Polyamine-controlled proliferation and protein biosynthesis are independent

- 2 determinants of hair follicle stem cell fate
- 3 Short Title: Polyamine control of stem cell fate
- 5 Kira Allmeroth¹, Christine S. Kim¹, Andrea Annibal¹, Andromachi Pouikli¹, Carlos
- 6 Andrés Chacón-Martínez¹, Christian Latza¹, Adam Antebi^{1,2}, Peter Tessarz^{1,2}, Sara A.
- 7 Wickström^{1,2,4,5,6} & Martin S. Denzel^{1,2,3*}
- ⁹ Max Planck Institute for Biology of Ageing, Joseph-Stelzmann-Str. 9b, D-50931
- 10 Cologne, Germany

1

4

8

11

14

17

20

23

26

- 12 ²CECAD Cluster of Excellence, University of Cologne, Joseph-Stelzmann-Str. 26, D-
- 13 50931 Cologne, Germany
- 15 ³Center for Molecular Medicine Cologne (CMMC), University of Cologne, Robert-Koch-
- 16 Str. 21, D-50931 Cologne, Germany
- ⁴Helsinki Institute for Life Science, Biomedicum Helsinki, Haartmaninkatu 8, FI-00290
- 19 Helsinki, Finland
- 21 ⁵Wihuri Research Institute, Biomedicum Helsinki, Haartmaninkatu 8, FI-00290
- 22 Helsinki, Finland
- ⁶Stem Cells and Metabolism Research Program, Faculty of Medicine, Yliopistonkatu
- 25 3, FI-00014 University of Helsinki, Finland
- 28 * Corresponding author: Martin S. Denzel
- 29 E-Mail: martin.denzel@age.mpg.de
- 30 phone: +49-(0)221 379 70 443
- 31 fax: +49-(0)221 379 70 88-443

Abstract

Stem cell differentiation is accompanied by an increase in mRNA translation. The rate of protein biosynthesis is influenced by the polyamines putrescine, spermidine, and spermine that are essential for cell growth and stem cell maintenance. However, the role of polyamines as endogenous effectors of stem cell fate and whether they act through translational control remains obscure. Here, we investigated the function of polyamines in stem cell fate decisions using hair follicle stem cell (HFSC) organoids. HFSCs showed lower translation rates than progenitor cells, and a forced suppression of translation by direct targeting of the ribosome or through specific depletion of natural polyamines elevated stemness. In addition, we identified N1-acetylspermidine as a novel parallel regulator of cell fate decisions, increasing proliferation without reducing translation. Overall, this study delineates the diverse routes of polyamine metabolism-mediated regulation of stem cell fate decisions.

Key Points

- Low mRNA translation rates characterize hair follicle stem cell (HFSC) state
- Depletion of natural polyamines enriches HFSCs via reduced translation
- N1-acetylspermidine promotes HFSC state without reducing translation
- N1-acetylspermidine expands the stem cell pool through elevated proliferation

Key Words

Hair follicle stem cells / mRNA translation / polyamines / cell fate

Introduction

54

55

56

57

58

59

60

61

62

63

64

65

66

67

68

69

70

71

72

73

74

75

76

77

78

79

Translation of messenger RNAs (mRNAs) is one of the most complex and energyconsuming processes in the cell (Roux & Topisirovic, 2018). It comprises initiation, elongation, termination, and ribosome recycling and plays a key role in gene expression regulation (Hershey et al., 2019). The rate of protein synthesis is tightly controlled by signaling pathways that sense various internal and external stimuli (Roux & Topisirovic, 2018). Deregulation of translation can manifest in a variety of diseases, including neurodegeneration and cancer (Tahmasebi et al., 2018b). The regulation of mRNA translation has also been implicated in early cell fate transitions (Ingolia et al., 2011, Kristensen et al., 2013, Lu et al., 2009). Several studies in both embryonic and somatic stem cells demonstrated that global translation is suppressed in stem cells to retain an undifferentiated state and increased in progenitor cells (Sampath et al., 2008, Signer et al., 2014, Zismanov et al., 2016). Hair follicle stem cells (HFSCs) represent an excellent paradigm to study adult somatic stem cell fate decisions, since they fuel cyclical rounds of hair follicle regeneration during the natural hair cycle (Blanpain & Fuchs, 2009, Fuchs et al., 2001). Interestingly, Blanco et al. (2016) showed that the activation of HFSCs during the transition from the resting phase, telogen, to proliferation in anagen during the hair cycle also coincides with increased translation. These data demonstrate a functional role for increased translation during differentiation in the epidermis. However, the mechanisms that regulate translation in stem cells in general and upon differentiation remain poorly understood. One important determinant of translation rates is the availability of the natural polyamines putrescine, spermidine, and spermine. These polycations are essential for cell growth and can bind to a variety of negatively charged cellular molecules, including nucleic acids (Dever & Ivanov, 2018). Polyamine homeostasis is critical to cell survival

81

82

83

84

85

86

87

88

89

90

91

92

93

94

95

96

97

98

99

100

101

102

103

104

and is achieved by coordinated regulation at different levels of their synthesis, degradation, uptake and excretion (Wallace et al., 2003). Interestingly, up to 15% of polyamines are stably associated with ribosomes in Escherichia coli, highlighting their importance in protein synthesis (Cohen & Lichtenstein, 1960). Additionally, the polyamine spermidine is converted to the amino acid hypusine, which posttranslationally modifies the elongation factor eIF5A (Park et al., 1981). This modification is critical for translation elongation, especially for difficult substrates like polyproline (Gutierrez et al., 2013, Saini et al., 2009). Given that increased polyamine levels positively regulate translation and that high translation rates promote stem cell differentiation, it is surprising that previous studies have implicated elevated polyamine levels in stem cell maintenance. For example, polyamines positively influence expression of MINDY1, a deubiquitinating enzyme, which promotes embryonic stem cell (ESC) self-renewal (James et al., 2018). Additionally, forced overexpression of the rate-limiting enzymes in polyamine biosynthesis adenosylmethionine decarboxylase (AMD1) or ornithine decarboxylase (ODC) in ESCs results in delayed differentiation upon removal of LIF and improves cellular reprogramming (Zhang et al., 2012, Zhao et al., 2012). In line with these observations, differentiation of human bone marrow-derived mesenchymal stem cells coincides with decreased polyamine levels (Tsai et al., 2015). Overall, these data suggest that changes in polyamine levels can affect cell fate. However, the link between polyamines and translation in cell fate decisions remains unclear. Do low polyamine levels endogenously reduce translation rates in stem cells? Does reduction of polyamine levels, and translation, affect stem cell maintenance and function? Might distinct polyamine species differ in their effects on stem cells through translation or other mechanisms?

To address these questions, we used an *in vitro* organoid HFSC culture system and investigated polyamines as key regulators of mRNA translation in stem cell fate decisions. We confirm reduced translation as a key property of HFSCs and demonstrate that forced reduction of translation promotes the stem cell state. We find that HFSCs display low levels of natural polyamines, which implicates polyamines as endogenous gatekeepers of translation in stem cells. Accordingly, specific depletion of natural polyamines using the polyamine analogue N1,N11-diethylnorspermine (DENSpm) improved stem cell maintenance through reduced translation. Surprisingly, depletion of all polyamines by difluoromethylornithine (DFMO) does not affect cell fate. Therefore, we hypothesized that the polyamine pathway can also modulate cell fate independently of reduced translation. Intriguingly, we identify a new function for N1-acteylspermidine (N1-AcSpd) in promoting proliferation, expanding the stem cell pool. Taken together, our data demonstrate that polyamines play a central role in stem cell fate decisions by controlling the key processes translation and proliferation to ensure stem cell maintenance and tissue homeostasis.

Results

Low translation rates mark the HFSC state and decreasing translation enhances

stemness in the 3D-3C organoids.

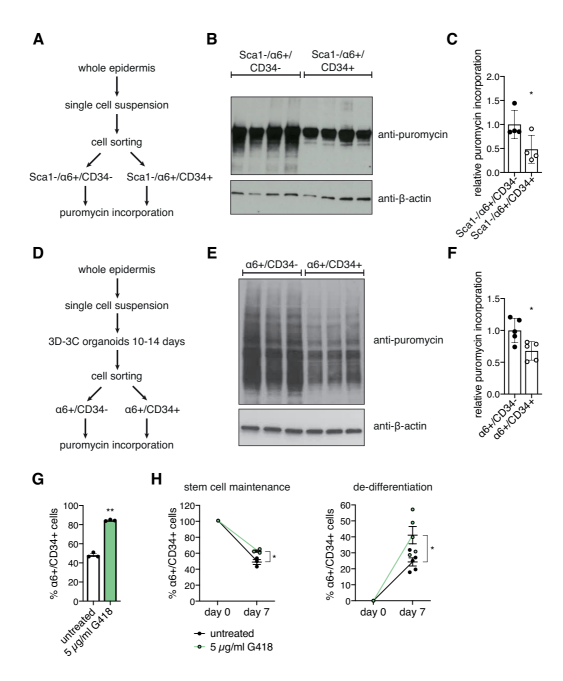


Figure 1: Low translation rates mark the HFSC state and decreasing translation enhances stemness in the 3D-3C organoids. (A) Schematic representation of the workflow for puromycin incorporation using freshly isolated cells. (B) Western blot analysis after puromycin incorporation in Sca1-/ α 6+/CD34- progenitor cells compared to Sca1-/ α 6+/CD34+ hair follicle stem cells (HFSCs). (C) Quantification of the Western blot in (B). Mean \pm SD (n=4); *p<0.05 (t test). (D) Schematic representation of the workflow for puromycin incorporation using 3D-3C cultured cells. (E) Representative Western blot analysis after puromycin incorporation in α 6+/CD34- progenitor cells compared to α 6+/CD34+ HFSCs. (F) Quantification of the Western blot in (E). Mean \pm SD (n=5); *p<0.05 (t test). (G) Ratio of α 6+/CD34+ cells after two weeks of 3D-3C culture with or without G418 treatment. Mean \pm SEM (n=3); **p<0.01 (t test). (H) Ratio of α 6+/CD34+ cells at day 0 and day 7 post-

