Mesenchymal Stem Cells: Time to Change the Name!

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ABSTRACT

Mesenchymal stem cells (MSCs) were officially named more than 25 years ago [1] to represent a class of cells from human and mammalian bone marrow and periosteum [3] that could be isolated and expanded in culture while maintaining their in vitro capacity to be induced to form a variety of mesodermal phenotypes and tissues. The in vitro capacity to form bone, cartilage, fat, etc., became an assay for identifying this class of multipotent cells and around which several companies were formed in the 1990s to medically exploit the regenerative capabilities of MSCs. Today, there are hundreds of clinics and hundreds of clinical trials using human MSCs with very few, if any, focusing on the in vitro multipotential capacities of these cells. Unfortunately, the fact that MSCs are called “stem cells” is being used to infer that patients will receive direct medical benefit, because they imagine that these cells will differentiate into regenerating tissue-producing cells. Such a stem cell treatment will presumably cure the patient of their medically relevant difficulties ranging from osteoarthritic (bone-on-bone) knees to various neurological maladies including dementia. I now urge that we change the name of MSCs to Medicinal Signaling Cells to more accurately reflect the fact that these cells home in on sites of injury or disease and secrete bioactive factors that are immunomodulatory and trophic (regenerative) meaning that these cells make therapeutic drugs in situ that are medicinal. It is, indeed, the patient’s own site-specific and tissue-specific resident stem cells that construct the new tissue as stimulated by the bioactive factors secreted by the exogenously supplied MSCs. Stem Cells Translational Medicine 2017;00:000–000

INTRODUCTION

Mesenchymal stem cells (MSCs) were officially named more than 25 years ago [1] to represent a class of cells from human and mammalian bone marrow and periosteum [3] that could be isolated and expanded in culture while maintaining their in vitro capacity to be induced to form a variety of mesodermal phenotypes and tissues (Fig. 1, The Mesengenic Process). The in vitro capacity to form bone, cartilage, fat, etc., became an assay for identifying this class of multipotent cells [9] and around which several companies (including Osi-ris Therapeutics, which my colleagues and I started,) were formed in the 1990s to medically exploit the regenerative capabilities of MSCs. Initially, the driving concept that a multipotent progenitor or “stem cell” existed in adult marrow was not only challenged, but was actively disregarded, especially by the orthopedic industry. Fast-forward to today and there are hundreds of clinics [10] and hundreds of clinical trials [11] using human MSCs (hMSCs) with very few, if any, focusing on the in vitro multipotential capacities of these cells.

Unfortunately, the fact that MSCs are called “stem cells” is being used to infer that patients will receive direct medical benefit, because they imagine that these cells will differentiate into the regenerating tissue-producing cells (i.e., these “stem cells” will be incorporated into and these differentiated cells will fabricate the diseased or missing tissue). Such a stem cell treatment will presumably cure the patient of their medically relevant difficulties ranging from osteoarthritic (bone-on-bone) knees to various neurological maladies, including dementia. I long ago urged, in print, that we change the name of MSCs to Medicinal Signaling Cells [12] to more accurately reflect the fact that these cells home in on sites of injury or disease and secrete bioactive factors [13] that are immunomodulatory and trophic (regenerative), meaning that these cells make therapeutic drugs [15] that are medicinal. It is, indeed, the patient’s own site-specific and tissue-specific resident stem cells that construct the new tissue as stimulated by the bioactive factors secreted by the exogenously supplied MSCs [16, 17].

