33

# Elevation of Serum Oncostatin M Level in Acute Ischaemic Stroke: A Review

Michael Christian and Youdong Wei

Department of Neurology, the First Affiliated Hospital of Chongqing Medical University, Chongqing, China

DOI: https://doi.org/10.17925/USN.2024.20.1.7

stroke, a neurovascular disorder, is one of the main causes of mortality and disability worldwide. There are several mechanisms of stroke, with inflammatory response being a key factor, with previous studies have concluding the involvement of oncostatin M (OSM) in inflammatory response. This review aims to analyze current knowledge to explain the elevation of serum OSM expression in acute ischaemic stroke (AIS). Information was gathered from the PubMed database on landmark studies on the relevance of serum OSM in AIS, and it was concluded that hypoxia and stagnant blood flow, which are involved in the typical pathophysiology of AIS, induced an inflammatory signalling pathway. OSM, a member of cytokine interleukin-6, participates in inflammatory responses in a dual manner: pro-inflammatory and anti-inflammatory. We concluded the elevation of serum OSM level in patients with AIS was due to an inflammatory response triggered by AIS. Cardioembolic stroke, a subtype of AIS, triggered a higher inflammatory substrate response in the circulatory system, explaining why serum OSM elevation is significant compared to other subtypes of acute ischaemic stroke.

#### Keywords

Acute ischaemic stroke, embolic stroke, neuroinflammation, oncostatin M, interleukin-6, glycated serum proteins

**Disclosures:** Michael Christian and Youdong Wei have no financial or non-financial relationships or activities to declare in relation to this article.

Review process: Double-blind peer review.

**Compliance with ethics:** This article involves a review of the literature and did not involve any studies with human or animal subjects performed by any of the authors.

**Data availability:** Data sharing is not applicable to this article as no datasets were generated or analyzed during the current study or during the writing of this article.

**Authorship:** The named authors meet the International Committee of Medical Journal Editors (ICMJE) criteria for authorship of this manuscript, take responsibility for the integrity of the work as a whole and have given final approval for the version to be published.

Access: This article is freely accessible at touchNEUROLOGY.com © Touch Medical Media 2024.

Received: 26 August 2023
Accepted: 31 October 2023
Published online: 26 January 2024

this article.

Citation: touchREVIEWS in Neurology. 2024;20(1):33-36

Corresponding author: Dr. Youdong Wei, Department of Neurology, The First Affiliated Hospital of Chongqing Medical University, Chongqing, China. E: havonewei@163.com

Support: No funding was received in the publication of

## Introduction

The term 'stroke' is used to describe an adverse clinical state involving interference of blood circulation to the brain due to obstruction or rupture of blood vessels. Stroke was previously categorized into a cardiovascular disorder until the release of the International Classification of Disease 11 (ICF-11) in 2018, when stroke was recognized as a neurological disorder.<sup>2</sup> For the past decade, it has been ranked the second most common cause of death, and a major cause of disability worldwide.3 Between the two subtypes of stroke, haemorrhagic and ischaemic, ischaemic stroke is the more common subtype, accounting for up to 87% of stroke cases in the United States. <sup>4</sup> The prevalence of stroke is also affected by demographics, being more prevalent in developing countries.3 In 2017, it was recorded that stroke and ischaemic heart disease was the leading cause of mortality and disability nationally in China.<sup>5</sup> Acute ischaemic stroke (AIS) is a type of stroke that progresses rapidly over a short period of time. In addition to symptomatic treatment and secondary prevention, the management of AIS is focused on reperfusion through intravenous thrombolysis and endovascular thrombectomy; both of which aim to reduce disability, but are highly dependent on time.<sup>6</sup> It is important that the thrombi derived from patients with AIS are highly heterogeneous, as this affects the choice of available treatment options and recovery prognosis. Thrombi with a higher composition of red blood cells (RBCs) tend to have a good prognosis, whereas thrombi with a higher fibrin composition have a tendency to be stiff, making it harder to treat and therefore leading to a poorer prognosis.<sup>7</sup>

