A2 receptor (A2A) on the surface of lymphocytes. It is worth noting that MSCs from diverse origins do not have the equivalent immunoregulatory ability.

In research comparing MSCs from various sources, Warton Jelly produced MSCs shown to be the most efficient in immunosuppression. According to research, activation of Toll-like receptors (TLRs) plays a critical role in immunomodulation mediated by cell-to-cell interaction as well as MSC-secreted soluble substances. TLRs have been discovered to have a crucial function in the innate immune system for recognizing pathogen-associated molecular patterns (PAMPs) and activating initial responses against pathogens. It is important to note that MSCs from different sources do not have the same immunoregulatory capacity. TLR3 and TLR4 ligation has been shown to boost the

Dependent on selecting the correct nanocarriers for the right medication candidate. Furthermore, nanocarriers circumvent the limits of current antiviral treatments. In COVID-19, nanoparticle-assisted regulation of antigen-presenting cells (APCs) is critical for vaccine development. Even though antimicrobial medicines such as chloroquine, rediiver, and favipiravir had shown promising results against SARS CoV-2, some patients had severe adverse effects. The association of nano-carriers offers the essential environment for these medicines to work without adverse effects. During the early phases of COVID-19, nano-macrophage mimetic systems neutralize viral activity and, afterward, decrease inflammation. The consequences of COVID-19 hematological pathology can be mitigated by Nano-Erythrocyte mimic medication delivery. Combination medication treatments are important in the treatment of COVID-19 because of their low side effects, low dose amount, and multiple targeting. Nanocarriers have been identified as promising candidates for multi-drug delivery, which is beneficial for combination medication therapy. Nanoparticle antigen delivery to dendritic cells enhances T-cell immunity. The use of nanomaterials such as Nanospheres, nanocarriers, liposomes, lipid nanoparticles, nanopages, and dendrimers can improve targeted medication delivery for the treatment of COVID-19. In addition to nanotechnology, stem cell therapy plays an important role in the treatment of COVID-19 illness.

2.2 Stem cell therapy and various methods of stem cell therapy

The term "stem cell" refers to a wide range of different cell types. They are either completely undifferentiated or partially differentiated cells that may proliferate continuously to generate more of the same kind of cell and can develop into a variety of cell types. The modifiers "embryonic" and "adult" are commonly used to distinguish stem cells based on the developmental stage of the animal from which they originate, but these terms are becoming obsolete as new research has discovered how to convert fully differentiated adult cells back into embryonic stem cells and, conversely, adult stem cells, more correctly termed "somatic" stem cells meaning "from the body", back into embryonic stem cells. Adult stem cells such as hematopoietic stem cells (MSCs) from adipose tissues have an excellent capacity to repair tissues because they can proliferate for longer periods while remaining undifferentiated and then differentiate into various types of cells. These cells are more commonly used in therapeutic procedures due to several ethical and regulatory constraints. MSCs' immunomodulatory properties have been reported to be one of the major elements imparting therapeutic benefits during the process of lung repair and regeneration in a variety of pathological conditions including bronchopulmonary dysplasia, asthma, acute lung injury, chronic obstructive pulmonary disease, and idiopathic pulmonary fibrosis. Several attempts are being made in this direction.

2.3 Outlook of mesenchymal stem cell therapy

MSCs can regenerate or renew cells on their own and they are differentiated by several lineages. MSCs can also be isolated from various adult tissues, such as bone marrow (BM), peripheral blood (PB), and adipose tissues (AT) (such as abdominal fat, infrapatellar fat pad, and buccal fat pad), as well as neonatal birth-associated tissues, such as placenta (PL), umbilical cord (UC), Warton jelly (WI), amniotic fluid (AF), and cord blood (CB), and then stored for future. As a result, it appears that MSC-based therapy, or at least a mix of treatments, might be an attractive option for clinical trials in

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immunosuppressive characteristics of MSCs in the absence of IFN-γ by enhancing tryptophan breakdown, which leads to increased kynurenine synthesis, which is known to be catalyzed by IDO and triggered by autocrine interferon [48]. MSCs may be polarized into two homogeneous phenotypes by downstream TLR signaling, known as MSC1 and MSC2. Lowered TLR4 against exposure polarises MSC1 toward pro-inflammatory activity via processes that are not completely understood. In contrast, for phenotype 2, MSCs can be polarised by TLR3 activation, and as a result, MSCs inhibit the immune response [49]. To minimize negative effects while utilizing MSCs as an anti-inflammatory treatment, it has been proposed to activate particular TLRs under culture conditions before using the cells in vivo [50]. Human leukocyte antigen (HLA) class I molecules are expressed at low levels in MSCs. This characteristic of stem cells in general aids them in evading natural killer (NK) cells. Furthermore, MSCs and other stem cells lack the expression of HLA class II molecules as well as costimulatory molecules such as CD40, CD40L, CD80, and CD86, which are important in the activation of T-lymphocyte-mediated immunological responses [6, 42, 43, 51]. MSCs with “immune privileged” properties, in addition to being no immunogenic, that is, do not elicit a rejection response due to less strong pro-inflammatory IFN-induced HLA-II expression, making them an appealing therapeutic tool for treating COVID-19 [52].

