

1 genes encoding glycolytic enzymes (*SLC2A3*, *TPI*, and *PGK1*) whereas the expression of these genes was
2 suppressed by Notch inhibition. In addition, Hes1 transduction induced mRNA expression of the same genes
3 (Figure 6). It was previously reported that *SLC2A3* expression was regulated by p65/NF- κ B signaling, and that
4 Notch/Hes1 is able to induce the activation of the NF- κ B pathway in human T-ALL lines and animal disease
5 models [54]. Espinosa et al. demonstrated that Hes1 directly targeted the deubiquitinase CYLD, resulting in
6 deubiquitination and inactivation of TAK1 and IKK, degradation of I κ B α , and activation of NF- κ B signaling [54]. In
7 our systems, however, we did not observe repression of *CYLD* mRNA in Hes1-overexpressing hADMPCs (data
8 not shown). While *PGK1* mRNA has been reported to be upregulated by NF- κ B, it has not clearly been shown to
9 be controlled by NF- κ B despite the presence of a NF- κ B site in the promoter [55]. Although modulation of *TPI*
10 expression by NF- κ B has not been reported, we found several NF- κ B binding sites on the human *TPI* promoter
11 (data not shown). As NF- κ B is likely to be one of the responsible signals for hypoxic-induced glycolysis [54],
12 further analysis will be required to determine the mechanism by which NF- κ B signaling is induced by Notch
13 signaling. Additionally, it will be important to investigate whether NF- κ B is really responsible for the observed
14 glycolysis and whether it regulates the expression of *SLC2A3*, *TPI*, and *PGK1* in hADMPCs under 5% oxygen.

15 In addition, *SCO2*, a positive modulator of aerobic respiration, and *TIGAR*, a negative regulator of
16 glycolysis, were transcriptionally downregulated in the 5% oxygen condition, whereas DAPT treatment partially
17 restored expression (Figure 6B). We observed some glycolysis and mitochondrial enzyme activity and found that
18 the activities of COX IV and PFK were consistent with gene expression data (Figure 6C and D). Adenoviral

1 expression of Hes1 dramatically reduced *SCO2* and *TIGAR* expression (Figure 6E), which suggests that
2 Notch-Hes1 signaling modulates the metabolic pathway. Intriguingly, our results also indicate that Hes1 could
3 suppress the expression of *TIGAR* and *SCO2*, a p53 target gene. It has been reported that Notch signaling
4 suppresses p53 in lymphomagenesis [47]. Moreover, Kim et al. reported that NICD1 inhibits p53 phosphorylation
5 and represses p53 transactivation by interacting with p53 [48]. In addition, DAPT treatment resulted in the
6 enhancement of p53 activity in the hypoxic conditions (Figure 4H and I). Therefore, it is possible that p53
7 activation was regulated by Notch signaling in hADMPCs, although we did not observe a decrease in p53 activity
8 in hypoxic conditions in this study (Figure 4). Further analysis will be required to determine whether p53 activity is
9 suppressed in hypoxic conditions over a longer period of culture.

10 Cells undergoing active proliferation utilize large amounts of glucose through glycolysis, producing
11 pyruvate for use in substrates (amino acids and lipids) and the pentose shunt for use in nucleic acid substrates,
12 and also producing NADPH as a reducing agent to counter oxidative stress [18,56]. In the current study, 5% O₂
13 actually increased proliferation and decreased the accumulation of ROS, which may be involved in the reduction
14 of senescence (Figure 1). Because accumulation of endogenous ROS might be a major reason for replicative
15 senescence [20], enhancing glycolysis in cultured cells may improve the quality of the cells by suppressing
16 premature senescence. Kondoh *et al.* demonstrated that enhanced glycolysis is involved in cellular
17 immortalization through reduction of intrinsic ROS production [14,18,19]. Therefore, it is possible that the
18 extension of lifespan observed in our experimental conditions was caused by the reduction of intracellular ROS

1 levels through enhanced glycolysis by Notch signaling. Our data indicate that aerobic glycolysis is utilized for
2 proliferation of hADMPs because the glycolytic inhibitor 2-DG attenuates the proliferation rate of hADMPs
3 (Figure 7A). Intriguingly, the aerobic respiration block by NaN_3 did not decrease the proliferation; rather, it
4 increased proliferation at a low concentration (Figure 7B), which may support our data indicating that the
5 metabolic switch from mitochondrial respiration to glycolysis provides a growth advantage to hADMPs. However,
6 the question of whether the enhanced glycolysis really contributes to the prolonged lifespan in hADMPs remains
7 to be determined in this study.

8 In the current study, the molecular mechanism for how Notch signaling is activated in 5% O_2
9 conditions was explored. It has been reported that Notch1 activity is influenced by oxygen concentration
10 [41,42,57]. In melanoma cells, hypoxia (2% O_2) resulted in increased expression of Notch1 by HIF-1 α and also by
11 Akt through NF- κ B activity [42]. Similarly, in hypoxic breast cancer cells, Notch ligand JAG2 was shown to be
12 transcriptionally activated by hypoxia (1% O_2) in a HIF-1 α dependent manner, resulting an elevation of Notch
13 signaling [41]. In contrast, in hESCs continuously cultured in 5% O_2 , alteration of the Notch pathway seems to be
14 independent of HIF-1 α [57]. In our system, Notch1 activation was not likely dependent on HIF-1 α and HIF-2 α
15 because these proteins did not accumulate in the Hx condition. In contrast, our results indicate that the 5% O_2
16 condition activated Akt and NF- κ B signaling (Figure 4), which suggests that these molecules may activate Notch
17 signaling in hADMPs. NF- κ B was previously shown to increase Notch1 activity indirectly by increasing the
18 expression of Notch ligand Jagged1 in HeLa, lymphoma, and myeloma cells [58]. In addition, Akt regulated

1 Notch1 by increasing Notch1 transcription through the activity of NF- κ B in melanoma cells [42]. Further analysis is
2 required to clarify the mechanism underlying this phenomenon.