Several methods have been established to classify the characteristics of stroke. The most widely used classification system is the Trial of Org 10 172 for Acute Stroke Treatment (TOAST), introduced in 1993. With the advancement of research and technologies, several classification systems have been assembled based on different approaches, such as the Causative Classification System (CCS) and Chinese Ischemic Stroke Subclassification (CISS). Amongst other classification methods not mentioned here, atherosclerosis and cardioembolic (CE) subtypes were mentioned in the above examples. Although the occurrence of the CE subtype is less than the AS subtype (accounting for about 25% of AIS cases on average), it is often more severe and more prone to early and late recurrences. In

Recent studies relating to inflammation and cancer inhibition have brought oncostatin M (OSM), a group of glycoproteins (gp), to researchers attention. OSM is a member of the interleukin 6 (IL-6) cytokine sharing a signalling pathway with gp130. The mechanism of engagement is through a second receptor, either through type 1 (Leukemia Inhibitory Factor Receptor  $\alpha$  [LIFRa]) or type 2 (OSM receptor  $\beta$  chain [OSMR $\beta$ ]), before engaging with gp130 and OSMR. The study reported that OSM had a neuroprotective function related to ischaemic-reperfusion brain injury, which leads to a potential therapeutic effect. This review aims to explore the current understanding behind the elevation of serum OSM in patients with AIS.

TOUCH MEDICAL MEDIA Journal Publication Date: 30 May 2024

### **Methods**

We searched the National Center for Biotechnology Information (NCBI) PubMed database for studies that link OSM and AIS. We explored the pathophysiology of AIS, the mechanism of action of OSM to find the cause of serum OSM elevation in AIS and the distinguished traits of the CE subtype that are associated with significant elevation of serum OSM level. An advanced search using MeSH terms was implemented using the search strategy ("acute ischemic stroke" OR "stroke" OR "AIS"), "neuroinflammation," and ("oncostatin M" OR "OSM"). A search filter was set for all types of articles in English with free full text. Studies that contained the information above were selected for both animal tests and human trials. We focused on the objectives of the studies, the test subjects and the findings. Results from the selected studies were analyzed and presented in this review article.

# **Findings**

# Neuroinflammatory in acute ischaemic stroke

The term 'neuroinflammation' has been widely used to refer to inflammatory responses found in neural cells following injuries, including stroke. Multiple processes have been proposed to explain the brain injury caused by ischaemia, including excitotoxicity, oxidative stress and inflammation.<sup>13</sup> Although the manifestation of post-ischaemic inflammation can occur days or weeks after the event, the inflammatory cascade is initiated as immediately after the occurrence of vessel occlusion. 14 In a study published by Anrather and Iadecola in 2016, the authors described the pathway from a hypoxic state to an inflammatory response in the stroke area in the brain. 15 To begin with, a stagnant blood flow triggers stress on the vascular endothelium and platelets. This signals an adhesion molecule P-selectin to accumulate on the surface of cells within minutes after activation. Selectins are important in controlling the circulation of leukocytes. 16 Inflammation cascades continue to occur in several areas, including the brain parenchyma. One of the earliest events is the release of danger-/damage-associated molecular patterns (DAMP) from injured and dying neurons, which together with other components finally activates the pattern recognition receptor on microglia, brain resident immune cells and astrocytes. 17-20 Microglia are activated in the brain due to the rise of extracellular ATP from depolarization of neurons.<sup>21</sup> These DAMPs and other cytokines can penetrate the blood brain barrier (BBB) and enter the circulation, which then triggers a systemic inflammatory response. This systemic inflammatory response is characterized by the increase of serum several cytokine expressions including IL-6, which is why IL-6 is found to be elevated in patients with AIS. 22,23 Neuroinflammation may increase potential additional harm, resulting in cell death, but at the same time, it has a helpful effect in promoting healing. This neuroinflammatory response is recorded to be present in all subtypes of stroke, but is deemed amplified in the CE subtype.24