2.5 Mesenchymal stem cell therapy treating other viral infectious diseases

When compared to highly differentiated cells, MSCs are usually resistant to viral infections. This ability is provided by the presence of IFN-stimulated genes (ISG), which can target at many stages of the viral cycle, preventing viruses from crossing the cell membrane, blocking the endocytic route, mRNA transcription, nuclear import of mRNAs, genome replication, protein translation, virus assembly and release [36, 53, 54]. PMAIP1, ISG15, IFI6, IFITM, SAT1, p21/CDKN1A, SERPINE1, and CCL2 are among the ISGs expressed in MSCs [36, 54]. These ISG work by inhibiting several viral infections in vitro, including dengue, Ebola, SARS, and influenza-A [53]. Wu and colleagues (2018) demonstrated the significance of certain MSC-derived-ISGs by silencing p21/CDKN1A expression, which resulted in enhanced MSC susceptibility to chikungunya virus infection. 74 Stem Cell Rev and Rep (2021) 17:71–93 IFITM3 silencing, however, made MSCs more vulnerable to yellow fever and zika virus infection [56, 55]. They discovered that IDO expressing MSC stimulated with IFN-γ in vitro decreased HIV-1/2 virion production. The authors hypothesized that this impact was caused by tryptophan deficiency, which inhibits the production of emerging viral proteins [56].

MSCs have been shown in pre-clinical trials to be helpful in a variety of viral infections [57]. Chan and colleagues found that influenza A/H5N1 infection induced alveolar fluid clearance (AFC) and increased alveolar protein permeability (APP) in human alveolar epithelial cells (AEC) in vitro, both of which are linked with acute lung damage [58]. Human BM-MSCs inhibited or decreased this impact, which was mediated in part by angiotensin-1 (Ang1) and keratinocyte growth factor (KGF) production in the co-culture experiment. In elderly A/H5N1-infected mice, MSC therapy on day 5 post-infection enhanced survival and body weight, as well as improved lung pathology and histopathology scores [57].

2.6 The mechanism behind mesenchymal stem cell therapy treating covid-19

China, the United States, Jordan, Iran, and numerous other countries have recently launched cell-based treatment clinical investigations, and some findings have been published (Table 1). Research on a female patient with acute COVID19 syndrome was published in China, and the findings of laboratory tests and CT scans revealed highly successful outcomes after 21 days of therapy with umbilical cord MSCs. A recent case study of a case report of a 65-year-old female patient diagnosed with COVID-19 in critical condition revealed the precise 2019nCoV variant now known as SARS-CoV-2 [59].

The unfortunate SARS-CoV-2 outbreak has imposed an urgent demand from scientists and clinicians worldwide to find effective therapeutic agents to alleviate sufferings in COVID-19 patients on the one hand, while also enforcing a demand for effective vaccines to mitigate its spread in the future and spared or unaffected present populations on the other. Cell therapy and gene therapy are currently advanced disciplines of research used to treat a wide range of illnesses. It is important to note that the inflammatory cytokine profiles generated by H5N1 and COVID-19 are comparable [60], including exceptionally high amounts of IL-6, GCSF, IP10, MCP-1, MIP1A, and TNF-α, which are likely to cause serious organ damage and mortality in large numbers [61, 62]. MSC recipients had considerably lower levels of TNF-α, a powerful pro-inflammatory cytokine, and higher levels of IL-10, an anti-inflammatory cytokine, as compared to placebo control individuals. When administered intravenously, a portion of the MSCs settle in the lungs as well, which may enhance the pulmonary microenvironment and, as a result, minimize immune system over-activation and encourage regeneration of damaged lung tissues [63, 64].

Cytokine dysregulation has also been linked to influenza pathogenesis produced by a variety of influenza A virus subtypes. The most prevalent etiology found in these viral infection situations is pneumonia and acute respiratory distress syndrome. Intravenous injection of MSCs to patients suffering from influenza and other pulmonary illnesses has shown promising results in restoring lung functions and correcting the lung functional and structural losses caused by the infection's cytokine storm. As a result, it is reasonable to think that MSC treatment can benefit COVID-19 patients via immunomodulatory pathways [65]. Its therapeutic properties like immune response inhibition play a major role in combating viruses. Their capacity to develop into type II alveolar epithelial (AT2) cells in vitro may also aid [66]. Lung damage may change the signals of the BMP4 pathway, resulting in abnormal alveolar progenitor function. TSP1, a substance produced by MSCs, may help to re-establish damaged vascular signals and promote epithelial healing [67]. As a result, avoidance of cytokine storm is seen to be the most important stage in COVID-19 therapy. MSCs, due to their potent immunomodulatory properties, may have positive effects in avoiding or attenuating the cytokine storm and assisting in the regeneration of injured lung tissues/other organs.

3. CONCLUSION

The infectious viral SARS-CoV-2 produces COVID-19, which predominantly affects the respiratory system and lungs, resulting in pneumonia and organ damage. These microorganisms induce a strong immunological response, resulting in excessive cytokine production, which might be referred to as a “cytokine storm.” Genetic tests revealed that this virus is closely linked to SARS-CoV and is most likely to be found in wild bats. Because of the constant interaction with wildlife in markets, the outbreak was likely triggered by a zoonotic process. Even though the pandemic just began a few months ago, much genomic and phylogenetic information on the causal culprit has already been published. SARS-CoV-2, like other previously discovered
coronaviruses, infects host cells by attaching to the ACE2 receptor, and it is closely related to bat-SL-CoVZC45 and bat-SL-CoVZXC21, both of which have been detected in wild bat populations.

HUC-MSC therapy, as a non-invasive treatment, is a very successful and promising way for clinical use and promotion at this critical time due to the absence of viable methods to treat severe COVID-19. However, the cost-effectiveness and timeliness of therapeutic preparation are the most capable discussed topics for MSC-based therapy for COVID-19, but human life is much more valuable, and COVID-19 is so dangerous. As a result, clinical trials of MSC treatment to treat COVID-19 are still some time away, however, there are some intriguing results to consider. Stem cell therapy, particularly MSCs, maybe one of the most effective therapies, or a combination of treatments, for COVID-19 patients. However, scientists are working tirelessly to create a COVID-19 vaccine as well as medicines to treat this condition.

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