135

136

137

138

139

140

141

142

143

144

145

146

147

148

149

150

151

152

153

154

155

156

157

158

159

sorting starting from 100 % α 6+/CD34+ cells (left) or 100 % α 6+/CD34- cells (right) with or without G418 treatment. Mean \pm SEM (n \geq 3); * p<0.05 (t test). It has been previously described that stem cells display lower translation rates than their differentiated counterparts (Tahmasebi et al., 2018a). To test if this was also true in epidermal stem cells, we sorted freshly isolated mouse epidermal cells using three different markers (Fig. 1A): α 6 integrin is expressed in all progenitors both in the interfollicular epidermis (IFE) and the hair follicle (HF) (Li et al., 1998, Sonnenberg et al., 1991); the hematopoietic stem cell marker CD34 marks the bulge stem cells of the HF (Trempus et al., 2003); stem cell antigen-1 (Sca1) expression can be detected in the infundibulum (IFD) region of the HF and the basal layer of the epidermis (Jensen et al., 2008). By gating for Sca1 negative cells, IFE and IFD progenitors can be separated from HF progenitors. Thus, we compared HF bulge stem cells (Sca1- $/\alpha6+/CD34+$) to HF outer root sheath cells (Sca1- $/\alpha6+/CD34-$). We performed puromycin incorporation to quantify the amount of newly synthesized proteins (Schmidt et al., 2009). Western blot analysis revealed a significant reduction of puromycin incorporation in Sca1- $/\alpha$ 6+/CD34+ stem cells (Fig. 1B-C), confirming reduced translation rates in HFSCs in vivo. To understand the mechanisms and functional consequences of this difference in translation, we made use of an ex vivo organoid culture system (3D-3C culture) established by Chacon-Martinez et al. (2017) allowing for the long-term culture and manipulation of HFSCs and their direct progeny. Importantly, in this culture cells maintain self-renewal capacity and multipotency and respond to the same signals as they do in vivo. In the 3D-3C organoids, a balance between α6+/CD34+ HFSCs and α 6+/CD34- progenitor cells is formed in a self-driven process. Based on their transcriptome and marker expression analysis, these progenitor cells represent HF outer root sheath (ORS) cells and inner bulge cells (Kim et al., 2019), both of which

161

162

163

164

165

166

167

168

169

170

171

172

173

174

175

176

177

178

179

180

181

182

183

184

represent HFSC progeny and act as niche cells for HFSCs in vivo (Hsu et al., 2014). To confirm cell identity of $\alpha6+/CD34+$ HFSCs and $\alpha6+/CD34-$ ORS progenitors, we sorted cells after two weeks of 3D-3C culture and analyzed gene expression by RNAsequencing. Comparing the two populations, the most enriched GO term was skin development (Fig. S1A). As expected, the differentiation markers Krt1 and Krt10 were enriched in progenitor cells, while the expression of the stem cell marker genes Id2, Lhx2, and Sox9 was higher in HFSCs (Fig. S1B). Thus, α 6+/CD34- progenitors and α6+/CD34+ HFSCs are clearly distinguishable at the gene expression level and represent distinct cellular states. Next, we investigated puromycin incorporation in the two cell populations after sorting and found reduced translation in α 6+/CD34+ HFSCs (Fig. 1D-F). Thus, the 3D-3C organoids maintain this key in vivo property of HFSCs, making them a suitable model to study the influence of translation on cell fate. Next, we asked whether reduced translation might drive cell fate decisions in HFSCs. To this end, we manipulated translation in the 3D-3C organoids by addition of low doses of G418, a well described inhibitor of translation elongation (Bar-Nun et al., 1983). We confirmed reduced translation after 4 h of G418 treatment in the 3D-3C organoids by measuring puromycin incorporation (Fig. S1C-D). Strikingly, a forced decrease in translation resulted in a significant increase of α 6+/CD34+ stem cells after two weeks of 3D-3C culture (Fig. 1G). To understand whether this increase had a functional consequence, we analyzed stem cell potency in a colony formation assay. For this, cells were seeded in clonal density on a feeder layer and cultured for 2 to 3 weeks (Fig. S1E). Quantification of total colony number revealed a significant increase upon G418 treatment in the organoid culture, indicating that inhibiting translation increased stem cell proliferative potential (Fig. S1F-G). The self-organizing plasticity of the organoids results in a balance between α 6+/CD34- and α 6+/CD34+ cells

186

187

188

189

190

191

192

193

194

195

196

197

198

199

200

201

202

203

204

205

206

207

208

209

210

(Chacon-Martinez et al., 2017). This balance is influenced by self-renewal and differentiation of HFSCs but also by proliferation and de-differentiation of progenitors back to the stem cell state. Thus, to investigate if the increase in stem cell potency was accompanied by increased self-renewal and de-differentiation in the G418-treated organoids, we sorted the cells after two weeks of 3D-3C culture to generate pure populations of either α 6+/CD34- or α 6+/CD34+ cells. These pure populations were cultured for one week with or without G418 treatment before analysis. Interestingly, G418 treatment prevented stem cell differentiation and promoted de-differentiation of progenitors back to the stem cell state (Fig. 1H). Taken together, these data suggest that reduced translation is not merely a consequence of stemness. Instead, decreasing translation actively promotes the stem cell state. Depletion of natural polyamines is a feature of stem cells and can drive stemness in vitro. To further analyze the effect of decreased translation rates on cell fate, we focused on the polyamine pathway (Fig. 2A). Reduction of the natural polyamines putrescine, spermidine, and spermine has been implicated in reduced translation (Dever & Ivanov, 2018, Pegg, 2016). We thus asked whether stem cells might display lower polyamine levels compared to their differentiated counterparts. Indeed, when we sorted cells after two weeks of 3D-3C culture and measured polyamines by LC-MS, we found reduced levels of putrescine, spermidine, and spermine in α 6+/CD34+ HFSCs compared to α 6+/CD34- progenitors (Fig. 2B). To probe polyamine function in stem cell fate decisions, we manipulated their levels using the spermine analogue N1,N11-diethylnorspermine (DENSpm), which increases spermidine/spermine N1-acetyltransferase (SSAT) expression (Coleman et al., 1995, Fogel-Petrovic et al., 1996, Parry et al., 1995). During polyamine catabolism, SSAT

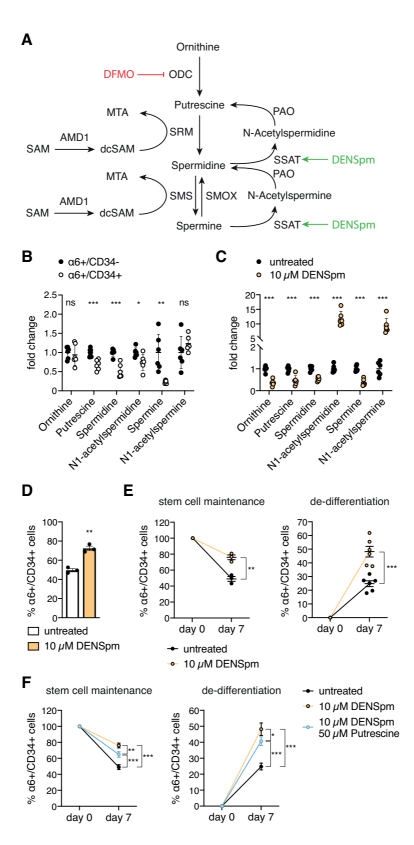


Figure 2: Reduction of translation by depletion of natural polyamines recapitulates the stem cell state and regulates cell fate decisions. (A) Schematic representation of the polyamine pathway. DENSpm is marked in green and DFMO is depicted in red. (B) Polyamine levels in sorted $\alpha6+/\text{CD}34$ -and $\alpha6+/\text{CD}34+$ cells after 3D-3C culture. Mean \pm SEM (n=6). (C) Polyamine levels in 3D-3C cultured cells with and without DENSpm treatment. Mean \pm SEM (n=6). (D) Ratio of $\alpha6+/\text{CD}34+$ cells after two weeks of 3D-3C culture with or without DENSpm treatment. Mean \pm SEM (n=3). (E) Ratio of $\alpha6+/\text{CD}34+$ cells at day 0 and day 7 post-sorting starting from 100 % $\alpha6+/\text{CD}34+$ cells (left) or 100 % $\alpha6+/\text{CD}34+$ cells (right) with or without DENSpm treatment. Mean \pm SEM (n \geq 3). (B-E) Statistical significance was

221

222 223

224

225

226

227

228

229

230

231

232

233

234

235

236

237

238

239

240

241

242

243

244

245

246

calculated by t test. *** p<0.001, ** p<0.01, * p<0.05, ns: not significant. (F) Ratio of α 6+/CD34+ cells at day 0 and day 7 post-sorting starting from 100 % α 6+/CD34+ cells (left) or 100 % α 6+/CD34- cells (right) with or without DENSpm or additional putrescine treatment. Mean ± SEM (n≥3). Two-way ANOVA Dunnett post-test. *** p<0.001, ** p<0.01, * p<0.05. acetylates spermidine and spermine using acetyl-CoA to form N1-acetylspermidine and N1-acetylspermine, respectively (Fig. 2A). Thus, DENSpm depletes natural polyamines, while the acetylated forms accumulate (Uimari et al., 2009), resulting in decreased mRNA translation (Landau et al., 2010). LC-MS analysis confirmed reduced natural polyamines and elevated acetylated forms upon DENSpm treatment in the 3D-3C organoids (Fig. 2C). Strikingly, supplementation with DENSpm resulted in a significant increase in α 6+/CD34+ stem cells after two weeks of 3D-3C culture (Fig. 2D). The increase in stemness was confirmed in the colony formation assay that revealed an elevated colony number upon DENSpm treatment in the 3D-3C organoids. suggesting increased proliferative potential (Fig. S2A-B). To investigate stem cell maintenance and de-differentiation separately, pure populations were sorted and cultured for one week with or without DENSpm treatment. Both mechanisms were affected by the depletion of natural polyamines, increasing α 6+/CD34+ stem cells (Fig. 2E). To rule out that the changes in stem cell number and proliferative potential were secondary to an effect on cell proliferation, we measured EdU incorporation in both α 6+/CD34+ and α 6+/CD34- cells. EdU incorporation was not affected by DENSpm treatment in either of the cell populations (Fig. S2C). Additionally, we analyzed apoptosis by annexin V staining. While annexin V staining was not changed in α 6+/CD34+ cells, DENSpm treatment slightly increased apoptosis in α 6+/CD34progenitors (Fig. S2D). However, this effect was not strong enough to account for the increase in α 6+/CD34+ HFSCs in treated organoid cultures. Furthermore, we demonstrated increased stem cell potency in the colony-forming assay (Fig. S2A-B). Thus, our data indicate that the promotion of the stem cell state by DENSpm was

through a direct effect on stem cell fate. Since the effect of DENSpm supplementation on the pure stem cell population was comparable to G418 treatment (Fig. 1H), we speculated that reduced translation might be the underlying mechanism of increased stemness. To test this hypothesis, we supplemented DENSpm treated cells with putrescine to elevate the natural polyamines. Indeed, this double treatment showed a partial rescue as stem cell maintenance and de-differentiation were significantly suppressed (Fig. 2F). Taken together, these data demonstrate that α 6+/CD34+ stem cells have lower polyamine levels, and that the reduction of natural polyamines influences HFSC fate through decreased translation.