3 In conclusion, the 5% oxygen condition conferred a growth advantage through a metabolic shift to
4 glycolysis, improved the proliferation efficiency, prevented the cellular senescence, and maintained the
5 undifferentiated status of hADMPCs. These observations thus provide new regulatory mechanisms for the
6 maintenance of stemness observed in 5% oxygen conditions. In addition, our study sheds new light on the
7 regulation of replicative senescence, which might have impact for quality control of hADMPC preparations used
8 for therapeutic applications.

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16 17 **Disclosure Statement**

1 The authors declare no conflict of interest. No competing financial interests exist.

3 References

- 4 1. Okura H, H Komoda, A Saga, A Kakuta-Yamamoto, Y Hamada, Y Fumimoto, CM Lee, A Ichinose, Y Sawa
5 and A Matsuyama. (2010). Properties of hepatocyte-like cell clusters from human adipose
6 tissue-derived mesenchymal stem cells. *Tissue engineering. Part C, Methods* 16:761-70.
- 7 2. Okura H, A Matsuyama, CM Lee, A Saga, A Kakuta-Yamamoto, A Nagao, N Sougawa, N Sekiya, K
8 Takekita, Y Shudo, S Miyagawa, H Komoda, T Okano and Y Sawa. (2010). Cardiomyoblast-like cells
9 differentiated from human adipose tissue-derived mesenchymal stem cells improve left ventricular
10 dysfunction and survival in a rat myocardial infarction model. *Tissue engineering. Part C, Methods*
11 16:417-25.
- 12 3. Okura H, H Komoda, Y Fumimoto, CM Lee, T Nishida, Y Sawa and A Matsuyama. (2009).
13 Transdifferentiation of human adipose tissue-derived stromal cells into insulin-producing clusters.
14 *Journal of artificial organs : the official journal of the Japanese Society for Artificial Organs* 12:123-30.
- 15 4. Safford KM, SD Safford, JM Gimble, AK Shetty and HE Rice. (2004). Characterization of neuronal/glia
16 lial differentiation of murine adipose-derived adult stromal cells. *Experimental neurology* 187:319-28.
- 17 5. Leu S, YC Lin, CM Yuen, CH Yen, YH Kao, CK Sun and HK Yip. (2010). Adipose-derived mesenchymal
18 stem cells markedly attenuate brain infarct size and improve neurological function in rats. *Journal of*
19 *translational medicine* 8:63.
- 20 6. Ikegame Y, K Yamashita, S Hayashi, H Mizuno, M Tawada, F You, K Yamada, Y Tanaka, Y Egashira, S
21 Nakashima, S Yoshimura and T Iwama. (2011). Comparison of mesenchymal stem cells from adipose
22 tissue and bone marrow for ischemic stroke therapy. *Cytotherapy* 13:675-85.
- 23 7. Tan B, Z Luan, X Wei, Y He, G Wei, BH Johnstone, M Farlow and Y Du. (2011). AMP-activated kinase
24 mediates adipose stem cell-stimulated neuritogenesis of PC12 cells. *Neuroscience* 181:40-7.
- 25 8. Reid AJ, M Sun, M Wiberg, S Downes, G Terenghi and PJ Kingham. (2011). Nerve repair with
26 adipose-derived stem cells protects dorsal root ganglia neurons from apoptosis. *Neuroscience*.
- 27 9. Rehman J, D Traktuev, J Li, S Merfeld-Clauss, CJ Temm-Grove, JE Bovenkerk, CL Pell, BH Johnstone, RV
28 Considine and KL March. (2004). Secretion of angiogenic and antiapoptotic factors by human adipose
29 stromal cells. *Circulation* 109:1292-8.
- 30 10. Lee EY, Y Xia, WS Kim, MH Kim, TH Kim, KJ Kim, BS Park and JH Sung. (2009). Hypoxia-enhanced
31 wound-healing function of adipose-derived stem cells: increase in stem cell proliferation and

1 up-regulation of VEGF and bFGF. Wound repair and regeneration : official publication of the Wound
2 Healing Society [and] the European Tissue Repair Society 17:540-7.

- 3 11. Moriyama M, H Moriyama, A Ueda, Y Nishibata, H Okura, A Ichinose, A Matsuyama and T Hayakawa.
4 (2012). Human adipose tissue-derived multilineage progenitor cells exposed to oxidative stress induce
5 neurite outgrowth in PC12 cells through p38 MAPK signaling. BMC Cell Biol 13:21.
- 6 12. Wu H, Z Ye and RI Mahato. (2011). Genetically modified mesenchymal stem cells for improved islet
7 transplantation. Mol Pharm 8:1458-70.
- 8 13. Wagner W, P Horn, M Castoldi, A Diehlmann, S Bork, R Saffrich, V Benes, J Blake, S Pfister, V Eckstein
9 and AD Ho. (2008). Replicative senescence of mesenchymal stem cells: a continuous and organized
10 process. PLoS One 3:e2213.
- 11 14. Kondoh H, ME Leonart, Y Nakashima, M Yokode, M Tanaka, D Bernard, J Gil and D Beach. (2007). A
12 high glycolytic flux supports the proliferative potential of murine embryonic stem cells. Antioxid Redox
13 Signal 9:293-9.
- 14 15. Prigione A, B Fauler, R Lurz, H Lehrach and J Adjaye. (2010). The senescence-related
15 mitochondrial/oxidative stress pathway is repressed in human induced pluripotent stem cells. Stem
16 Cells 28:721-33.
- 17 16. Varum S, AS Rodrigues, MB Moura, O Momcilovic, CA Easley, J Ramalho-Santos, B Van Houten and G
18 Schatten. (2011). Energy metabolism in human pluripotent stem cells and their differentiated
19 counterparts. PLoS One 6:e20914.
- 20 17. Warburg O, F Wind and E Negelein. (1927). The Metabolism of Tumors in the Body. J Gen Physiol
21 8:519-30.
- 22 18. Kondoh H. (2008). Cellular life span and the Warburg effect. Exp Cell Res 314:1923-8.
- 23 19. Kondoh H, ME Leonart, J Gil, J Wang, P Degan, G Peters, D Martinez, A Carnero and D Beach. (2005).
24 Glycolytic enzymes can modulate cellular life span. Cancer Res 65:177-85.
- 25 20. Beckman KB and BN Ames. (1998). The free radical theory of aging matures. Physiol Rev 78:547-81.
- 26 21. Ezashi T, P Das and RM Roberts. (2005). Low O₂ tensions and the prevention of differentiation of hES
27 cells. Proc Natl Acad Sci U S A 102:4783-8.
- 28 22. Forristal CE, KL Wright, NA Hanley, RO Oreffo and FD Houghton. (2010). Hypoxia inducible factors
29 regulate pluripotency and proliferation in human embryonic stem cells cultured at reduced oxygen
30 tensions. Reproduction 139:85-97.
- 31 23. Yoshida Y, K Takahashi, K Okita, T Ichisaka and S Yamanaka. (2009). Hypoxia enhances the generation of
32 induced pluripotent stem cells. Cell Stem Cell 5:237-41.
- 33 24. Takubo K, N Goda, W Yamada, H Iriuchishima, E Ikeda, Y Kubota, H Shima, RS Johnson, A Hirao, M
34 Suematsu and T Suda. (2010). Regulation of the HIF-1alpha level is essential for hematopoietic stem
35 cells. Cell Stem Cell 7:391-402.