## Cardioembolic subtype

While stroke has been recategorized as a neurological disorder, it is still a common practice to observe cardiac injury following cerebrovascular disease.<sup>25</sup> The second leading cause of mortality following ischaemic stroke has been reported to be cardiovascular complications. It has also been reported that the risk of cardiac complications in proportional to the severity of ischaemia.<sup>26</sup> A study by Maida et al. focused on neuroinflammatory mechanisms in CE ischaemic stroke. The authors reported an accumulation of evidence demonstrating that ischaemic stroke initiates a complex process involving genetic, molecular and cellular changes. Phlogosis plays a critical part in this mechanism, both in the central and peripheral nervous systems.<sup>22</sup> Another study recorded

that systemic inflammation participated in the pathological process that led to cardioembolism, suggesting that a higher inflammatory substrate could be found in CE stroke compared to the other subtypes.<sup>27</sup> With growing evidence indicating neuroinflammation preceding AIS leads to an increase of cytokines and chemokines that stimulate infiltration of leukocytes into the cerebral parenchyma. If an extensive area of cerebral tissue is affected it could induce a higher neuroinflammatory response.<sup>22</sup>

#### Oncostatin M

There have been studies into OSM since 1986 when it was first identified in the conditioned media of phorbol 12-myristate 13-acetate (PMA)-treated U937 monocytic cells.<sup>28</sup> OSM belongs to the IL-6 family of cytokines which contribute to communication and signalling pathways. Several studies have recorded the involvement of OSM in physiological and pathophysiological activities, including haematopoiesis, mesenchymal stem cell differentiation, fibrosis, nociception, inflammation, metabolism and cancer. 29-33 The biological activities of OSM vary in different species, according to the concentration of ligands and specific cell types.<sup>29,34</sup> In animal models, mouse OSM (mOSM) only binds to OSMRB/gp130 in mice, whereas human OSM (hOSM) signals both LIFR\$ and OSMR\$, specifically called type I receptor complex (LIFR\$/gp130) or type II receptor complex (OSMRB/gp130). 35,36 Hermanns et al. reported that OSM possessed the broadest signalling profile, which comprised Janus kinase (JAK)/signal transducer and activator of transcription (STAT), mitogen-activated protein kinases (MAPK) - including extracellular signal-regulated kinase (ERK), p38 and Jun NH<sub>2</sub>-terminal kinase (JNK) – the phosphatidylinositol-3kinase (PI3K)/Akt and the protein kinase C delta (PKCδ) pathways.<sup>29</sup> This broad signalling profile was due to OSM comprising BC loops, a unique helical loop on OSM between its B and C helices that is not found on other IL- 6 family cytokines, that act as a steric barrier for OSMRB and LIFRB. As a result, OSM initially binds with gp130 before signalling OSMRB or LIFR<sub>B</sub>.37

The production of OSM in the human body is derived from multiple sources, mainly the cells of the immune system: dendritic cells, neutrophils, monocytes/macrophages and T cells.38-40 Regardless of any existing inflammation, haematopoietic cells of the bone marrow also produce OSM, explaining why a low concentration of serum OSM can be found in a normal healthy body. 41 While previous studies focus on the role of OSM with regard to inflammation, cell proliferation and haematopoiesis, increasing evidence reported the involvement of OSM in the nervous system. In the central nervous system, OSM is mainly expressed by neurons, astrocytes and microglia.42-44 OSM has been implicated in the homeostasis of neural progenitor cells (NPCs) in physiological settings, a pool of cells for the production of new neural cells located in the subventricular zone, hippocampus and olfactory bulb in the adult mammalian brain. 45-48 It has been reported that the number of OSMR\$-positive neurons in the site of injury was significantly decreased following a sciatic nerve axotomy.<sup>49</sup> OSM has been shown to decrease NMDA-induced neuronal death and inhibit excitotoxic injury in vivo.50 OSM-based treatment also showed a reduction in lesion size after a spinal cord hemisection and improved functional recovery and neurite outgrowth in an animal model.<sup>51</sup> Although several studies reported neuroprotective properties of OSM, some studies also reported contradictory findings. A study reported OSM derived from mononuclear cells induced apoptosis in primary neurons.<sup>52</sup> Contradictory roles of OSM might be correlated with different experimental settings, cell types, stimuli and recruitment of receptors.35