N1-acetylspermidine is a novel determinant of HFSC fate acting independently

of reduced translation.

To further test if reduction of translation by depletion of natural polyamines caused the observed cell fate changes upon DENSpm treatment, we made use of difluoromethylornithine (DFMO), an irreversible ODC inhibitor (Metcalf et al., 1978) (Fig. 2A) that depletes all polyamines. In contrast to ODC inhibition by DFMO, DENSpm supplementation caused specific reduction of the natural polyamines putrescine, spermidine, and spermine, while the acetylated forms accumulated (Fig. 2C). Surprisingly, DFMO treatment did not affect cell fate in the 3D-3C organoids (Fig. 3A). Thus, we hypothesized that reduced mRNA translation through decreased polyamine availability was not the only determinant of the cell fate changes observed upon DENSpm treatment. Since additional putrescine supplementation only partially rescued the DENSpm effect on stem cell maintenance and de-differentiation, we speculated that putrescine itself might affect cell fate. Indeed, putrescine treatment in the 3D-3C organoids resulted in an increase in the number of α 6+/CD34+ stem cells (Fig. 3B). Increased stemness was confirmed by elevated colony-forming ability.

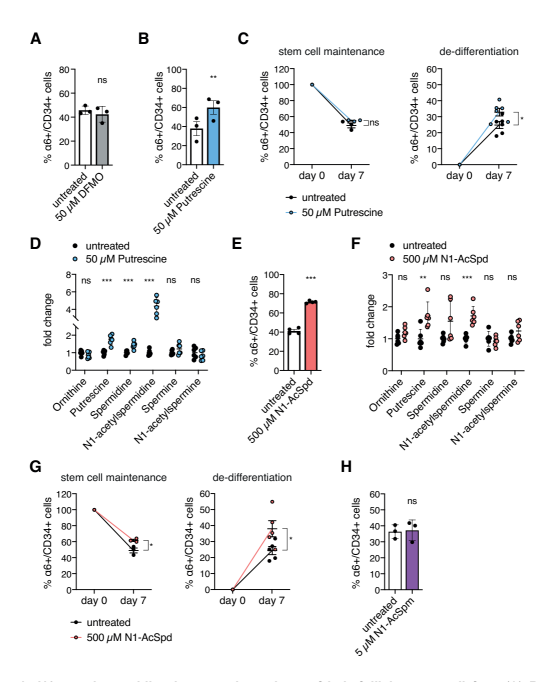


Figure 3: N1-acetylspermidine is a novel regulator of hair follicle stem cell fate. (A) Ratio of α 6+/CD34+ cells after two weeks of 3D-3C culture with or without DFMO treatment. Mean \pm SEM (n=3). (B) Ratio of α 6+/CD34+ cells after two weeks of 3D-3C culture with or without putrescine treatment. Mean \pm SEM (n=3). (C) Ratio of α 6+/CD34+ cells at day 0 and day 7 post-sorting starting from 100 % α 6+/CD34+ cells (left) or 100 % α 6+/CD34- cells (right) with or without putrescine treatment. Mean \pm SEM (n≥3). (D) Polyamine levels in 3D-3C cultured cells with and without putrescine treatment. Mean \pm SEM (n=6). (E) Ratio of α 6+/CD34+ cells after two weeks of 3D-3C culture with or without N1-AcSpd treatment. Mean \pm SEM (n=3). (F) Polyamine levels in 3D-3C cultured cells with and without N1-AcSpd treatment. Mean \pm SEM (n=6). (G) Ratio of α 6+/CD34+ cells at day 0 and day 7 post-sorting starting from 100 % α 6+/CD34+ cells (left) or 100 % α 6+/CD34- cells (right) with or without N1-AcSpd treatment. Mean \pm SEM (n≥3). (H) Ratio of α 6+/CD34+ cells after two weeks of 3D-3C culture with or without N1-AcSpm treatment. Mean \pm SEM (n=3). ns: not significant (t test).

showing enhanced proliferative potential (Fig. S3A-B). Surprisingly, putrescine supplementation specifically enhanced de-differentiation of α 6+/CD34- progenitors to

289

290

291

292

293

294

295

296

297

298

299

300

301

302

303

304

305

306

307

308

309

310

311

312

α6+/CD34+ HFSCs, while stem cell maintenance was not affected (Fig. 3C). However, the effect was not as pronounced as with DENSpm treatment (Fig. 2E). To further investigate the effect of putrescine supplementation, we measured intracellular polyamine levels in the 3D-3C organoids. Spermidine and N1-acetylspermidine (N1-AcSpd) levels were increased in response to putrescine treatment (Fig. 3D). In addition, an elevation of putrescine confirmed the effectiveness of our treatment. Since both DENSpm and putrescine supplementation led to elevated N1-AcSpd levels (Fig. 2C and Fig. 3D), we speculated that increased N1-AcSpd might cause the cell fate changes. An increase in any polyamine would not be expected to reduce translation and would then be mediated through a distinct mechanism. Intriguingly, N1-AcSpd treatment significantly increased the number of α 6+/CD34+ stem cells in the 3D-3C organoids (Fig. 3E). Stemness was confirmed by enhanced proliferative potential in the colony formation assay (Fig. S3C-D). As expected, analysis of puromycin incorporation confirmed that the effect on stemness occurred without a reduction of translation. Instead, translation rates were increased upon short-term N1-AcSpd treatment (72 h) (Fig. S3E-F), which did not affect the ratio of HFSCs in the 3D-3C organoids (Fig. S3G). Next, we measured intracellular polyamine levels upon N1-AcSpd treatment and found that apart from N1-AcSpd, also putrescine was increased (Fig. 3F). Thus, the effect of N1-AcSpd treatment on intracellular polyamines resembled the putrescine supplementation (Fig. 3D). However, in contrast to putrescine addition, N1-AcSpd treatment enhanced not only de-differentiation, but also stem cell maintenance when starting from pure populations (Fig. 3G). In stark contrast, supplementation with N1-acetylspermine (N1-AcSpm) did not influence the ratio of HFSCs (Fig. 3H). Thus, elevation of polyamines in general was not sufficient; rather, putrescine and N1-AcSpd treatment had specific effects on cell fate. Overall, these

data implicate N1-AcSpd as a novel cell fate regulator that acts independently of reduced translation.

N1-acetylspermidine treatment affects hair follicle stem cell fate by increasing

proliferation.

313

314

315

316

317

318

319

320

321

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

338

To investigate the molecular mechanism by which N1-AcSpd supplementation promotes stemness, we performed RNA-sequencing of 3D-3C cultured cells upon short-term N1-AcSpd treatment (72 h). HFSCs and progenitor cells were purified and analyzed separately (Fig. S4A). We compared α 6+/CD34- cells with α 6+/CD34+ cells to identify differentially expressed genes for control (CTR) and N1-AcSpd treatment (Fig. S4B). The two groups of differentially expressed genes were subjected to further analysis. Around 2.200 genes with differential expression between cell types were not affected by N1-AcSpd treatment (Fig. S4C). 851 genes were differentially expressed between HFSCs and progenitors only in the presence of N1-AcSpd, while 540 genes were differentially expressed between cell types exclusively in untreated control cells (Fig. S4C). GO term analysis was performed with the three different groups using Metascape (Zhou et al., 2019). Strikingly analysis of the genes differentially expressed between HFSCs and progenitors only during N1-AcSpd treatment revealed that most of the ten highest ranked GO terms were linked to cell cycle progression (Fig. 4A). The most significant GO term was cell division, of which 54 genes were present among the 851 differentially expressed genes in N1-AcSpd treatment (Fig. S4D). Interestingly, expression of the majority of these 54 genes was higher in α 6+/CD34+ stem cells (Fig. 4B). Importantly, N1-AcSpd treatment affected the amplitude of the fold change; the expression of these genes was higher also in the untreated α 6+/CD34+ stem cells (Fig. 4B). Consistently, EdU incorporation was higher in untreated α 6+/CD34+ HFSCs compared to progenitor cells (Fig. S4E). Together, these data suggest that N1-AcSpd