- 1 25. Santilli G, G Lamorte, L Carlessi, D Ferrari, L Rota Nodari, E Binda, D Delia, AL Vescovi and L De Filippis.
2 (2010). Mild hypoxia enhances proliferation and multipotency of human neural stem cells. *PLoS One*
3 5:e8575.
- 4 26. Tsai CC, YJ Chen, TL Yew, LL Chen, JY Wang, CH Chiu and SC Hung. (2011). Hypoxia inhibits senescence
5 and maintains mesenchymal stem cell properties through down-regulation of E2A-p21 by HIF-TWIST.
6 *Blood* 117:459-69.
- 7 27. Takubo K, G Nagamatsu, CI Kobayashi, A Nakamura-Ishizu, H Kobayashi, E Ikeda, N Goda, Y Rahimi, RS
8 Johnson, T Soga, A Hirao, M Suematsu and T Suda. (2013). Regulation of glycolysis by pdk functions as a
9 metabolic checkpoint for cell cycle quiescence in hematopoietic stem cells. *Cell Stem Cell* 12:49-61.
- 10 28. Grayson WL, F Zhao, R Izadpanah, B Bunnell and T Ma. (2006). Effects of hypoxia on human
11 mesenchymal stem cell expansion and plasticity in 3D constructs. *J Cell Physiol* 207:331-9.
- 12 29. Wang DW, B Fermor, JM Gimble, HA Awad and F Guilak. (2005). Influence of oxygen on the
13 proliferation and metabolism of adipose derived adult stem cells. *J Cell Physiol* 204:184-91.
- 14 30. Moriyama M, M Osawa, SS Mak, T Ohtsuka, N Yamamoto, H Han, V Delmas, R Kageyama, F Beermann,
15 L Larue and S Nishikawa. (2006). Notch signaling via Hes1 transcription factor maintains survival of
16 melanoblasts and melanocyte stem cells. *J Cell Biol* 173:333-9.
- 17 31. Chiba S. (2006). Notch signaling in stem cell systems. *Stem Cells* 24:2437-47.
- 18 32. Moriyama M, H Moriyama, A Ueda, Y Nishibata, H Okura, A Ichinose, A Matsuyama and T Hayakawa.
19 (2012). Human adipose tissue-derived multilineage progenitor cells exposed to oxidative stress induce
20 neurite outgrowth in PC12 cells through p38 MAPK signaling. *BMC Cell Biol* 13:21.
- 21 33. Okura H, A Saga, Y Fumimoto, M Soeda, M Moriyama, H Moriyama, K Nagai, CM Lee, S Yamashita, A
22 Ichinose, T Hayakawa and A Matsuyama. (2011). Transplantation of human adipose tissue-derived
23 multilineage progenitor cells reduces serum cholesterol in hyperlipidemic Watanabe rabbits. *Tissue Eng*
24 *Part C Methods* 17:145-54.
- 25 34. Saga A, H Okura, M Soeda, J Tani, Y Fumimoto, H Komoda, M Moriyama, H Moriyama, S Yamashita, A
26 Ichinose, T Daimon, T Hayakawa and A Matsuyama. (2011). HMG-CoA reductase inhibitor augments
27 the serum total cholesterol-lowering effect of human adipose tissue-derived multilineage progenitor
28 cells in hyperlipidemic homozygous Watanabe rabbits. *Biochem Biophys Res Commun* 412:50-4.
- 29 35. Moriyama H, M Moriyama, K Sawaragi, H Okura, A Ichinose, A Matsuyama and T Hayakawa. (2013).
30 Tightly regulated and homogeneous transgene expression in human adipose-derived mesenchymal
31 stem cells by lentivirus with tet-off system. *PLoS One* 8:e66274.
- 32 36. Sekiya I, BL Larson, JR Smith, R Pochampally, JG Cui and DJ Prockop. (2002). Expansion of human adult
33 stem cells from bone marrow stroma: conditions that maximize the yields of early progenitors and
34 evaluate their quality. *Stem Cells* 20:530-41.
- 35 37. Wagner W, F Wein, A Seckinger, M Frankhauser, U Wirkner, U Krause, J Blake, C Schwager, V Eckstein,
36 W Ansorge and AD Ho. (2005). Comparative characteristics of mesenchymal stem cells from human
37 bone marrow, adipose tissue, and umbilical cord blood. *Exp Hematol* 33:1402-16.