In inflammatory conditions, OSM is significantly involved in both acute and chronic phases, leading to tissue fibrosis and cancer, and is also

34

expressed by immune cells after a variety of soluble mediators.<sup>53</sup> A double-side effect of OSM can act as an anti- or pro-inflammatory action. This is due to the ability of OSM to either directly recruit or activate innate immune cells, or indirectly regulate stromal cells located in the area of injury.54,55 In pro-inflammatory activities, OSM has been recognized to induce target cells to release cytokines and chemokines, such as CXCL3, CCL2, CCL5, and CCL20,56 which proceed with neutrophils and monocyte/macrophage recruitment, creating a positive amplification of OSM's effect. In chronic inflammatory conditions, an overexpression of OSM/OSMBR is often found, where OSM sustains inflammation and promotes fibrosis.<sup>36</sup> OSM concentration is relatively modest in a normal healthy condition and multiple studies have confirmed a significant increase in OSM from a few hours after the onset of stroke that can persist for up to 90 days following cerebral ischaemia.<sup>29,57</sup> Although previous studies identified OSMB expression in several cell types in the brain (neurons and astrocytes, but not microglia), the OSMB expression profile proceeding onset of stroke is still unclear. 58 OSMβ protects against ischaemic/reperfusion (I/R)-induced cerebral injury by activating the JAK2/STAT3 cascade, but it is not clear whether JAK2/STAT3 is necessary for OSMβ-mediated protection in vivo. Still, OSM is considered to be a promising therapeutic agent in rats and humans.<sup>35</sup>

## **Discussion**

Everything related to AIS, from onset to treatment, is time critical. Understanding the mechanism of AIS helps to identify the pathophysiological nature of AIS, as well as appropriate responses in order to develop target-specific treatments. This review article reported the three major mechanisms of neurological damage following an infarct and stroke: loss of neurons, excessive production of reactive oxygen species and inflammation. <sup>59-61</sup> Among the three mechanisms, we focused on the inflammation response. In an ischaemic environment, glial cells such as astrocytes and microglia, together with endothelial cells and leukocytes, release various pro-inflammatory agents and initiate signalling cascades facilitated by chemokines, cytokines and other enzymes. During the review process, we specifically focused on OSM.

OSM stands out due to its wide range of cascading capabilities and its neuroprotective properties. In addition to that, hOSM is correlated with

both type 1 and type 2 signalling pathways, whereas mOSM is only associated with type 2 signalling pathways. This allows researchers to conclude a certain degree of representation between animal models and humans. It has been known that OSM exists in a normal, healthy human body and is significantly elevated in inflammatory conditions, including AIS. Unfortunately, the present understanding of the minimum threshold that causes OSM elevation is still unknown. Due to its nature, OSM can act as an anti- and pro-inflammatory agent, which causes a dilemma in finding the appropriate usage of OSM related to AIS. OSM can be measured through enzyme linked immunosorbent assay (ELISA), which is a strength for its practical accessibility.

We identified several limitations reported in past studies. It was reported that a close interaction between pro-inflammatory and anti-inflammatory molecules during the early stages of stroke was present, but the precise understanding in the process that modulates the interaction was still lacking. Studies about the correlation between OSM and its properties in the nervous system are still very limited. There is also a limitation of data availability on the neuroinflammation association with cardioembolism, meaning that future studies are needed to accurately determine the reasoning behind CE stroke.