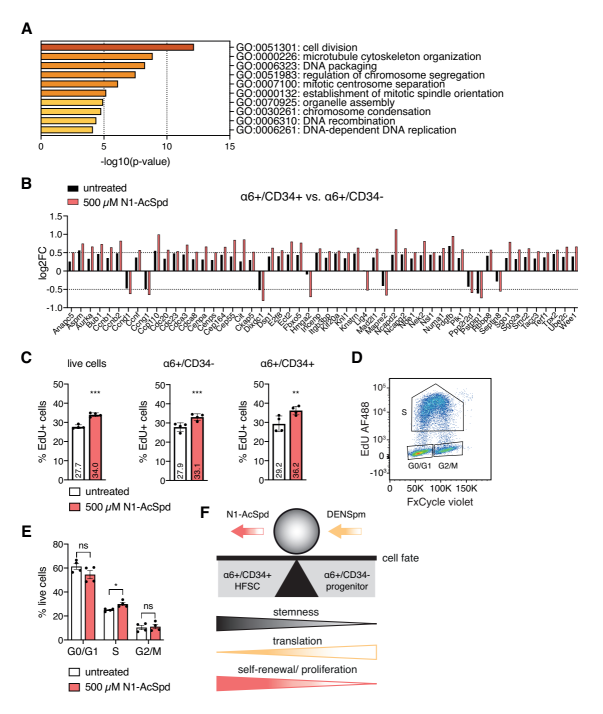


Figure 4: N1-acetylspermidine treatment affects cell fate by increasing cell cyle progression. (A) GO term analysis of differentially regulated genes between cell types (p-value < 0.05, log2FC > \pm 0.5) only upon N1-AcSpd treatment (depicted in red in Fig. S4A, biological process, metascape.org). (B) log2FC of the genes associated with the GO term "cell division" in untreated and N1-AcSpd treated cells. The ratio of the log2FC in α 6+/CD34+ vs. α 6+/CD34- cells is shown. (C) Ratio of EdU+ cells with or without N1-AcSpd treatment (72 h). EdU was incorporated for 2 h. Mean \pm SEM (n=4). *** p<0.01; ** p<0.01 (t test). (D) Dot plot of FxCycle violet intensity and EdU AF488 intensity, separating the phases of the cell cycle (G0/G1, S, G2/M). (E) Ratio of live cells with or without N1-AcSpd treatment (72 h) according to their cell cycle phase distribution. Mean \pm SEM (n=4). * p<0.05; ns: not significant (t test). (F) Model of polyamine-controlled effects on cell fate. Manipulation of translation or proliferation by DENSpm or N1-AcSpd, respectively, results in the enrichment of HFSCs.

selectively promoted self-renewal of HFSCs. GO term analysis of the around 2.200 genes that differed between HFSCs and progenitors without being affected by N1-

354

355

356

357

358

359

360

361

362

363

364

365

366

367

368

369

370

371

372

373

374

375

376

377

378

AcSpd revealed skin development as the most significant GO term (Fig. S4F). This suggests that N1-AcSpd treatment did not affect general cell identity in the 3D-3C organoids. Consistently, a comparison of the gene expression changes between cell types in a heat map showed a clear separation of α 6+/CD34+ HFSCs and α 6+/CD34progenitors (Fig. S4G). Although N1-AcSpd treated cells clustered separately of control cells, their gene expression was very similar to the respective cell type in the control (Fig. S4G). Analysis of the 540 genes differentially expressed between cell types only in untreated cells showed low significance for all GO terms. The most significant GO terms were intraciliary transport and cilium organization (Fig. S4H). Since the analysis of global gene expression changes clearly indicated that N1-AcSpd treatment would affect proliferation, we measured EdU incorporation. Importantly, short-term N1-AcSpd supplementation (72 h) increased the number of EdU+ cells (Fig. 4C). The effect on α 6+/CD34+ HFSCs was higher compared to α 6+/CD34progenitor cells (Fig. 4C). Still, consistent with increased de-differentiation and elevated stem cell maintenance, proliferation of both populations was affected by the treatment. Furthermore, we analyzed the cell cycle phase distribution of the cells using FxCycle violet staining. Plotting of FxCycle violet intensity on the x-axis and EdU AF488 intensity on the y-axis enabled clear separation of the different cell cycle phases (Fig. 4D). Consistent with increased EdU incorporation, we observed faster cell cycle progression with fewer cells in G0/G1 phase and more cells in S phase upon shortterm N1-AcSpd treatment (72 h) (Fig. 4E). The number of cells in G2/M phase was not changed. Taken together, these data suggest that the acetylated polyamine N1-AcSpd fulfills a functional role in cell cycle progression and that it affects cell fate by increasing proliferation. Overall, in this study we decipher the different and independent polyamine-controlled effects on HFSC fate. We demonstrate that decreasing translation rates by the depletion of natural polyamines through DENSpm, as well as

- 379 enhancing proliferation by N1-AcSpd shifts the balance towards the stem cell state in
- 380 the 3D-3C organoids (Fig. 4F).

Discussion

381

382

383

384

385

386

387

388

389

390

391

392

393

394

395

396

397

398

399

400

401

402

403

404

405

406

In this study we delineate the different routes of polyamine-mediated regulation of HFSC maintenance and function. First, we show that reduced translation is not only a marker of the stem cell state. Instead, a forced decrease of translation was sufficient to enhance stemness. Next, we noted low polyamine levels in α 6+/CD34+ stem cells and depletion of the natural polyamines with DENSpm resulted in enhanced HFSC maintenance. Strikingly, full depletion of polyamines by DFMO did not recapitulate our findings with DENSpm, suggesting that reduced translation was not the only relevant consequence of polyamine changes in cell fate. Indeed, putrescine supplementation had a beneficial effect on progenitor cell de-differentiation. This was mediated by N1-AcSpd that we identify as novel regulator of HFSC fate, acting independently of reduced translation by increasing proliferation. Previously, Blanco et al. (2016) showed that translation becomes elevated during the hair cycle growth phase when cells become activated and differentiate. In contrast, quiescent HFSCs in the resting phase show low translation. This agrees with other studies showing that translation is upregulated during differentiation (Sampath et al., 2008, Signer et al., 2014, Zismanov et al., 2016). Importantly, we can reproduce these findings in vivo and in the 3D-3C HFSC organoid culture; α6+/CD34+ stem cells in the organoids recapitulate key features of HFSCs in vivo. Strikingly, forced inhibition of translation by loss of NOP2/Sun RNA methyltransferase family member 2 (NSUN2) in the mouse blocks the differentiation of HFSCs to progenitor cells (Blanco et al., 2011), clearly showing that upregulation of translation is necessary for differentiation in the HF. Using G418 treatment or the depletion of natural polyamines to reduce translation in the organoids, we confirm promotion of the stem cell state. We further show that the number of stem cells was elevated due to enhanced self-renewal rather than attenuated differentiation using EdU incorporation during DENSpm treatment.

408

409

410

411

412

413

414

415

416

417

418

419

420

421

422

423

424

425

426

427

428

429

430

431

Additionally, we see an effect on progenitor cell de-differentiation, further supporting the idea that a forced decrease of translation is sufficient to promote stemness in the 3D-3C organoids. Previously, the acetylated polyamines were mostly described as the major group of polyamines exported from the cell (Seiler & Dezeure, 1990). Extending this notion, we find that the intracellular accumulation of N1-AcSpd has an effect on stemness. This occurs without reducing translation, which is consistent with previous observations that acetylated polyamines have no effect on protein synthesis in vitro (Kakegawa et al., 1991). Instead, RNA-sequencing of purified cell populations from the organoids revealed that N1-AcSpd treatment affected expression of cell cycle-associated genes. Interestingly, these same genes were found expressed at higher levels in α 6+/CD34+ stem cells, which also display higher EdU incorporation compared to progenitor cells in the organoid cultures. Importantly, increased proliferation and stemness have been linked before. A study in ESCs suggests that elevated CDK activity, and thus increased cell cycle progression, contributes to stem cell maintenance (Liu et al., 2017). Strikingly, several studies have described that cells in the G1 phase are more susceptible to cell fate changes and that differentiation is associated with G1 phase lengthening (Calder et al., 2013, Clegg et al., 1987, Sela et al., 2012). Consistently, reprogramming efficiency seems to be linked to successful acceleration of the cell cycle (Guo et al., 2014, Ruiz et al., 2011). Of note, forced overexpression of AMD1 or ODC in mouse fibroblasts, resulting in the accumulation of polyamines, improves reprogramming efficiency (Zhang et al., 2012, Zhao et al., 2012). Our results suggest that this effect is caused by increased proliferation. Collectively, these data indicate that both stem cell maintenance and de-differentiation might be improved by elevated cell cycle progression upon N1-AcSpd treatment in our 3D-3C organoid culture.

433

434

435

436

437

438

439

440

441

442

443

444

445

446

447

448

449

450

451

452

453

454

455

456

457

Polyamines have been implicated in cell cycle progression before. Under normal growth conditions, polyamine levels are dynamically regulated during the cell cycle (Sunkara et al., 1981), which is due to cyclical changes in activity of ODC and AMD1 (Fredlund et al., 1995). Consistently, several studies have shown that polyamine depletion results in cell cycle arrest (Odenlund et al., 2009, Ray et al., 1999, Yamashita et al., 2013). Intriquingly, ODC activity was shown to be increased by N1-AcSpd, while N1-AcSpm, which did not influence cell fate in the organoid culture, does not affect ODC activity (Canellakis et al., 1989). Thus, activation of ODC might be an important step in cell fate regulation via increased cell cycle progression. Strikingly, expression of ODC is also dynamically regulated during the hair cycle. Nancarrow et al. (1999) showed that ODC is abundantly expressed in proliferating bulb cells of anagen follicles, while entry of the follicle into catagen is accompanied by down-regulation of ODC. These data implicate that polyamine levels, which are controlled by ODC, might track proliferation rates in the hair follicle. Additionally, the authors revealed expression of ODC in ORS cells in vicinity of the follicle bulge region (Nancarrow et al., 1999). After the anagen growth phase, some ORS cells return to the bulge region, where they resume guiescence and CD34 expression. These cells become the primary stem cells for the initiation of the next hair cycle (Hsu et al., 2011). This process is comparable to the de-differentiation of α 6+/CD34- progenitor cells back to the stem cell state in the 3D-3C organoids, which was enhanced by putrescine and N1-AcSpd supplementation. Overall, these data suggest that ODC activity, cell cycle progression, and cell fate are tightly linked. In the HF, quiescent stem cells ensure reduced translation rates by low polyamine levels, which increase upon activation due to elevated ODC activity, resulting in accelerated cell cycle progression. Consistently, skin-specific ODC overexpression results in hair loss beginning 2 to 3 weeks after birth, demonstrating the need for low polyamine levels in HFSC quiescence (Soler et al., 1996). Our data

459

460

461

462

463

464

465

466

467

468

469

470

471

472

473

474

suggest that enhanced cell cycle progression promotes stem cell self-renewal upon activation, while differentiation is decreased. At the same time, cell cycle progression might be required for successful de-differentiation of ORS cells back to HFSCs, a notion supported by elevated ODC expression in ORS cells. Overall, this study dissects independent routes of polyamine-controlled regulation of cell fate: While depletion of natural polyamines endogenously reduced translation. addition of N1-AcSpd accelerated cell cycle progression. Previously, acetylated polyamines were described primarily as the major polyamine species exported from the cell. Here, we demonstrate that N1-AcSpd directly influenced cell fate decisions via increased proliferation. Thus, our study explains why elevated polyamine levels and low translation rates are not mutually exclusive in stem cell maintenance. Instead, they regulate different aspects of cell fate. While low translation rates favor guiescence in vivo, enhanced cell cycle progression ensures stem cell self-renewal upon activation. ODC activity might function as the molecular switch to regulate polyamine availability and thus cell fate in the HF. Over the long-term, N1-AcSpd treatment could be a viable intervention to tackle age-associated diseases caused by a decline in tissue homeostasis due to stem cell exhaustion.