- 1 38. Hass R, C Kasper, S Bohm and R Jacobs. (2011). Different populations and sources of human
2 mesenchymal stem cells (MSC): A comparison of adult and neonatal tissue-derived MSC. *Cell Commun*
3 *Signal* 9:12.
- 4 39. Gustafsson MV, X Zheng, T Pereira, K Gradin, S Jin, J Lundkvist, JL Ruas, L Poellinger, U Lendahl and M
5 Bondesson. (2005). Hypoxia requires notch signaling to maintain the undifferentiated cell state. *Dev*
6 *Cell* 9:617-28.
- 7 40. Zheng X, S Linke, JM Dias, X Zheng, K Gradin, TP Wallis, BR Hamilton, M Gustafsson, JL Ruas, S Wilkins,
8 RL Bilton, K Brismar, ML Whitelaw, T Pereira, JJ Gorman, J Ericson, DJ Peet, U Lendahl and L Poellinger.
9 (2008). Interaction with factor inhibiting HIF-1 defines an additional mode of cross-coupling between
10 the Notch and hypoxia signaling pathways. *Proc Natl Acad Sci U S A* 105:3368-73.
- 11 41. Pietras A, K von Stedingk, D Lindgren, S Pahlman and H Axelson. (2011). JAG2 induction in hypoxic
12 tumor cells alters Notch signaling and enhances endothelial cell tube formation. *Mol Cancer Res*
13 9:626-36.
- 14 42. Bedogni B, JA Warneke, BJ Nickoloff, AJ Giaccia and MB Powell. (2008). Notch1 is an effector of Akt and
15 hypoxia in melanoma development. *J Clin Invest* 118:3660-70.
- 16 43. Beitner-Johnson D, RT Rust, TC Hsieh and DE Millhorn. (2001). Hypoxia activates Akt and induces
17 phosphorylation of GSK-3 in PC12 cells. *Cell Signal* 13:23-7.
- 18 44. Culver C, A Sundqvist, S Mudie, A Melvin, D Xirodimas and S Rocha. (2010). Mechanism of
19 hypoxia-induced NF-kappaB. *Mol Cell Biol* 30:4901-21.
- 20 45. Rohwer N, C Dame, A Haugstetter, B Wiedenmann, K Detjen, CA Schmitt and T Cramer. (2010).
21 Hypoxia-inducible factor 1alpha determines gastric cancer chemosensitivity via modulation of p53 and
22 NF-kappaB. *PLoS One* 5:e12038.
- 23 46. Espinosa L, S Cathelin, T D'Altri, T Trimarchi, A Statnikov, J Guiu, V Rodilla, J Ingles-Esteve, J Nomdedeu,
24 B Bellosillo, C Besses, O Abdel-Wahab, N Kucine, SC Sun, G Song, CC Mullighan, RL Levine, K Rajewsky, I
25 Aifantis and A Bigas. (2010). The Notch/Hes1 pathway sustains NF-kappaB activation through CYLD
26 repression in T cell leukemia. *Cancer Cell* 18:268-81.
- 27 47. Beverly LJ, DW Felsher and AJ Capobianco. (2005). Suppression of p53 by Notch in lymphomagenesis:
28 implications for initiation and regression. *Cancer Res* 65:7159-68.
- 29 48. Kim SB, GW Chae, J Lee, J Park, H Tak, JH Chung, TG Park, JK Ahn and CO Joe. (2007). Activated Notch1
30 interacts with p53 to inhibit its phosphorylation and transactivation. *Cell Death Differ* 14:982-91.
- 31 49. Landor SK, AP Mutvei, V Mamaeva, S Jin, M Busk, R Borra, TJ Gronroos, P Kronqvist, U Lendahl and CM
32 Sahlgren. (2011). Hypo- and hyperactivated Notch signaling induce a glycolytic switch through distinct
33 mechanisms. *Proc Natl Acad Sci U S A* 108:18814-9.
- 34 50. Ciofani M and JC Zuniga-Pflucker. (2005). Notch promotes survival of pre-T cells at the beta-selection
35 checkpoint by regulating cellular metabolism. *Nat Immunol* 6:881-8.
- 36 51. Welford SM, B Bedogni, K Gradin, L Poellinger, M Broome Powell and AJ Giaccia. (2006). HIF1alpha
37 delays premature senescence through the activation of MIF. *Genes Dev* 20:3366-71.

- 1 52. Zhou D, J Xue, JC Lai, NJ Schork, KP White and GG Haddad. (2008). Mechanisms underlying hypoxia
2 tolerance in *Drosophila melanogaster*: hairy as a metabolic switch. *PLoS Genet* 4:e1000221.
- 3 53. Funes JM, M Quintero, S Henderson, D Martinez, U Qureshi, C Westwood, MO Clements, D Bourboulia,
4 RB Pedley, S Moncada and C Boshoff. (2007). Transformation of human mesenchymal stem cells
5 increases their dependency on oxidative phosphorylation for energy production. *Proc Natl Acad Sci U S*
6 *A* 104:6223-8.
- 7 54. Kawauchi K, K Araki, K Tobiume and N Tanaka. (2008). p53 regulates glucose metabolism through an
8 IKK-NF-kappaB pathway and inhibits cell transformation. *Nat Cell Biol* 10:611-8.
- 9 55. Carter KL, E Cahir-McFarland and E Kieff. (2002). Epstein-barr virus-induced changes in B-lymphocyte
10 gene expression. *J Virol* 76:10427-36.
- 11 56. Ak P and AJ Levine. (2010). p53 and NF-kappaB: different strategies for responding to stress lead to a
12 functional antagonism. *FASEB J* 24:3643-52.
- 13 57. Prasad SM, M Czepiel, C Cetinkaya, K Smigielska, SC Weli, H Lysdahl, A Gabrielsen, K Petersen, N Ehlers,
14 T Fink, SL Minger and V Zachar. (2009). Continuous hypoxic culturing maintains activation of Notch and
15 allows long-term propagation of human embryonic stem cells without spontaneous differentiation. *Cell*
16 *Prolif* 42:63-74.
- 17 58. Bash J, WX Zong, S Banga, A Rivera, DW Ballard, Y Ron and C Gelinis. (1999). Rel/NF-kappaB can trigger
18 the Notch signaling pathway by inducing the expression of Jagged1, a ligand for Notch receptors.
19 *EMBO J* 18:2803-11.
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- 21