#### Conclusion

This review was designed to assess current knowledge about the relationship between serum OSM and AIS. Hypoxia and stagnant blood flow trigger inflammatory responses from the tissue. Inflammatory responses following AIS can be found as early as a few hours after onset of AID and can last for days. This inflammatory response triggers the signalling pathway of OSM through the activation of cytokines. There are multiple producers of OSM in the body. OSM is known to have a dual effect: to induce inflammation and to respond to inflammation. Several studies have reported the elevation of OSM levels in AIS, notably in CE. The significant increase in the CE subtype is due to a higher concentration of inflammatory substrates in the circulation expressing the OSM. With a lack of published studies around OSM levels in humans, further research should be carried out to further investigate the findings reported in animal studies, as well as the association between neuroinflammation and cardioembolism.  $\Box$ 

- Lo EH, Dalkara T, Moskowitz MA. Mechanisms, challenges and opportunities in stroke. Nat Rev Neurosci. 2003;4:399–415. DOI: 10.1038/nrn1106.
- Shakir R. The struggle for stroke reclassification. Nat Rev Neurol. 2018;14:447–8. DOI: 10.1038/s41582-018-0036-5.
- Kuriakose D, Xiao Z. Pathophysiology and treatment of stroke: Present status and future perspectives. Int J Mol Sci. 2020;21:7609. DOI: 10.3390/ijms21207609.
- Barthels D, Das H. Current advances in ischemic stroke research and therapies. *Biochim Biophys Acta Mol Basis Dis.* 2020;1866:165260. DOI: 10.1016/j.bbadis.2018.09.012.
- Zhou M, Wang H, Zeng X, et al. Mortality, morbidity, and risk factors in China and its provinces, 1990–2017: A systematic analysis for the global burden of disease study 2017. The Lancet. 2019;394:1145–58. DOI: 10.1016/S0140-6736(19)30427-1
- Walter K. What is acute ischemic stroke JAMA. 2022;327:885. DOI: 10.1001/jama.2022.1420.
- Jolugbo P, Ariéns RAS. Thrombus composition and efficacy of thrombolysis and thrombectomy in acute ischemic stroke. Stroke. 2021;52:1131–42. DOI: 10.1161/ STROKEAHA.120.032810.
- Adams HP, Bendixen BH, Kappelle LJ, et al. Classification of subtype of acute ischemic stroke. definitions for use in a multicenter clinical trial. TOAST. trial of Org 10172 in acute stroke treatment. Stroke. 1993;24:35–41. DOI: 10.1161/01.str.24 1.35.
- Chen PH, Gao S, Wang YJ, et al. Classifying ischemic stroke, from TOAST to CISS. CNS Neuroscience & Therapeutics. 2012;18:452–6 DOI: 10.1111/j.1755-5949.2011.00292.x
- 2012;18:452–6. DOI: 10.1111/j.1755-5949.2011.00292.x.
   Pillai AA, Tadi P, Kanmanthareddy A. Cardioembolic stroke. In: StatPearls. Treasure Island. FL. USA: StatPearls Publishing. 2023

- Auguste P, Guillet C, Fourcin M, et al. Signaling of type II oncostatin M receptor. J Biol Chem. 1997;272:15760–4. DOI: 10.1074/ibc.272.25.15760.
- Guo S, Li ZZ, Gong J, et al. Oncostatin M confers neuroprotection against ischemic stroke. J Neurosci. 2015;35:12047–62. DOI: 10.1523/JNEUROSCI.1800-15.2015.
- Moskowitz MA, Lo EH, ladecola C. The science of stroke: Mechanisms in search of treatments. *Neuron*. 2010;67:181–98 DOI: 10.1016/i.neuron.2010.07.002.
- Iadecola C, Anrather J. The immunology of stroke: From mechanisms to translation. *Nat Med*. 2011;17:796–808. DOI: 10.1038/nm.2399.
- Anrather J, ladecola C. Inflammation and stroke: an overview. Neurotherapeutics. 2016;13:661–70. DOI: 10.1007/s13311-016-0483-x
- De Meyer SF, Denorme F, Langhauser F, et al. Thromboinflammation in stroke brain damage. Stroke. 2016;47:1165-72. DOI: 10.1161/STROKEAHA.115.011238.
   Petrovic-Djergovic D, Goonewardena SN, Pinsky
- DJ. Inflammatory disequilibrium in stroke. *Circ Res*. 2016;119:142–58. DOI: 10.1161/CIRCRESAHA.116.308022.