475 Methods 476 477 Mouse husbandry 478 Animals were housed on a 12:12 h light:dark cycle with ad libitum access to food under 479 pathogen-free conditions in individually ventilated cages. All animals were kept in 480 C57BL/6J background. Animal care and experimental procedures were in accordance 481 with the institutional and governmental guidelines. 482 Isolation of epidermal cells 483 484 Isolation of epidermal cells from mice in telogen stage was performed as described 485 previously (Chacon-Martinez et al., 2017). In brief, mouse skin was incubated on 0.8 % 486 trypsin (ThermoFisher Scientific) for 50 min at 37 °C. The skin was transferred to 8 ml 487 KGM medium and the epidermis was separated from the dermis. After centrifugation, 488 the keratinocytes were resuspended in ice cold KGM medium and embedded in growth 489 factor-reduced matrigel (Corning Inc.). 490 491 Culture of hair follicle stem cell organoids 492 Hair follicle stem cell organoids were cultured in KGM medium: MEM medium 493 (Spinner's modification, Sigma-Aldrich), supplemented with 8 % fetal bovine serum 494 (chelated, ThermoFisher Scientific), penicillin/streptavidin (ThermoFisher Scientific), L-495 glutamine (ThermoFisher Scientific), insulin (5 µg/ml, Sigma-Aldrich), hydrocortisone 496 (0.36 μg/ml, Calbiochem), EGF (10 ng/ml, Sigma-Aldrich), transferrin (10 μg/ml, 497 Sigma-Aldrich), phosphoethanolamine (10 µM, Sigma-Aldrich), ethanolamine (10 µM, 498 Sigma-Aldrich), CaCl₂ (14.5 μM, Sigma-Aldrich). 5 μM Y27632, 20 ng/ml mouse 499 recombinant VEGF, 20 ng/ml human recombinant FGF2 (all Miltenyi Biotech) were 500 added to KGM medium. The cells were grown at 37 °C in 5 % CO₂.

502

503

504

505

506

507

508

509

510

511

512

513

514

515

516

517

518

519

520

521

522

523

524

525

526

Cell maintenance NIH3T3 fibroblasts were grown in DMEM containing 4.5 g/L glucose supplemented with 10 % fetal bovine serum and penicillin/streptavidin (all ThermoFisher Scientific) at 37 °C in 5 % CO₂ on non-coated tissue culture plates. **Colony formation assay** NIH3T3 fibroblasts were seeded on collagen G-coated (30 µg/ml in PBS, Biochrom AG) tissue culture plates. The cells were grown at 37 °C in 5 % CO₂ for two days before proliferation was inhibited with mitomycin C (Sigma-Aldrich). Hair follicle stem cells were grown in 3D-3C organoids for two weeks and analyzed by flow cytometry. 4000 cells were seeded per 6-well on the fibroblast layer in MEM/HAM's F12 (FAD) medium with low Ca²⁺ (50 μM, Biochrom AG), supplemented with 10% fetal bovine serum (chelated, ThermoFisher Scientific), penicillin/streptavidin (ThermoFisher Scientific), L-glutamine (ThermoFisher Scientific), ascorbic acid (50 μg/ml, Sigma-Aldrich), adenine (0.18 mM, Sigma-Aldrich), insulin (5 μg/ml, Sigma-Aldrich), hydrocortisone (0.5 µg/ml, Sigma-Aldrich), EGF (10 ng/ml, Sigma-Aldrich), and cholera enterotoxin (10 ng/ml, Sigma-Aldrich). The cells were grown for 2-3 weeks at 32 °C in 5 % CO₂ until colonies were formed. Remaining fibroblasts were removed using 0.25 % trypsin-ETDA (ThermoFisher Scientific) for 2 min at 37 °C. Trypsin was stopped using supplemented DMEM. After washing the plates with PBS, keratinocytes were fixed with 4 % PFA (Sigma-Aldrich) for 15 min at RT. The cells were washed with PBS twice and the colonies were stained using 1 % crystal violet (Sigma-Aldrich in PBS) for 1 h at RT on an orbital shaker. The wells were washed with tap water until no stain was released and air dried. The plates were scanned and the number of colonies was counted manually.

Flow cytometry

Matrigel droplets were degraded in 0.5 % trypsin, 0.5 mM EDTA in PBS for 8 min at 37 °C. Trypsin was neutralized using cold KGM. After centrifugation for 5 min at 2000 rpm cells were washed with FACS buffer (2 % FBS, 2 mM EDTA in PBS). Surface marker staining was performed for 30 min on ice with the following antibodies: CD34-eFluor 660 (eBioscience, clone RAM34, 1:100) and ITGA6-PE/Cy7 (Biolegend, clone GoH3, 1:1000) for sorting or CD34-eFluor 660 (eBioscience, clone RAM34, 1:100) and ITGA6-pacific blue (Biolegend, clone GoH3, 1:200) for analysis. Freshly isolated keratinocytes were stained additionally with Sca1-Pacific blue (Biolegend, clone D7, 1:400). 7AAD or PI was used to exclude dead cells. Cells were analyzed on FACSCantolI (BD Biosciences) or sorted on FACSAria IIIu Svea and FACSAria Fusion sorters (BD Biosciences). Sorted cells were collected in 15 ml tubes containing KGM at 4 °C.

EdU incorporation and detection

Cell proliferation was assessed using the Click-iT[™] Plus EdU Alexa Fluor[™] 488 Flow Cytometry Assay Kit (ThermoFisher) following the manufacturer's instructions. Cells were grown for 9 days in 3D-3C culture before 10 µM EdU was added to the medium and incubated for 2 h or 24 h. Matrigel droplets were degraded, the cells were washed with PBS and subsequently stained with fixable LIVE/DEAD-violet (ThermoFisher, 1:500) for 20 min at RT. The cells were washed with FACS buffer and surface marker staining was performed for 30 min on ice using CD34-eFluor 660 (eBioscience, clone RAM34, 1:100) and ITGA6-PE/Cy7 (Biolegend, clone GoH3, 1:1000). The cells were washed with 1 % BSA in PBS, fixed with 4 % PFA for 10 min at RT and permeabilized using Click-iT saponin-based permeabilization buffer for 15 min at RT. The EdU

reaction cocktail was prepared following the manufacturer's instructions and incubated for 30 min at RT in the dark after washing the cells. The cells were analyzed on FACSCantoll.

Cell cycle analysis

Cell proliferation was assessed using the Click-iT™ Plus EdU Alexa Fluor™ 488 Flow Cytometry Assay Kit (ThermoFisher) as described above. Cells were grown for two weeks in 3D-3C culture before 10 µM EdU was added to the medium and incubated for 2 h. Matrigel droplets were degraded, the cells were washed with PBS and stained with fixable Zombie NIR dye (Biolegend, 1:500) for 20 min at RT. The cells were washed, fixed, and permeabilized and the EdU reaction cocktail was incubated as described above. Subsequently, the cells were permeabilized using 0.1 % triton X-100 in PBS for 15 min at RT and stained with FxCycle violet (ThermoFisher, 1:500) for 30 min at RT. The cells were analyzed on FACSCantoll without further washing.

Annexin V staining

Matrigel droplets were degraded in 0.5 % trypsin, 0.5 mM EDTA in PBS for 8 min at 37 °C. Trypsin was neutralized using cold KGM. After centrifugation for 5 min at 2000 rpm cells were washed with FACS buffer. Surface marker staining was performed for 30 min on ice using CD34-eFluor 660 (eBioscience, clone RAM34, 1:100) and ITGA6-pacific blue (Biolegend, clone GoH3, 1:200). Cells were washed with PBS and stained with annexin V-AF488 (ThermoFisher Scientific, 1:20 in 100 μl annexin-binding buffer per sample) for 15 min at RT. 100 μl annexin-binding buffer containing PI were added per sample. The cells were analyzed on FACSCantolI.

Puromycin incorporation

Cells were incubated with 10 µg/ml puromycin in the medium for exactly 10 min at 37 °C. The cells were washed with PBS and collected in RIPA buffer (50 mM TrisHCl, 120 mM NaCl, 0.1 % SDS, 1 % NP40, 0.5 % deoxycholate) with proteinase and phosphatase inhibitors (Roche Diagnostics GmbH).

Western Blot analysis

Protein concentration of cell lysates was determined using the PierceTM BCA protein assay kit according to manufacturer's instructions (ThermoFisher Scientific). Samples were subsequently subjected to SDS-PAGE and blotted on a nitrocellulose membrane. The following antibodies were used in 5 % low-fat milk in TBS-Tween buffer: Puromycin (ms, Millipore, 12D10, 1:10.000), β-actin (ms, Sigma, AC-74, 1:25.000). After incubation with HRP-conjugated secondary antibody (Invitrogen, 1:5000), the blot was developed using ECL solution (Merck Millipore). Films were used for detection (Amersham Biosciences).