1 **Figure legends**

2 **Figure 1. Hypoxia increases proliferation capacity and decreases senescence in hADMPs.** (A) Growth
3 profiles of hADMPs under normoxic (red square) and hypoxic (blue square) conditions. The population doubling
4 level (PDL) was determined to be 0 when cells were isolated from human adipose tissue. Cells were maintained
5 until they reached PDL13–15 (passage 3) and then split into four aliquots of equal cell densities. PDL was
6 calculated based on the total cell number at each passage. (B) Detection of normoxic (Nx) and hypoxic (Hx) cells
7 by flow cytometry following incorporation of EdU. (C) Percentages of apoptotic cells with sub-G1 DNA under Nx
8 and Hx conditions. The results are presented as the mean of 3 independent experiments. (D) hADMPs cultured
9 under Nx and Hx conditions were harvested by trypsin-EDTA and then imaged using a phase-contrast
10 microscope. Arrowheads indicate cells with a larger and more irregular shape. (E) Cells expanded under Nx and
11 Hx conditions were stained with SA- β -gal. (F) Cellular ROS detection by the oxidative stress indicator
12 CM-H2DCFDA in hADMPs under Nx or Hx. Data are presented as the mean fluorescence intensity of 3
13 independent experiments. Error bars indicate SD. *P < 0.05 and **P < 0.01 indicate significant difference
14 (independent *t*-test) between Nx and Hx. Scale bars; 100 μ m.

15
16 **Figure 2. Hypoxic culture maintains mesenchymal stem cell properties.** hADMPs cultured under normoxia
17 (20% O₂) or hypoxia (5% O₂) were labeled with antibodies against the indicated antigens and analyzed by flow
18 cytometry. Representative histograms are shown. The respective isotype control is shown as a gray line.

1
2 **Figure 3. Hypoxic culture enhances stem cell properties.** hADMPCs were expanded under normoxic and
3 hypoxic conditions. **(A)** Normoxic (20% O₂) and hypoxic (5% O₂) cells at passage 8 were induced for 3 weeks to
4 differentiate into osteoblasts and adipocytes and stained with alizarin red and Oil-red O, respectively. The stained
5 dye was extracted and OD values were measured and plotted as the means of 3 independent experiments ± SD.
6 *P < 0.05. Scale bars, 200 μm. **(B)** Normoxic (20% O₂) and hypoxic (5% O₂) cells at passage 8 were induced for 3
7 weeks to differentiate to chondrocytes, and immunofluorescent analysis of collagen II (red) and Alucian blue
8 staining were performed. The blue signals indicate nuclear staining. Scale bars, 100 μm. Non-induced control
9 cultures in growth medium without adipogenic, osteogenic or chondrogenic differentiation stimuli are shown
10 (Undifferentiated).

11
12 **Figure 4. Hypoxic culture condition activates Notch signaling but not HIF proteins.** hADMPCs were
13 expanded under normoxic (20% O₂) and hypoxic (5% O₂) conditions. DAPT (1 μM) was added to inhibit Notch
14 signaling. **(A)** Western blot analysis of intracellular domain of Notch1 (Notch1 ICD) expression. Actin served as
15 the loading control. Numbers below blots indicate relative band intensities as determined by ImageJ software. **(B)**
16 Q-PCR analysis of *HES1*. Each expression value was calculated with the $\Delta\Delta C_t$ method using *UBE2D2* as an
17 internal control. **(C)** Western blot analysis of *HES1* in nuclear fractions of hADMPCs. Lamin A/C served as the
18 loading control. **(D, E)** Western blot analysis of HIF-1 α **(D)** and HIF-2 α **(E)**. Cobalt chloride (CoCl₂) was added at a
19 concentration of 100 μM to stabilize HIF proteins (positive control). **(F)** Western blot analysis of phosphorylated

1 Akt (p-Akt) and Akt. Actin served as the loading control. Numbers below blots indicate relative band intensities as
2 determined by ImageJ software. (G) Western blot analysis of nuclear localization of p65. Lamin A/C served as the
3 loading control. Numbers below blots indicate relative band intensities as determined by ImageJ software. (H)
4 Western blot analysis of phosphorylated p53 (p-p53) and p53. Actin served as the loading control. (I) Activity of
5 p53 was measured by the p53-luciferase reporter assay. Relative luciferase activity was determined from 3
6 independent experiments and normalized to pGL4.74 activity.

7
8 **Figure 5. Notch signaling is indispensable for acquisition of the advantageous properties of hADMPCs.**

9 hADMPCs were expanded under normoxic (20% O₂; Nx) and hypoxic (5% O₂; Hx) conditions. DAPT (1 μM) was
10 added to inhibit Notch signaling. (A) Growth profiles of hADMPCs under Nx (red) and Hx (blue) conditions. Solid
11 lines represent control cells and dotted lines represent DAPT-treated cells. The number of population doublings
12 was calculated based on the total cell number at each passage. (B) Percentages of apoptotic cells with sub-G1
13 DNA. Results are presented as the mean of 3 independent experiments ± SD. (C-D) hADMPCs at passage 8
14 were induced for 3 weeks to differentiate into adipocytes (C) and osteoblasts (D) and stained with Oil Red O and
15 Alizarin Red, respectively. The stained dye was extracted, and OD values were measured and plotted as the
16 means of 3 independent experiments ± SD. (E) hADMPCs at passage 8 were induced for 3 weeks to differentiate
17 into chondrocytes, and an immunofluorescent analysis of collagen II (red) was performed. The blue signals
18 indicate nuclear staining. (F) hADMPCs were stained with SA-β-gal. *P < 0.05 and **P < 0.01 indicate significant
19 difference (independent *t*-test) between Nx and Hx. Scale bars; 100 μm.