HISTORY OF MSCS FROM A CAPLAN PERSPECTIVE

In the early 1970s into the 1980s, my colleagues and I published a number of papers based on the culturing of stage 24, embryonic chick limb bud mesodermal cells (ECLBMCs) that were observed to differentiate into cartilage, muscle, and bone under certain culture conditions [18–22]. These in vitro studies were correlated with a variety of in vivo studies that focused on the cellular and molecular events associated with the formation of embryonic limb bone [23, 24], cartilage [25], and muscle [26] in which several very prominent dogmas-of-the-day were challenged. For example, the concept that “cartilage is replaced by bone” led to the implication that if one could form cartilage in culture from embryonic mesodermal progenitor cells, one could observe the transition of that new cartilage into...
bone. Moreover, the endochondral replacement of cartilage by bone implied that the cartilage so formed in culture would differentiate into hypertrophic cartilage, which would calcify and provide the calcified matrix for subsequent bone formation. Although we documented that the ECLBMCs formed cartilage in culture [22, 25, 26] and that hypertrophic chondrocytes could be identified by the production of type X collagen, the only mineral that formed in culture was observed in the noncartilage, connective tissue valleys between mounds of cartilage [23, 24]. Reducing the initial plating densities of the freshly isolated ECLBMCs (where no cartilage formed) allowed us to observe the differentiation of a maximum number of cells into calcified matrix-producing osteoblasts [27].

Only when we went back to the developing chick embryo and carefully completed rigorous histology of the mid-diaphysis of the developing embryonic tibia, did we firmly establish that the new bone that formed came from a progenitor cell layer (stacked cell layer) outside and away from the already formed and expanding cartilage core (or cartilage model as it was called [28]) [29, 30]. Importantly, the hypertrophic cartilage core was replaced by invading vasculature and then marrow, not bone. Moreover, others [31, 32] clearly showed that these embryonic hypertrophic chondrocytes could be isolated, cultured, and maintained for many weeks in vitro, documenting that hypertrophic chondrocytes were not “programmed” to die (i.e., their demise was due to the nutrient and oxygen deprivation by a collar of calcified bone that was outside and away from the cartilage core) [29].

In addition, our early pioneering studies on the synthesis of proteoglycans of cartilage by the ECLBMC cells in culture with Vincent Hascall’s [33, 34] group brought us into areas of detailing the extracellular matrix (ECM) of first cartilage [35, 36], then muscle [37, 38] and bone and played an important role in our current interest in the basement membrane surrounding all blood vessels (to be discussed below).

We spent considerable time and effort in optimizing these stage 24 ECLBMC-cultures which, incidentally, we never called mesenchymal or mesodermal “stem cells,” although the evidence strongly suggested that they were multipotent. During this same time period and especially in the early 1980s, Marshal Urist and others were isolating molecular agents from the matrix of demineralized bone [39–41]. The phenomenological basis for such efforts stemmed from implantations of demineralized bone pieces into muscle or subcutaneous pockets in rodents, which eventually caused bone to de novo form from host cells [39]. Urist coined the term “bone morphogenetic proteins (BMPs)” to summarily refer to the bioactive agents released from demineralized bone.
matrix that could cause de novo bone to form in nonosseous tissues, such as muscle, or subcutaneously.

Stimulated by the public lectures and publications of Urist and because of a talented postdoctoral fellow, Glenn Syftestad, who had worked in Dr. Urist’s lab at UCLA before coming to my lab in 1981, we joined the race to purify the BMPs. Our first approach was to take high salt extracts of demineralized bone exactly as published by Urist and to put them on cultures of stage 24 ECLBMCs arranged to just form bone [42, 43]. To our great surprise, these extracts caused the cells to form mounds of cartilage. We named the presumed active agents as chondrogenic stimulating activity, which we purified, and the university filed patents [44, 45], which for reasons that could be challenged, they stopped maintaining. In the mid-1980s and certainly by 1987, it became known that Dr. John Wozney and his colleagues at Genetics Institute, Inc. (Cambridge, MA) had cloned BMP2 and had patented the BMP-family of molecules [46, 47]. The race for the BMPs was over, and my colleagues and I had failed to win, much less “place or show.”