RNA isolation and qRT-PCR

Matrigel droplets were degraded in 0.5 % trypsin, 0.5 mM EDTA in PBS for 8 min at 37 °C. Trypsin was neutralized using cold KGM. After centrifugation for 5 min at 2000 rpm cells were washed with PBS. The pellet was resuspended in QIAzol (Qiagen) and frozen in liquid nitrogen. Sorted and freshly isolated keratinocytes were collected in QIAzol and frozen in liquid nitrogen. Samples were subjected to three freeze/thaw cycles (liquid nitrogen/ 37 °C water bath). Half of the total volume of QIAzol was added per sample. After incubation for 5 min at RT, 200 µl chloroform were added per 1 ml QIAzol. The samples were vortexed, incubated for 2 min at RT, and centrifuged at 10.000 rpm and 4 °C for 15 min. The aqueous phase was mixed with an equal volume of 70 % ethanol and transferred to a Rneasy Mini spin column (Qiagen). Total RNA

was isolated with the Rneasy Mini Kit (Qiagen) according to manufacturer's instructions. cDNA was generated using the iScript cDNA Synthesis Kit (Bio-Rad Laboratories Inc.). qRT-PCR was performed using Power SYBR Green master mix (Applied Biosystems) on a ViiA 7 Real-Time PCR System (Applied Biosystems). Primer sequences are listed below.

Table 1: Primer sequences for gRT-PCR.

Primer/gene name	Sequence (5'→ 3')		
huTBP_fwd	GGGTTTTCCAGCTAAGTTCTTG		
huTBP_rev	CTGTAGATTAAACCAGGAAATAACTCTG		
huSSAT_fwd	GGTTGCAGAAGTGCCGAAAG		
huSSAT_rev	GTAACTTGCCAATCCACGGG		

Preparation of 3' RNA-sequencing libraries

Sorted cells were collected in QIAzol and frozen in liquid nitrogen. Total RNA was isolated as described above. 50 ng RNA per sample were used for cDNA synthesis with Maxima H Minus reverse transcriptase (ThermoFisher Scientific). During reverse transcription, unique barcodes including unique molecular identifiers (UMI) were attached to each sample. After cDNA synthesis, all samples were pooled and processed in one single tube. DNA was purified using AmpureXP beads (Beckman Coulter) and the eluted cDNA was subjected to Exonuclease I treatment (New England Biolabs). cDNA was PCR-amplified for 12 cycles and subsequently purified. After purification, cDNA was tagmented in 5 technical replicates of 1 ng cDNA each using the Nextera XT Kit (Illumina), according to manufacturer's instructions. The final library was purified and concentration and size were validated by Qubit and High Sensitivity TapeStation D1000 analyses. A detailed protocol for library preparation is available on request. Sequencing was carried out on an Illumina NovaSeq system.

Bioinformatic analysis of 3' RNA-sequencing data

The raw reads were demultiplexed and the UMI-tag was added to each read name using FLEXBAR (Dodt et al., 2012). The second end of the read-pairs were mapped using Kallisto creating pseudo-alignments to the mouse reference genome mm10 (Bray et al., 2016). UMI-tags were assigned using BEDTools intersect and bash reformatting (Quinlan & Hall, 2010). An R script was used to count the unique UMI-tags per gene. Differential gene expression was calculated using the R package Deseq2 (Love et al., 2014). Only genes with an average of more or equal to 5 reads over all samples were used for the analysis. Only genes with an adjusted p-value ≤ 0.05 ($\alpha6+/CD34-$ vs. $\alpha6+/CD34+$; untreated or treated) are shown in the heatmap. Venn diagrams were generated using the R package VennDiagram. Functional enrichment analysis was performed using the Metascape tool [http://metascape.org] (Zhou et al., 2019). Only genes with a p-value < 0.05 and a log2FC $> \pm 0.5$ ($\alpha6+/CD34-$ vs. $\alpha6+/CD34+$; untreated or treated) were defined as differentially expressed and included in the functional enrichment analysis.

Polyamine extraction from cells

Matrigel droplets were degraded in 0.5 % trypsin, 0.5 mM EDTA in PBS for 8 min at 37 °C. Trypsin was neutralized using cold KGM. After centrifugation for 5 min at 2000 rpm cells were washed with PBS. Sorted cells were centrifuged and washed with PBS. The pellet was flash frozen in liquid nitrogen. The cells were lysed in ddH2O by freeze/thaw cycles and the protein concentration was determined using the Pierce[™] BCA protein assay kit (ThermoFisher Scientific). Polyamines were extracted by Bligh and Dyer extraction (Noga et al., 2012, Pieragostino et al., 2015). In brief, a methanol:chloroform mixture 2:1 (v/v) was added to a volume of cell lysate, which

654

655

656

657

658

659

660

661

662

663

664

665

666

667

668

669

670

671

672

673

674

675

676

677

678

corresponded to 100 µg of protein and incubated 1 h at 4 °C. Samples were centrifuged for 5 min at 3000 rcf at 4 °C and the liquid phase was transferred and dried in a SpeedVac Vacuum Concentrator (Genevac). Samples were reconstituted with 15 µL aqueous acetonitrile 2:3 (v/v) and 10 µl were injected into the LC-MS system. Targeted analysis of polyamines by LC-MS The identification of polyamines was performed on a triple stage quadrupole (TSQ Altis, ThermoFisher Scientific GmbH) coupled with a binary pump system (Vanguish, ThermoFisher Scientific GmbH). Polyamine species were separated using a reverse column (Xselect column, 2.1x100mm, 2,5 µm) using solvent A (water with 0.1 % formic acid) and B (acetonitrile with 0.1 % formic acid) as previously reported (Hakkinen et al., 2008, Hakkinen et al., 2013, Sanchez-Lopez et al., 2009). The gradient started from 0.1 % eluent B and ramped to 0.3 % eluent B in 0.5 min. It ramped further to 0.5 % eluent B in 0.5 min and to 1 % B in the next 0.5 min. The gradient increased to 2 % eluent B in the next minute, then it ramped to 5 % eluent B in 1 min. In the following 3 min it went to 95 % eluent B and stayed constant for 3 min. Afterwards the gradient decreased to 0.1 % eluent B in 2 min and stayed constant for 3 min adding up to a total time of 15 min. The column was heated to 30 °C using a flow rate of 100 µl/min. The LC system was flushed in between runs with isopropanol:water 75:25 (v/v) with 0.1 % FA. Polyamines were detected using heated electrospray ionization (HESI) with the following parameters: 10 (a.u.) sheath gas, 5 (a.u.) auxiliary, 200 °C transfer ion capillary and 3 kV for spray voltage. Relative quantification was performed using selected ion monitoring chromatogram mode (SIM-Q1) in positive ion mode using a scan rate of 1000 Da/sec, Q1 resolution was set to 0.7 m/z and calibrated RF lenses were used. The following ions were

monitored: arginine \rightarrow 175.2, ornithine \rightarrow 133.17, putrescine \rightarrow 89.17, spermidine \rightarrow 146.23, spermine \rightarrow 203.32, N1-acetylspermidine \rightarrow 188.25, N1-acetylspermine \rightarrow 245.34, N1,N11-diethylnorspermine \rightarrow 245.39. The relative response for each polyamine was calculated using Spermidine-(butyl-d8) as internal standard.

Table 2: Detailed description of the detected metabolites.

Name	Elemental composition	retention time (min)	Mass (amu)	[M+H] ⁺ (m/z)
Arginine	$C_6H_{14}N_4O_2$	2.38	174.111	175.118
Ornithine	$C_5H_{12}N_2O_2$	2.73	132.089	133.097
Putrescine	$C_4H_{12}N_2$	2.41	88.099	89.107
Spermidine	$C_7H_{19}N_3$	2.22	145.157	146.165
Spermine	$C_{10}H_{26}N_4$	2.46	202.215	203.223
N1-acetylspermidine	$C_9H_{21}N_3O$	3.14	187.167	188.175
N1-acetylspermine	$C_{12}H_{28}N_4O$	3.05	244.225	245.233
N1,N11-diethylnorspermine	$C_{13}H_{32}N_4$	2.30	244.262	245.269

Data Availability

The raw RNA-sequencing data in this publication are available upon request from the authors.

Acknowledgements

688

689

690

691

692

693

694

695

696

697

698

699

700

701

702

703

704

We thank all M.S.D. laboratory members for lively and helpful discussions. We thank F.A.M.C. Mayr for critical reading of the manuscript. We thank K. Folz-Donahue and L. Schumacher from the FACS and imaging core facility, F. Metge, A. Iqbal and J. Boucas from the Bioinformatics core facility and the Comparative Biology Facility at the Max Planck Institute for Biology of Ageing. We thank Patrick Wollek for expert mouse work. This study was supported by the Cologne Graduate School for Ageing Research (to K.A.), by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) – Projektnummer 73111208 – SFB 829 (to S.A.W. and M.S.D.), by the Jane and Aatos Erkko Foundation (to S.A.W.), by the ERC-StG 640254 and ERC-PoC 768524 (to M.S.D.), and by the Max Planck Society (to Ad.An., P.T., S.A.W., and M.S.D.).

Conflict of Interest

The authors declare no potential competing interests.

Authorship

- 705 K.A., C.S.K., An.An., A.P., C.A.C.-M., P.T., S.A.W. and M.S.D designed the research.
- Ad.An. gave feedback throughout the project. K.A. (original draft), C.S.K., A.P., P.T.,
- 707 S.A.W. and M.S.D (review & editing) wrote the manuscript. An.An. and C.L. performed
- 708 LC-MS analysis. A.P. prepared the library for 3' RNA-sequencing. K.A. performed all
- other experiments. Funding acquisition: Ad.An., P.T., S.A.W. and M.S.D.