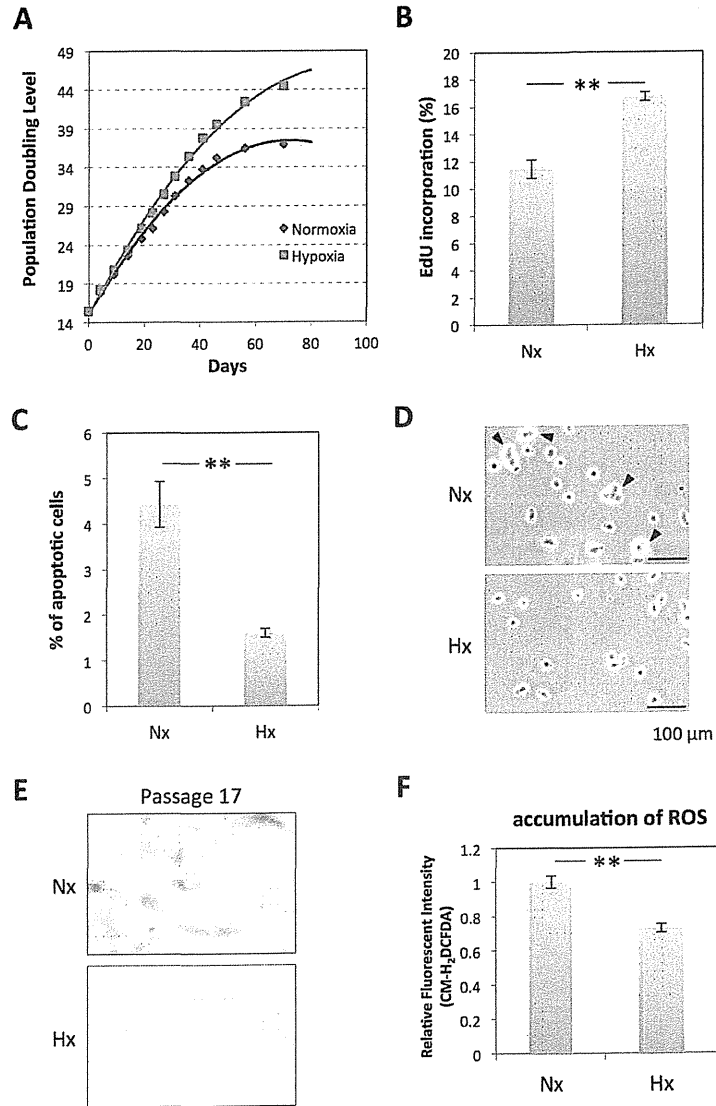
1
2 **Figure 6. Glycolysis is enhanced under 5% oxygen through Notch signaling.** (A-D) hADMPCs were
3 expanded under normoxic (20% O₂) and hypoxic (5% O₂) conditions. DAPT (1 μM) was added in to inhibit Notch
4 signaling. (A) Glucose consumption and lactate production of hADMPCs were measured and plotted as the
5 means of 3 independent experiments ± SD. (B) Relative mRNA expression of *SLC2A3*, *TPI*, *PGK1*, *TIGAR*, and
6 *SCO2* in hADMPCs. Each expression value was calculated with the $\Delta\Delta C_t$ method using *UBE2D2* as an internal
7 control. (C, D) Hexokinase (HK), phosphofructokinase (PFK), lactate dehydrogenase (LDH) (C), pyruvate
8 dehydrogenase (PDH), and Complex IV (Cox IV) (D) activities were measured and the value of relative activity
9 was plotted as the means of 3 independent experiments ± SD. (E, F) hADMPCs were transduced with either
10 mock (Cont) or HES1 and then cultured for 3 days. (E) Relative mRNA expression of *SLC2A3*, *TPI*, *PGK1*,
11 *TIGAR*, and *SCO2* in hADMPCs. Each expression value was calculated with the $\Delta\Delta C_t$ method using *UBE2D2* as
12 an internal control. (F) Glucose consumption and lactate production of hADMPCs were measured and plotted as
13 the means of 3 independent experiments ± SD. (G) hADMPCs were transduced with either scrambled control
14 RNAi (Cont) or RNAi against HES1 (HES1-KD), and then cultured for 3 days. Glucose consumption and lactate
15 production of hADMPCs were measured and plotted as the means of 3 independent experiments ± SD. **P <
16 0.01. * 0.01 < P < 0.05.

17
18 **Figure 7. Glycolysis supports proliferation of hADMPCs.** hADMPCs were treated with 0, 0.2, 0.4 and 1 mM
19 2-deoxy-D-glucose (2-DG) (A) or 0, 1 and 5 mM sodium azide (NaN₃) (B) for 24 h. Cells were then allowed to

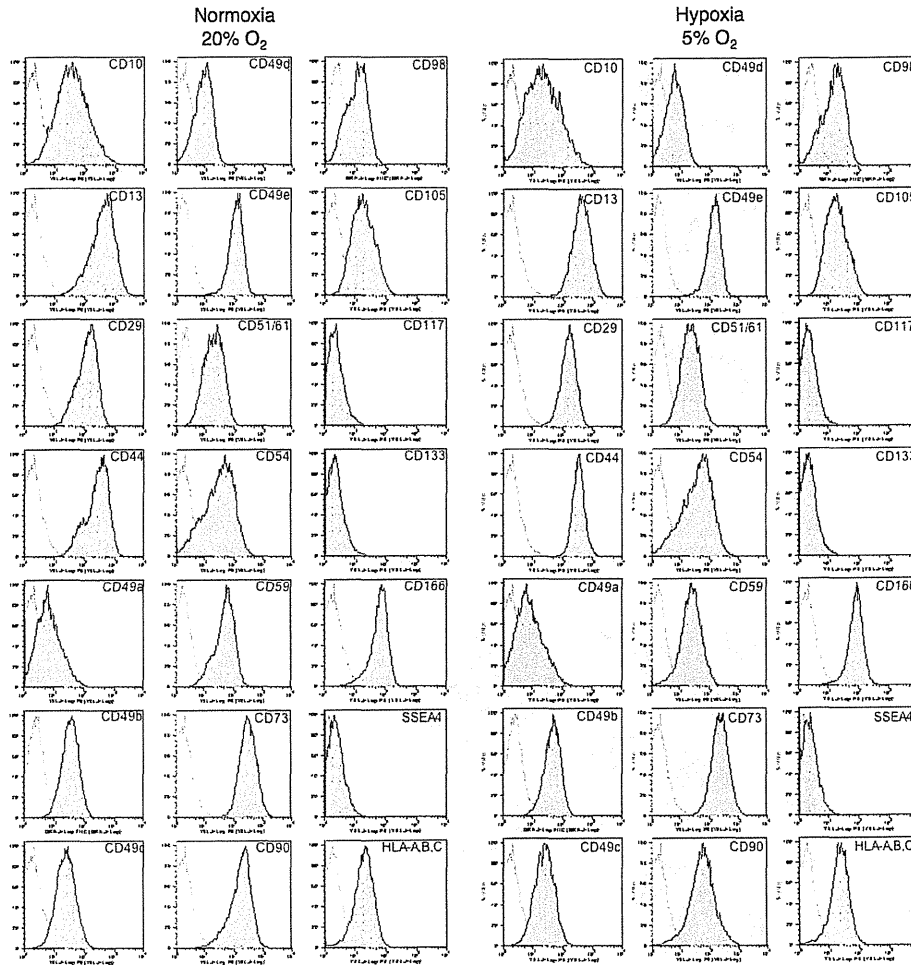
1 incorporate EdU for 2 h, and the EdU-positive cells were analyzed by flow cytometry. The percentages for the 0
2 mM control were plotted as the means of 3 independent experiments \pm SD. ** $P < 0.01$. * $0.01 < P < 0.05$.