In one of the demineralized bone implantation systems, Dr. Hari Reddi purified one member of this BMP-family and, importantly, characterized the in vivo temporal events caused by these factors [48–50]. These temporal events involved the invasion of the implant and cell division of host mesenchymal cells followed by their differentiation into cartilage which became hypertrophic and which was replaced by vascularized and marrowized bone [50]. Using Dr. Reddi’s histology slides of these subcutaneous implantation specimens, which he generously provided, I suggested that the temporal sequence of cartilage replacement by bone was identical to that which we described in the developing embryonic chick tibia [5]—essentially, that the implanted demineralized bone particles were surrounded by mesenchymal progenitor cells, which were attracted to the demineralized particles and formed cartilage. Since the implant was walled-off, encysted by a layer of these mesenchymal cells comparable to the stack cell layer of the embryonic chick tibia, all blood vessels were excluded. The blood vessels outside the layer of surrounding and encysting mesenchymal cells caused the bottom layer of encircling cells to differentiate into a layer of osteoblasts, which fabricated a layer of osteoid that became mineralized. The deprivation of nutrients and oxygen caused the encased chondrocytes to form hypertrophic cartilage (Reddi documented the production of type X collagen) whose cells expired, releasing large quantities of vascular endothelial growth factor (VEGF), which caused the external vascular tree to invade just as occurs in the mid-dyaphasis of the embryonic chick tibia, all blood vessels were excluded. These invading vessels brought a fresh supply of mesenchymal progenitor cells, whose cells expired, releasing large quantities of vascular endothelial growth factor (VEGF), which caused the external vascular tree to invade just as occurs in the mid-dyaphasis of the embryonic chick tibia [5, 29, 30]. These invading vessels brought a fresh supply of mesenchymal progenitor cells, which then formed vascularized and marrowized bone.

Without going into details, the central fact that comes from the above is that upon jamming the demineralized bone into muscle or the subcutaneous sites, the release and clustering of mesenchymal progenitor cells could be documented in these adult rodent hosts. In concert with these facts was the realization that adult bone marrow contained the same or similar primitive osteochondral progenitors [51–57]. The presence of these mesenchymal progenitors could be deduced from many avenues of exploration: (a) since the days of Aristotle, bone marrow was known to enhance orthopedic/bone healing [51]; (b) in modern terms, Connolly et al. [52, 53] and more recently Hernigou [58], documented the direct osteochondral potency of bone marrow or bone marrow aspirates; (c) Friedenstein et al., as rediscovered and popularized by Owen, showed that clones of adherent osteogenic progenitor cells could be isolated and propagated in culture from adult marrow [54–57]; and last (d) Owen herself imagined a crude mesenchymal lineage comparable to that described for descendants of hematopoietic stem cell (HSC) [57]. It is important to stress that in the 1980s and early 1990s, the dogma-of-the-day was that the only stem cells that existed in the adult body were HSCs.

The above facts (especially the demineralized bone implantations into adult hosts) led Dr. Stephen Haynesworth and me to see if we could isolate and purify the mesenchymal progenitor cells from adult human bone marrow [1, 59–63]. At that time, we were not aware of the work of Friedenstein and of Owen, which was lucky because we had the ECLBMC system, which was quite different from the culture conditions of Friedenstein and Owen. We had long before optimized this ECLBMC system, in particular by optimally choosing the batch of fetal bovine serum (FBS) used to culture these chick embryonic cells [64]. This lucky batch of serum was later shown to be optimal for the attachment, propagation, and maintenance of the multipotency of the culture adherent cells from human adult marrow [65, 66]. Indeed, one in 10–20 batches of FBS was shown to be optimal for marrow-derived hMSCs by the ECLBMC culture assay system, which eventually was replaced by other criteria [66]. This assayed batch of FBS allowed MSCs to optimally attach to the culture dishes, to expand to form colonies (referred to as colony form units/fibroblast, CFU-f by Friedenstein [55], that could be counted to give MSC titers, which ranged from 1 in 10,000 marrow cells in newborns to 1 in 2 million marrow cells in 80-year-old marrow donors [67]. Given all of the above, I named these propagated cells that were multipotent in culture assay: MSCs [1].

**MSCs: Various Names Mean the Same**

Given the historic outline above, various names for these culture adherent and passaged adult marrow-derived, multipotent mesenchymal cells came to mind:

**Marrow Stromal Cells**

The term “stroma” is an older morphological term meaning from connective tissue or the structural component of tissue. As defined by Owen in 1988 [57], these are fibroblastic cells that adhere to plastic and expand, forming colonies (CFU-f) that are osteogenic. One could also envision that bone marrow stroma was a unique scaffold that supports different lineage arms of hematopoiesis. Such a three-dimensional connective tissue scaffold does not exist in marrow, although the vision of such a specialized framework is enchanting.