  18. Garcia-Bonilla L, ladecola C. Peroxiredoxin sets the brain on fire
- after stroke. *Nat Med*. 2012;18:858–9. DOI: 10.1038/nm.2797.

  19. Benakis C. Garcia-Bonilla I. Jadecola C. Anrather J. The role
- Benakis C, Garcia-Bonilla L, Iadecola C, Anrather J. The role of microglia and myeloid immune cells in acute cerebral ischemia. Front Cell Neurosci. 2014;8:461. DOI: 10.3389/ fncel.2014.00461.
- Singh V, Roth S, Veltkamp R, Liesz A. Hmgb1 as a key mediator of immune mechanisms in ischemic stroke. Antioxid Redox Signal. 2016;24:635–51. DOI: 10.1089/ars.2015.6397.
- Davalos D, Grutzendler J, Yang G, et al. ATP mediates rapid microglial response to local brain injury in vivo. Nat Neurosci. 2005;8:752–8. DOI: 10.1038/nn1472.

- Ferrarese C, Mascarucci P, Zoia C, et al. Increased cytokine release from peripheral blood cells after acute stroke. J Cereb Blood Flow Metab. 1999;19:1004–9. DOI: 10.1097/00004647-1999/9000-00008.
- Basic Kes V, Simundic AM, Nikolac N, et al. Pro-inflammatory and anti-inflammatory cytokines in acute ischemic stroke and their relation to early neurological deficit and stroke outcome. Clin Biochem. 2008;41:1330–4. DOI: 10.1016/j. clinbiochem.2008.08.08.
- Maida CD, Norrito RL, Daidone M, et al. Neuroinflammatory mechanisms in ischemic stroke: Focus on cardioembolic stroke, background, and therapeutic approaches. Int J Mol Sci. 2020;21:6454. DOI: 10.3390/ijms21186454.
- Ay H, Koroshetz WJ, Benner T, et al. Neuroanatomic correlates of stroke-related myocardial injury. Neurology. 2006;66:1325–9. DOI: 10.1212/01.wnl.0000206077.13705.6d.
- Yoshimura S, Toyoda K, Ohara T, et al. Takotsubo cardiomyopathy in acute ischemic stroke. *Ann Neurol*. 2008;64:547–54. DOI: 10.1002/ana.21459.
- Tuttolomondo A, Di Raimondo D, Pecoraro R, et al. Inflammation in ischemic stroke subtypes. Curr Pharm Des 2012;18:4289–310. DOI: 10.2174/138161212802481200.
- Zarling JM, Shoyab M, Marquardt H, et al. Oncostatin M: A growth regulator produced by differentiated histiocytic lymphoma cells. Proc. Natl Acad Sci U S A. 1986;83:9739–43. DOI: 10.1073/pnas.83.24.9739.
- Hermanns HM. Oncostatin M and interleukin-31: Cytokines, receptors, signal transduction and physiology. Cytokine Growth Factor Rev. 2015;26:545–58. DOI: 10.1016/j. cytogfr.2015.07.006.
- Yu Z, Li Z, Wang C, et al. Oncostatin M receptor, positively regulated by Sp1, promotes gastric cancer growth and

touchREVIEWS in Neurology 35

- metastasis upon treatment with oncostatin M. *Gastric Cancer* 2019;22:955–66. DOI: 10.1007/s10120-019-00934-y.

  31. Matsuda M, Tsurusaki S, Miyata N, et al. Oncostatin M causes
- Matsuda M, Tsurusaki S, Miyata N, et al. Oncostatin M causes liver fibrosis by regulating cooperation between hepatic stellate cells and macrophages in mice. *Hepatology*. 2018;67:296–312. DOI: 10.1002/hep.29421.
- Pothoven KL, Schleimer RP. The barrier hypothesis and oncostatin M: Restoration of epithelial barrier function as a novel therapeutic strategy for the treatment of type 2 inflammatory disease. *Tissue Barriers*. 2017;5:e1341367. DOI: 10.1080/21688370.2017.1341367.