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MoriyamaFig1
170x237mm (300 x 300 DPI)



MoriyamaFig2
169x174mm (300 x 300 DPI)

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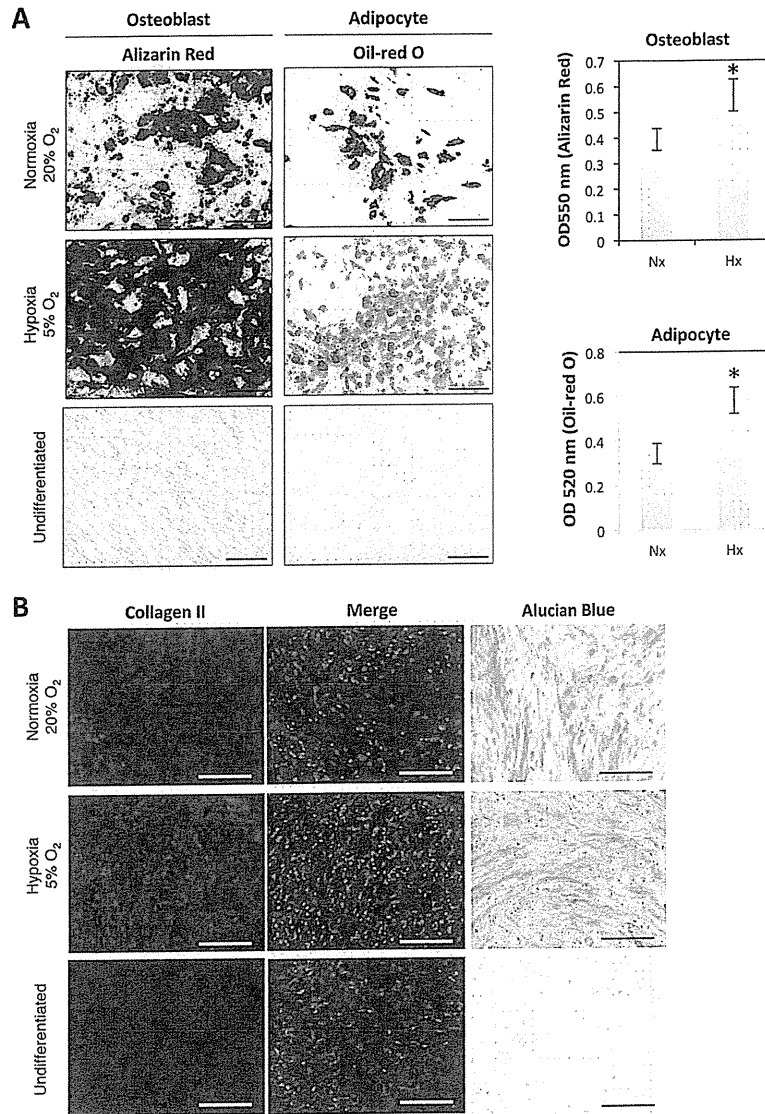
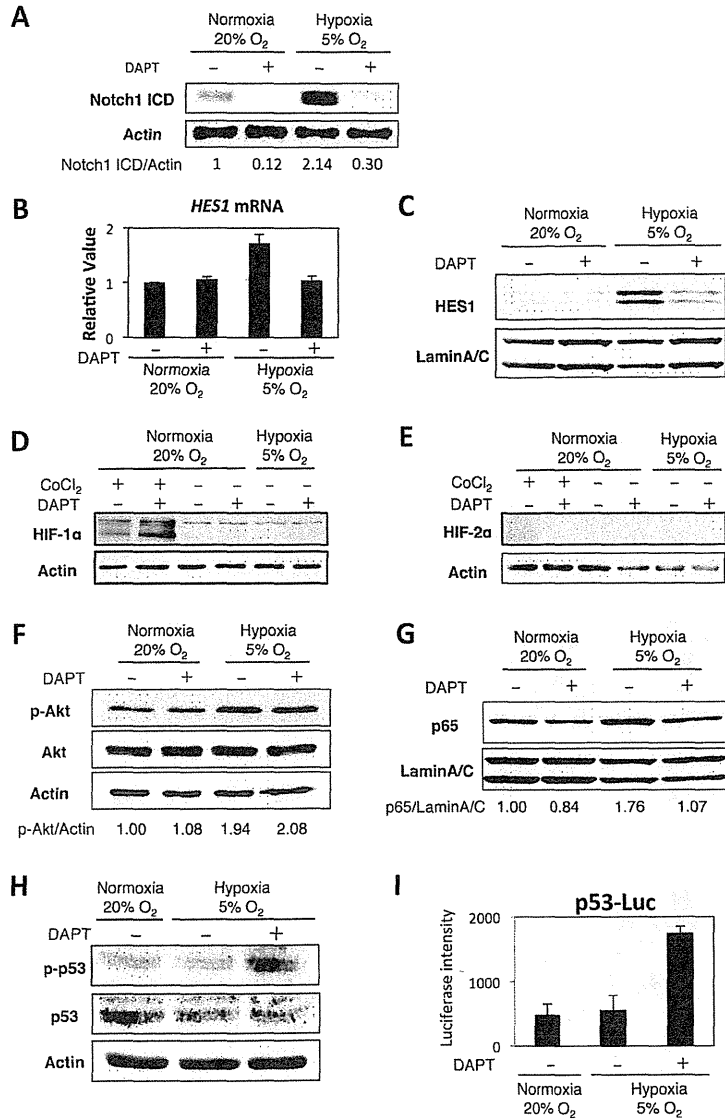
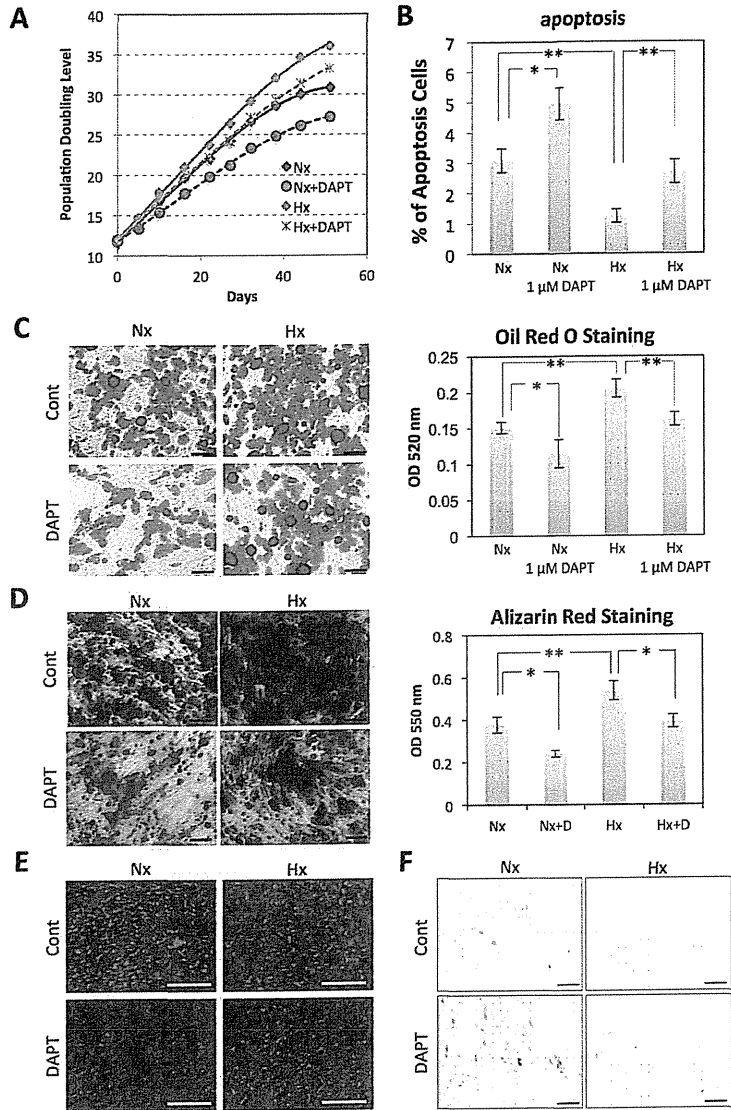