**Multipotent Stromal Cells**

MSCs can be multipotent, as documented in various culture circumstances.

**Mesodermal Stem Cells**

Because of our studies of ECLBMC cells, this term was highly favored, especially because all of the induced or bioactive factor-treated cells and tissue formed in culture were of mesodermal (middle layer of the embryo) origin.
MSCs
I chose this term because mesenchyme is a type of tissue characterized by loosely associated cells that lack polarity and are surrounded by a large ECM. Because of their in vitro multipotency and clonability [68], I, provocatively, called them “stem cells” to especially appeal to the orthopedic community. As defined by hematologists, all stem cells must be capable of serial transplantation and unlimited doublings. Indeed, there are published reports that support this definition [69, 70].

Mesenchymal Stromal Cell
A group of scientists at an international meeting termed the MSC as a “stromal” cell because they did not favor the stem cell classification and imagined, incorrectly, that the origin of MSCs, from a variety of tissues, was the connective tissue layer of that tissue [9].

Medicinal Signaling Cell
Because the function of MSCs in vivo is secretory and primarily functional at sites of injury, disease, or inflammation, I now favor this terminology [12].

The New Science: MSCs Are Derived from Pericytes
Central to the renaming strategy is the fact that most, if not all, MSCs are derived from the differentiation of perivascular or mural cells, pericytes [71]. The studies of Dr. Bruno Péault and colleagues [72] clearly document that pericytes isolated from a variety of tissues give rise to MSCs, as identified by cell surface antigens and their in vitro multipotency. Importantly, MSCs can be isolated from every vascularized tissue [73] and even from menstrual flow [74, 75] (i.e., broken blood vessels release the perivascular cells that differentiate into MSCs). The perivascular location as the origin of MSCs and their functional capacity to be immunomodulatory and trophic (including fabricating and secreting antibiotic proteins [76]) challenges the “stromal” name and origin of the MSCs [9, 77, 78].

Based on the above, we have assembled the new and current information on the pericyte MSC (pMSC) into a poster, which has a number of interesting and unusual pieces of information not previously appreciated [79]. These include the fact that each separate tissue-specific stem cell is both in communication with its underlying vascular endothelial cells and neighboring specific pericyte/MSC [Universal Stem Cell Niche]. These pMSCs are specific to each stem cell, including a chemically different pMSC next to the active versus quiescent HSC in marrow [80, 81]. In every tissue examined in detail, the marrow, neural tissue [82], liver [83], heart [84], etc., tissue-specific stem cells are next to its specific pMSC on a blood vessel. These observations further support the concept that all pMSCs have both MSC-common and MSC-unique chemical and functional features. In the in vitro multipotency assays, the assay must be optimized for each tissue specific MSC. For example, hMSCs of marrow were shown to be induced in culture into the chondrogenic lineage by TGF-β [85], while fat-derived hMSCs require both TGF-β and BMP-6 [86]. The main in vivo functional differences of MSCs from different tissues or organs remain largely unknown, even though the major therapeutic functionality of MSCs at various sites of disease or injury are very similar when comparing these different MSCs [87].

Changing Names
Since the main functionality in vivo of MSCs [88] is not multipotency and, thus, not as a stem cell [89–91], I propose that its name be changed. The precedent for changing medical terms is not new. For example, names of many diseases have been changed: ablepsy was changed to blindness; ague to malarial fever; American plague to yellow fever; anasarca to generalized massive edema; aphi to laryngitis; aphi to thrush in infants; and apoplexy to paralysis due to stroke [92]. Of course, there is great stigma associated with the accepted names for some diseases; multiple sclerosis was once called hysterical paralysis when people believe this was caused by stress linked with oedipal fixations. Chronic fatigue syndrome is a serious ailment, yet 85% of clinicians view it as a psychiatric disorder; activists are currently trying to change the name to remove the bias and stigma. There is no stigma associated with the term MSC except, for me, the implied promise that it is a true “stem cell,” which it is not in vivo.