   West NR, Hegazy AN, Owens BMJ, et al. Oncostatin M
- West NR, Hegazy AN, Owens BMJ, et al. Oncostatin M drives intestinal inflammation and predicts response to tumor necrosis factor-neutralizing therapy in patients with inflammatory bowel disease. Nat Med. 2017;23:579–89. DOI: 10.1038/nm0617-788d
- Richards CD. The enigmatic cytokine oncostatin M and roles in disease. ISRN Inflamm. 2013;2013:512103. DOI: 10.1155/2013/512103.
- Guo S, Li ZZ, Gong J, et al. Oncostatin M confers neuroprotection against ischemic stroke. J Neurosci. 2015;35:12047–62. DOI: 10.1523/INFUROSCI.1800-15.201
- 2015;35:12047–62. DOI: 10.1523/JNEUROSCI.1800-15.2015.
   Stawski L, Trojanowska M. Oncostatin M and its role in fibrosis. *Connect Tissue Res*. 2019;60:40–9. DOI: 10.1080/03008207.2018.1500558.
- Chollangi S, Mather T, Rodgers KK, Ash JD. A unique loop structure in oncostatin M determines binding affinity toward oncostatin M receptor and leukemia inhibitory factor receptor. J Biol Chem. 2012;287:32848–59. DOI: 10.1074/ibc.M112.387324.
- Biol Chem. 2012;287:32848–59. DOI: 10.1074/jbc.M112.387324.
  38. Grenier A, Combaux D, Chastre J, et al. Oncostatin M production by blood and alveolar neutrophils during acute lung injury. Lab Invest. 2001;81:133–41. DOI: 10.1038/labinvest.3780220.
- Suda T, Chida K, Todate A, et al. Oncostatin M production by human dendritic cells in response to bacterial products. Cytokine. 2002;17:335–40. DOI: 10.1006/cyto.2002.1023.
- Brown TJ, Lioubin MN, Marquardt H. Purification and characterization of cytostatic lymphokines produced by activated human T lymphocytes. Synergistic antiproliferative activity of transforming growth factor beta 1, interferongamma, and oncostatin M for human melanoma

- cells. J Immunol. 1987;139:2977–83. DOI: 10.4049/immunol.139.9.2977.
- Sato F, Miyaoka Y, Miyajima A, Tanaka M. Oncostatin M maintains the hematopoietic microenvironment in the bone marrow by modulating adipogenesis and osteogenesis. *PLoS One*. 2014;9:e116209. DOI: 10.1371/journal.pone.0116209.
   Ruprecht K, Kuhlmann T, Seif F, et al. Effects of oncostatin
- Ruprecht K, Kuhlmann T, Seif F, et al. Effects of oncostatin M on human cerebral endothelial cells and expression in inflammatory brain lesions. J Neuropathol Exp Neurol. 2001;60:1087–98. DOI: 10.1093/jnen/60.11.1087.
- Moidunny S, Matos M, Wesseling E, et al. Oncostatin M promotes excitotoxicity by inhibiting glutamate uptake in astrocytes: Implications in HIV-associated neurotoxicity. J Neuroinflammation. 2016;13:144. DOI: 10.1186/s12974-016-0613-8
- Znoyko I, Sohara N, Spicer SS, et al. Comparative studies of oncostatin M expression in the tissues of adult rodents. Anat Rec A Discov Mol Cell Evol Biol. 2005;283:182–6. DOI: 10.1002/ 262-20150.
- Houben E, Hellings N, Broux B. Oncostatin M, an underestimated player in the central nervous system. Front Immunol. 2019;10:1165. DOI: 10.3389/fimmu.2019.01165.
- Morshead CM, Reynolds BA, Craig CG, et al. Neural stem cells in the adult mammalian forebrain: A relatively quiescent subpopulation of subependymal cells. Neuron. 1994;13:1071–82. DOI: 10.1016/0896-6273(94)90046-9.
- Bull ND, Bartlett PF. The adult mouse hippocampal progenitor is neurogenic but not a stem cell. J Neurosci. 2005;25:10815–21. DOI: 10.1523/JNEUROSCI.3249-05.2005.
- Liu Z, Martin LI. Olfactory bulb core is a rich source of neural progenitor and stem cells in adult rodent and human. J Comp Neurol. 2003;459:368–91. DOI: 10.1002/cne.10664.
- Morikawa Y. Oncostatin M in the development of the nervous system. *Anat Sci Int*. 2005;80:53–9. DOI: 10.1111/j.1447-073x.2005.00100.x.
- Weiss TW, Samson AL, Niego B, et al. Oncostatin M is a neuroprotective cytokine that inhibits excitotoxic injury in vitro and in vivo. FASEB J. 2006;20:2369–71. DOI: 10.1096/fj.06-5850fie
- 51. Slaets H, Nelissen S, Janssens K, et al. Oncostatin M reduces lesion size and promotes functional recovery and