710 References

- 711 Bar-Nun S, Shneyour Y, Beckmann JS (1983) G-418, an elongation inhibitor of 80 S
- 712 ribosomes. Biochim Biophys Acta 741: 123-7
- 713 Blanco S, Bandiera R, Popis M, Hussain S, Lombard P, Aleksic J, Sajini A, Tanna H,
- 714 Cortes-Garrido R, Gkatza N, Dietmann S, Frye M (2016) Stem cell function and stress
- response are controlled by protein synthesis. Nature 534: 335-40
- 716 Blanco S, Kurowski A, Nichols J, Watt FM, Benitah SA, Frye M (2011) The RNA-
- 717 methyltransferase Misu (NSun2) poises epidermal stem cells to differentiate. PLoS
- 718 Genet 7: e1002403
- 719 Blanpain C, Fuchs E (2009) Epidermal homeostasis: a balancing act of stem cells in
- 720 the skin. Nat Rev Mol Cell Biol 10: 207-17
- 721 Bray NL, Pimentel H, Melsted P, Pachter L (2016) Erratum: Near-optimal probabilistic
- 722 RNA-seq quantification. Nat Biotechnol 34: 888
- 723 Calder A, Roth-Albin I, Bhatia S, Pilquil C, Lee JH, Bhatia M, Levadoux-Martin M,
- 724 McNicol J, Russell J, Collins T, Draper JS (2013) Lengthened G1 phase indicates
- 725 differentiation status in human embryonic stem cells. Stem Cells Dev 22: 279-95
- 726 Canellakis ZN, Marsh LL, Bondy PK (1989) Polyamines and their derivatives as
- 727 modulators in growth and differentiation. Yale J Biol Med 62: 481-91
- 728 Chacon-Martinez CA, Klose M, Niemann C, Glauche I, Wickstrom SA (2017) Hair
- 729 follicle stem cell cultures reveal self-organizing plasticity of stem cells and their
- 730 progeny. EMBO J 36: 151-164
- 731 Clegg CH, Linkhart TA, Olwin BB, Hauschka SD (1987) Growth factor control of
- 732 skeletal muscle differentiation: commitment to terminal differentiation occurs in G1
- 733 phase and is repressed by fibroblast growth factor. J Cell Biol 105: 949-56
- Cohen SS, Lichtenstein J (1960) Polyamines and ribosome structure. J Biol Chem 235:
- 735 2112-6

- 736 Coleman CS, Huang H, Pegg AE (1995) Role of the carboxyl terminal MATEE
- 737 sequence of spermidine/spermine N1-acetyltransferase in the activity and stabilization
- by the polyamine analog N1,N12-bis(ethyl)spermine. Biochemistry 34: 13423-30
- 739 Dever TE, Ivanov IP (2018) Roles of polyamines in translation. J Biol Chem 293:
- 740 18719-18729
- 741 Dodt M, Roehr JT, Ahmed R, Dieterich C (2012) FLEXBAR-Flexible Barcode and
- 742 Adapter Processing for Next-Generation Sequencing Platforms. Biology (Basel) 1:
- 743 895-905
- 744 Fogel-Petrovic M, Vujcic S, Brown PJ, Haddox MK, Porter CW (1996) Effects of
- 745 polyamines, polyamine analogs, and inhibitors of protein synthesis on spermidine-
- spermine N1-acetyltransferase gene expression. Biochemistry 35: 14436-44
- 747 Fredlund JO, Johansson MC, Dahlberg E, Oredsson SM (1995) Ornithine
- 748 decarboxylase and S-adenosylmethionine decarboxylase expression during the cell
- 749 cycle of Chinese hamster ovary cells. Exp Cell Res 216: 86-92
- 750 Fuchs E, Merrill BJ, Jamora C, DasGupta R (2001) At the roots of a never-ending
- 751 cycle. Dev Cell 1: 13-25
- Guo S, Zi X, Schulz VP, Cheng J, Zhong M, Koochaki SH, Megyola CM, Pan X,
- 753 Heydari K, Weissman SM, Gallagher PG, Krause DS, Fan R, Lu J (2014)
- Nonstochastic reprogramming from a privileged somatic cell state. Cell 156: 649-62
- 755 Gutierrez E, Shin BS, Woolstenhulme CJ, Kim JR, Saini P, Buskirk AR, Dever TE
- 756 (2013) eIF5A promotes translation of polyproline motifs. Mol Cell 51: 35-45
- 757 Hakkinen MR, Keinanen TA, Vepsalainen J, Khomutov AR, Alhonen L, Janne J,
- 758 Auriola S (2008) Quantitative determination of underivatized polyamines by using
- 759 isotope dilution RP-LC-ESI-MS/MS. Journal of pharmaceutical and biomedical
- 760 analysis 48: 414-21
- Hakkinen MR, Roine A, Auriola S, Tuokko A, Veskimae E, Keinanen TA, Lehtimaki T,
- Oksala N, Vepsalainen J (2013) Analysis of free, mono- and diacetylated polyamines
- 763 from human urine by LC-MS/MS. Journal of chromatography B, Analytical
- technologies in the biomedical and life sciences 941: 81-9

- Hershey JWB, Sonenberg N, Mathews MB (2019) Principles of Translational Control.
- 766 Cold Spring Harb Perspect Biol 11
- 767 Hsu YC, Li L, Fuchs E (2014) Emerging interactions between skin stem cells and their
- 768 niches. Nat Med 20: 847-56
- Hsu YC, Pasolli HA, Fuchs E (2011) Dynamics between stem cells, niche, and progeny
- 770 in the hair follicle. Cell 144: 92-105
- 771 Ingolia NT, Lareau LF, Weissman JS (2011) Ribosome profiling of mouse embryonic
- stem cells reveals the complexity and dynamics of mammalian proteomes. Cell 147:
- 773 789-802
- James C, Zhao TY, Rahim A, Saxena P, Muthalif NA, Uemura T, Tsuneyoshi N, Ong
- 775 S, Igarashi K, Lim CY, Dunn NR, Vardy LA (2018) MINDY1 Is a Downstream Target
- of the Polyamines and Promotes Embryonic Stem Cell Self-Renewal. Stem Cells 36:
- 777 1170-1178
- 778 Jensen UB, Yan X, Triel C, Woo SH, Christensen R, Owens DM (2008) A distinct
- population of clonogenic and multipotent murine follicular keratinocytes residing in the
- 780 upper isthmus. J Cell Sci 121: 609-17
- 781 Kakegawa T, Guo Y, Chiba Y, Miyazaki T, Nakamura M, Hirose S, Canellakis ZN,
- 782 Igarashi K (1991) Effect of acetylpolyamines on in vitro protein synthesis and on the
- 783 growth of a polyamine-requiring mutant of Escherichia coli. J Biochem 109: 627-31
- 784 Kim CS, Ding X, Allmeroth K, Kolenc OI, L'Hoest N, Chacon-Martinez CA, Edlich-Muth
- 785 C, Giavalisco P, Quinn KP, Denzel MS, Eming SA, Wickstrom SA (2019) An mTORC2-
- 786 dependent switch to glutamine metabolism controls stem cell fate reversibility and
- 787 long-term maintenance in the hair follicle. Manuscript submitted for publication
- 788 Kristensen AR, Gsponer J, Foster LJ (2013) Protein synthesis rate is the predominant
- 789 regulator of protein expression during differentiation. Mol Syst Biol 9: 689
- 790 Landau G, Bercovich Z, Park MH, Kahana C (2010) The role of polyamines in
- 791 supporting growth of mammalian cells is mediated through their requirement for
- 792 translation initiation and elongation. J Biol Chem 285: 12474-81

- 793 Li A, Simmons PJ, Kaur P (1998) Identification and isolation of candidate human
- keratinocyte stem cells based on cell surface phenotype. Proc Natl Acad Sci U S A 95:
- 795 3902-7
- 796 Liu L, Michowski W, Inuzuka H, Shimizu K, Nihira NT, Chick JM, Li N, Geng Y, Meng
- 797 AY, Ordureau A, Kolodziejczyk A, Ligon KL, Bronson RT, Polyak K, Harper JW, Gygi
- 798 SP, Wei W, Sicinski P (2017) G1 cyclins link proliferation, pluripotency and
- 799 differentiation of embryonic stem cells. Nat Cell Biol 19: 177-188
- 800 Love MI, Huber W, Anders S (2014) Moderated estimation of fold change and
- dispersion for RNA-seq data with DESeq2. Genome Biol 15: 550
- 802 Lu R, Markowetz F, Unwin RD, Leek JT, Airoldi EM, MacArthur BD, Lachmann A,
- 803 Rozov R, Ma'ayan A, Boyer LA, Troyanskaya OG, Whetton AD, Lemischka IR (2009)
- 804 Systems-level dynamic analyses of fate change in murine embryonic stem cells.
- 805 Nature 462: 358-62
- 806 Metcalf BW, Bey P, Danzin C, Jung MJ, Casara P, Vevert JP (1978) Catalytic
- 807 irreversible inhibition of mammalian ornithine decarboxylase (E.C.4.1.1.17) by
- 808 substrate and product analogs. Journal of the American Chemical Society 100: 2551-
- 809 2553
- 810 Nancarrow MJ, Nesci A, Hynd PI, Powell BC (1999) Dynamic expression of ornithine
- 811 decarboxylase in hair growth. Mech Dev 84: 161-4
- Noga MJ, Dane A, Shi S, Attali A, van Aken H, Suidgeest E, Tuinstra T, Muilwijk B,
- 813 Coulier L, Luider T, Reijmers TH, Vreeken RJ, Hankemeier T (2012) Metabolomics of
- 814 cerebrospinal fluid reveals changes in the central nervous system metabolism in a rat
- 815 model of multiple sclerosis. Metabolomics: Official journal of the Metabolomic Society
- 816 8: 253-263
- Odenlund M, Holmqvist B, Baldetorp B, Hellstrand P, Nilsson BO (2009) Polyamine
- 818 synthesis inhibition induces S phase cell cycle arrest in vascular smooth muscle cells.
- 819 Amino Acids 36: 273-82
- Park MH, Cooper HL, Folk JE (1981) Identification of hypusine, an unusual amino acid,
- in a protein from human lymphocytes and of spermidine as its biosynthetic precursor.
- 822 Proc Natl Acad Sci U S A 78: 2869-73