It has been argued, because MSC science and clinical use is so strong and, indeed, positive with almost 700 clinical trials listed on clinicaltrials.gov, that the MSC nomenclature should remain. The problem is not with the “mesenchymal” part of the name; it is the “stem cell” part of the name that is the issue. As outlined in our poster, the pMSC functions quite differently from the released pericyte that forms an activated, site-specific MSC. Infused autologous or allogenic MSCs appear to home in on active vascular sites of injury or inflammation [93]. At such disease sites, the MSC rarely or never differentiate into the tissue at that site [13, 88], but they secrete bioactive factors (some of the names of these factors we know [94]) and their therapeutic effects can be analyzed as site-specific clinical outcome parameters. Outcomes for graft-versus-host disease, acute myocardial infarct, low back pain, osteoarthritic knees, tendonitis, and aspects of inflammatory bowel disease or Crohn’s disease have been reported (www.mesoblast.com). Again, for emphasis, these MSC-effects are medicinal.

MSCs Are Not Stem Cells
The science and commercialization of adult MSCs were enhanced by the popularization of embryonic stem cells (ESCs) and made more attractive by President Bush’s prohibition of the use and study of ESCs [95]. This popularization of ESCs also served as a disadvantage because all “stem cells” have been viewed by the public as being pluripotent or multipotent. Thus, the infusion of hMSCs in an osteoarthritic knee is imagined to contribute directly into the regeneration of cartilage tissue by the infused MSCs forming functional chondrocytes that fabricate functional cartilage tissue. The infusion into cardiac patients of hMSCs assumes that these cells will directly convert into functional heart muscle cells to replace the cells that die from the ischemia of the heart attack. And so on and so on: stem cells directly convert into the diseased or injured tissue in question. Although we, in this field, all have our own favorite explanation for the mechanisms that govern the observed positive therapeutic outcomes, the in vivo effects of infused hMSCs are best described as medicinal and most likely not associated with the infused cells differentiating into regenerative or replacement tissue [96–99]. These stem cell misconceptions have led some practitioners in the United States and worldwide to advertise the availability of stem cell-treatments (i.e., MSCs can cure the blind, make the lame walk, and make old tissue young again [10]). I, of course,
want the MSC nomenclature to remain in use, but not as stem cells. Perhaps we should call them magic signaling cells, more strategic cells, maxi secreting cells, most sensitive cells, main secreting cells, or message secreting cells. I propose to change the name of MSCs to reflect our new understanding that they do not function in the body as progenitors for tissues, neither in the normal steady-state nor in disease or injury circumstances; they are not stem cells.

**MSCs and Metastasis**

Last, we recently published a treatise which documents that the pMSC actively binds to and pulls circulating melanoma cells into the marrow of bone [100]. This grab/pull mechanism for melanoma metastasis is counter to the current concept that metastatic cells secrete digestive enzymes that allow the melanoma to erode its way into bone. We further hypothesized that the laminin identity in the basement membrane ECM of the blood vessels plays both an active and permissive role in the extravasation of melanoma into bone. Thus, the melanoma must pass through the endothelial cell layer, its basal lamina or basement membrane and past the dense covering of mural cells. The active pMSC not only facilitates this extravasation, but is actively and molecularly controlling this translocation from the circulation into the marrow of bone. Clearly, the pMSC is not medicinal in this context even though its differentiated progeny, the MSC, can provide powerful medicinal benefit given other circumstances. Last, the pMSC is not multipotent nor does it, itself, cause tumors to form. The pMSC is corrupted by the cancer cell; it does not corrupt normal cells to become cancerous.

**Conclusions**

It should be permissible for the person named the MSCs to drop the stem cell nomenclature because it is scientifically and therapeutically misleading. In 2010, I proposed that we call them medicinal signaling cells [12]. That is what these do, and the culture plasticity of most mesenchymal cells [we can induce adult human chondrocytes to make a bone or fat in cell cultures [101]] means that the stem cell moniker is inappropriate. I was wrong. I take back the name that I gave these hugely important cells. Call them MSCs, but please, not stem cells.

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**Disclosure of Potential Conflicts of Interest**

Case Western Reserve University receives royalties from Osiris Therapeutics, Inc. (now by Mesoblast Limited) which they share with AIC for MSC technology transferred in 1992. No other conflicts exist.

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