- neurite outgrowth after spinal cord injury. *Mol Neurobiol*. 2014;50:1142–51. DOI: 10.1007/s12035-014-8795-5.
- Ensoli F, Fiorelli V, DeCristofaro M, et al. Inflammatory cytokines and HIV-1-associated neurodegeneration: Oncostatin-M produced by mononuclear cells from HIV-1-infected individuals induces apoptosis of primary neurons. *J Immunol*. 1999;162:6268–77. DOI: 10.4049/jimmunol.162.10.6268.
- Wallace PM, MacMaster JF, Rouleau KA, et al. Regulation of inflammatory responses by oncostatin M. J Immunol. 1999:162:5547–55. DOI: 10.4049/iimmunol.162.9.5547.
- 1999;162:5547–55. DOI: 10.4049/jimmunol.162.9.5547.
   Wahl AF, Wallace PM. Oncostatin M in the anti-inflammatory response. *Ann Rheum Dis*. 2001;60 Suppl 3:iii75–80. DOI: 10.1136/ard.60.90003.iii75.
- Shin SH, Han SK, Jeong SH, Kim WK. Potential of oncostatin M to accelerate diabetic wound healing. *Int Wound J.* 2014;11:398–403. DOI: 10.1111/j.1742-481X.2012.01107.x.
- Pohin M, Guesdon W, Mekouo AAT, et al. Oncostatin M overexpression induces skin inflammation but is not required in the mouse model of Imiquimod-induced psoriasis-like inflammation. Eur J Immunol. 2016;46:1737–51. DOI: 10.1002/ eli 201546216.
- Loddick SA, Turnbull AV, Rothwell NJ. Cerebral interleukin-6 is neuroprotective during permanent focal cerebral ischemia in the rat. *Jereb Blood Flow Metab*. 1998;18:176–9. DOI: 10.1097/00004647-199802000-00008.
- Hsu MP, Frausto R, Rose-John S, Campbell IL. Analysis of IL-6/ Gp130 family receptor expression reveals that in contrast to astroglia, microglia lack the oncostatin M receptor and functional responses to oncostatin M. Glia. 2015;63:132–41. DOI: 10.1002/glia.22739.
- Chen Y-C, Ma N-X, Pei Z-F, et al. A Neurod1 AAV-based gene therapy for functional brain repair after ischemic injury through in vivo Astrocyte-to-neuron conversion. Mol Ther. 2020;28:217–34. DOI: 10.1016/j.ymthe.2019.09.003
- 2020;28:217–34. DOI: 10.1016/j.ymthe.2019.09.003.
  60. Hu X, Wu D, He X, et al. CircGSK3ß promotes metastasis in esophageal squamous cell carcinoma by augmenting B-catenin signaling. *Mol Cancer*. 2019;18:160. DOI: 10.1186/s12943-019-1095-v
- Ren X, Hu H, Farooqi I, Simpkins JW. Blood substitution therapy rescues the brain of mice from ischemic damage. Nat Commun. 2020;11:4078. DOI: 10.1038/s41467-020-17930-x.

36 touchREVIEWS in Neurology