Figure 3
101x144mm (300 x 300 DPI)

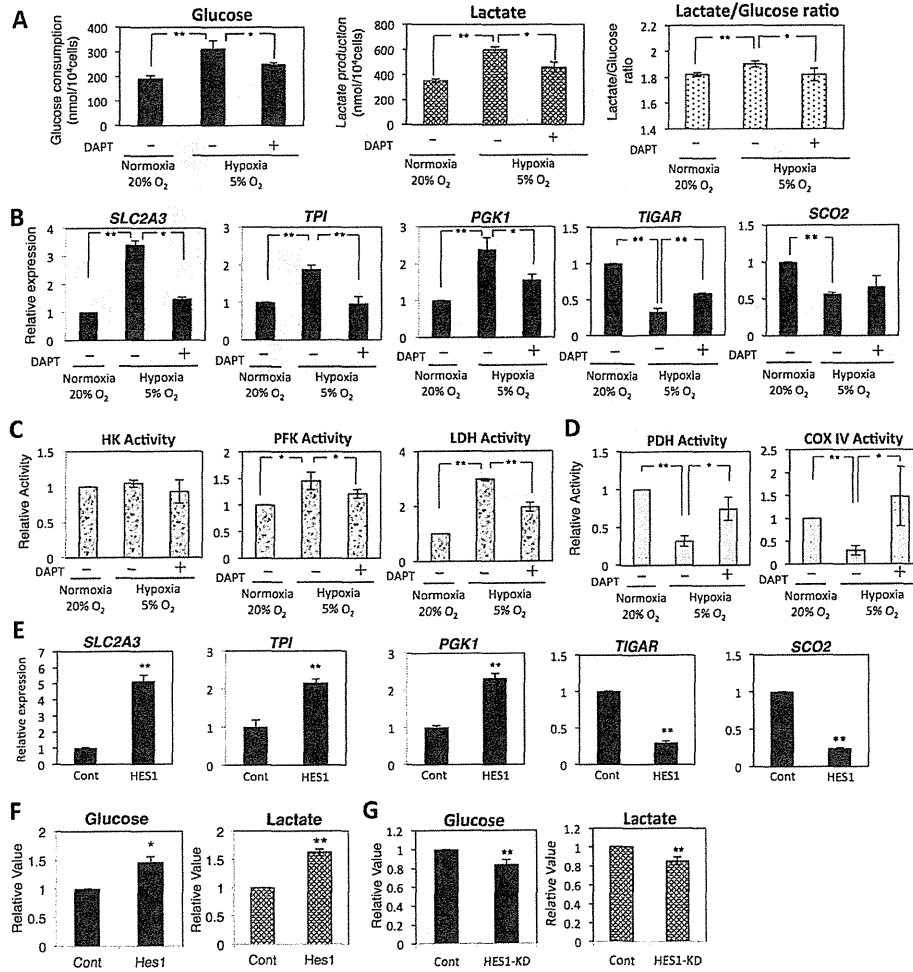


MoriyamaFig4
163x246mm (300 x 300 DPI)

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MoriyamaFig5
171x247mm (300 x 300 DPI)



MoriyamaFig6
169x176mm (300 x 300 DPI)