- Parry L, Balana Fouce R, Pegg AE (1995) Post-transcriptional regulation of the content
- of spermidine/spermine N1-acetyltransferase by N1N12-bis(ethyl)spermine. Biochem
- 825 J 305 (Pt 2): 451-8
- Pegg AE (2016) Functions of Polyamines in Mammals. J Biol Chem 291: 14904-12
- Pieragostino D, D'Alessandro M, di Ioia M, Rossi C, Zucchelli M, Urbani A, Di Ilio C,
- 828 Lugaresi A, Sacchetta P, Del Boccio P (2015) An integrated metabolomics approach
- 829 for the research of new cerebrospinal fluid biomarkers of multiple sclerosis. Molecular
- 830 bioSystems 11: 1563-72
- Quinlan AR, Hall IM (2010) BEDTools: a flexible suite of utilities for comparing genomic
- 832 features. Bioinformatics 26: 841-2
- 833 Ray RM, Zimmerman BJ, McCormack SA, Patel TB, Johnson LR (1999) Polyamine
- depletion arrests cell cycle and induces inhibitors p21(Waf1/Cip1), p27(Kip1), and p53
- 835 in IEC-6 cells. Am J Physiol 276: C684-91
- 836 Roux PP, Topisirovic I (2018) Signaling Pathways Involved in the Regulation of mRNA
- 837 Translation. Mol Cell Biol 38
- 838 Ruiz S, Panopoulos AD, Herrerias A, Bissig KD, Lutz M, Berggren WT, Verma IM,
- 839 Izpisua Belmonte JC (2011) A high proliferation rate is required for cell reprogramming
- and maintenance of human embryonic stem cell identity. Curr Biol 21: 45-52
- 841 Saini P, Eyler DE, Green R, Dever TE (2009) Hypusine-containing protein eIF5A
- 842 promotes translation elongation. Nature 459: 118-21
- 843 Sampath P, Pritchard DK, Pabon L, Reinecke H, Schwartz SM, Morris DR, Murry CE
- 844 (2008) A hierarchical network controls protein translation during murine embryonic
- stem cell self-renewal and differentiation. Cell Stem Cell 2: 448-60
- 846 Sanchez-Lopez J, Camanes G, Flors V, Vicent C, Pastor V, Vicedo B, Cerezo M,
- 847 Garcia-Agustin P (2009) Underivatized polyamine analysis in plant samples by ion pair
- 848 LC coupled with electrospray tandem mass spectrometry. Plant physiology and
- 849 biochemistry: PPB 47: 592-8
- 850 Schmidt EK, Clavarino G, Ceppi M, Pierre P (2009) SUnSET, a nonradioactive method
- 851 to monitor protein synthesis. Nat Methods 6: 275-7

- 852 Seiler N, Dezeure F (1990) Polyamine transport in mammalian cells. Int J Biochem 22:
- 853 211-8
- 854 Sela Y, Molotski N, Golan S, Itskovitz-Eldor J, Soen Y (2012) Human embryonic stem
- 855 cells exhibit increased propensity to differentiate during the G1 phase prior to
- phosphorylation of retinoblastoma protein. Stem Cells 30: 1097-108
- 857 Signer RA, Magee JA, Salic A, Morrison SJ (2014) Haematopoietic stem cells require
- a highly regulated protein synthesis rate. Nature 509: 49-54
- 859 Soler AP, Gilliard G, Megosh LC, O'Brien TG (1996) Modulation of murine hair follicle
- function by alterations in ornithine decarboxylase activity. J Invest Dermatol 106: 1108-
- 861 13
- 862 Sonnenberg A, Calafat J, Janssen H, Daams H, van der Raaij-Helmer LM, Falcioni R,
- Kennel SJ, Aplin JD, Baker J, Loizidou M, et al. (1991) Integrin alpha 6/beta 4 complex
- is located in hemidesmosomes, suggesting a major role in epidermal cell-basement
- 865 membrane adhesion. J Cell Biol 113: 907-17
- 866 Sunkara PS, Ramakrishna S, Nishioka K, Rao PN (1981) The relationship between
- levels and rates of synthesis of polyamines during mammalian cell cycle. Life Sci 28:
- 868 1497-506
- Tahmasebi S, Amiri M, Sonenberg N (2018a) Translational Control in Stem Cells. Front
- 870 Genet 9: 709
- 871 Tahmasebi S, Khoutorsky A, Mathews MB, Sonenberg N (2018b) Translation
- deregulation in human disease. Nat Rev Mol Cell Biol 19: 791-807
- 873 Trempus CS, Morris RJ, Bortner CD, Cotsarelis G, Faircloth RS, Reece JM, Tennant
- 874 RW (2003) Enrichment for living murine keratinocytes from the hair follicle bulge with
- the cell surface marker CD34. J Invest Dermatol 120: 501-11
- 876 Tsai YH, Lin KL, Huang YP, Hsu YC, Chen CH, Chen Y, Sie MH, Wang GJ, Lee MJ
- 877 (2015) Suppression of ornithine decarboxylase promotes osteogenic differentiation of
- human bone marrow-derived mesenchymal stem cells. FEBS Lett 589: 2058-65
- Uimari A, Keinanen TA, Karppinen A, Woster P, Uimari P, Janne J, Alhonen L (2009)
- 880 Spermine analogue-regulated expression of spermidine/spermine N1-

882

883

884

885

886

887

888

889

890

891

892

893

894

895

896

897

898

899

acetyltransferase and its effects on depletion of intracellular polyamine pools in mouse fetal fibroblasts. Biochem J 422: 101-9 Wallace HM, Fraser AV, Hughes A (2003) A perspective of polyamine metabolism. Biochem J 376: 1-14 Yamashita T, Nishimura K, Saiki R, Okudaira H, Tome M, Higashi K, Nakamura M, Terui Y, Fujiwara K, Kashiwagi K, Igarashi K (2013) Role of polyamines at the G1/S boundary and G2/M phase of the cell cycle. Int J Biochem Cell Biol 45: 1042-50 Zhang D, Zhao T, Ang HS, Chong P, Saiki R, Igarashi K, Yang H, Vardy LA (2012) AMD1 is essential for ESC self-renewal and is translationally down-regulated on differentiation to neural precursor cells. Genes Dev 26: 461-73 Zhao T, Goh KJ, Ng HH, Vardy LA (2012) A role for polyamine regulators in ESC selfrenewal. Cell Cycle 11: 4517-23 Zhou Y, Zhou B, Pache L, Chang M, Khodabakhshi AH, Tanaseichuk O, Benner C, Chanda SK (2019) Metascape provides a biologist-oriented resource for the analysis of systems-level datasets. Nat Commun 10: 1523 Zismanov V, Chichkov V, Colangelo V, Jamet S, Wang S, Syme A, Koromilas AE, Crist C (2016) Phosphorylation of elF2alpha Is a Translational Control Mechanism Regulating Muscle Stem Cell Quiescence and Self-Renewal. Cell Stem Cell 18: 79-90

Supplemental Figures

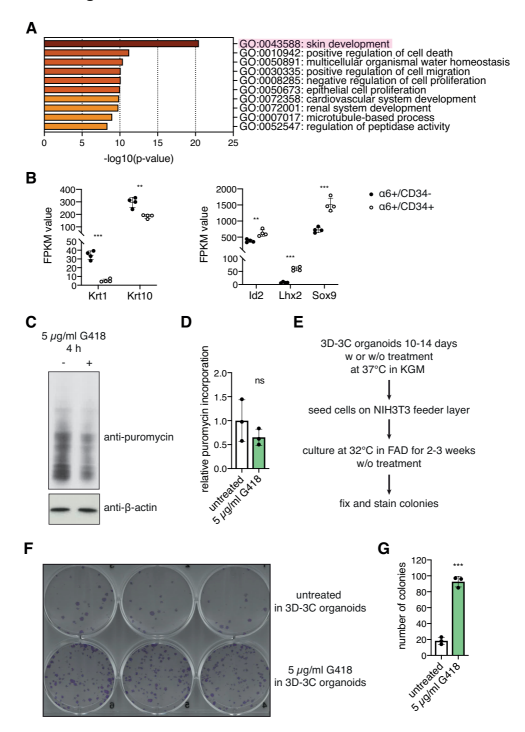


Figure S1: RNA-Sequencing confirms cell identity, which can be influenced by G418 treatment in the 3D-3C organoids. (A) GO term analysis of differentially expressed genes (p-value < 0.05, log2FC > ± 0.5) from RNA-Seq experiment comparing $\alpha 6+$ /CD34- progenitor cells with $\alpha 6+$ /CD34+ stem cells (biological process, metascape.org). (B) FPKM values of differentiation (K1, K10) and stem cell marker genes (Id2, Lhx2, Sox9). Mean \pm SD (n=4). *** p<0.001; ** p<0.01 (t test). (C) Representative Western blot analysis after puromycin incorporation in 3D-3C cultured cells. G418 treatment was performed for the last 4 h of the culture. (D) Quantification of the Western blot in (C). Mean \pm SD (n=3). ns: not significant (t test). (E) Schematic representation of the workflow for colony formation assay after 3D-3C organoid culture. (F) Representative image of tissue culture plate after colony formation assay using cells with or without G418 treatment in 3D-3C organoids (n=3). (G) Quantification of colony number in (F). Mean \pm SD (n=3). **** p<0.001 (t test).

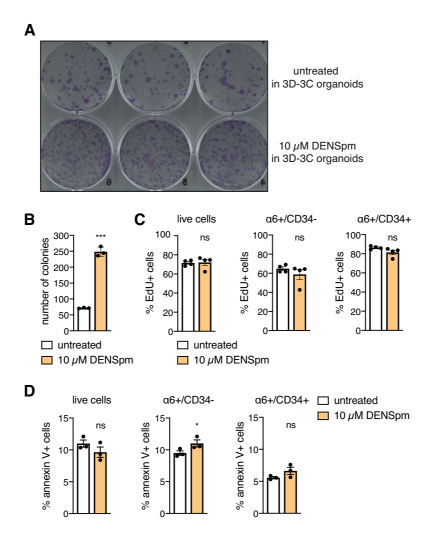


Figure S2: DENSpm treatment increases stem cell potency without major effects on proliferation or apoptosis. (A) Representative image of tissue culture plate after colony formation assay using cells with or without DENSpm treatment in 3D-3C organoids (n=2). (B) Quantification of colony number in (A). Mean \pm SD (n=3). *** p<0.001 (t test). (C) Ratio of EdU+ cells with or without DENSpm treatment. EdU was incorporated for 24 h. Mean \pm SEM (n=4). ns: not significant (t test). (D) Ratio of annexin V+ cells with or without DENSpm treatment. Mean \pm SEM (n=4). * p<0.05; ns: not significant (t test).

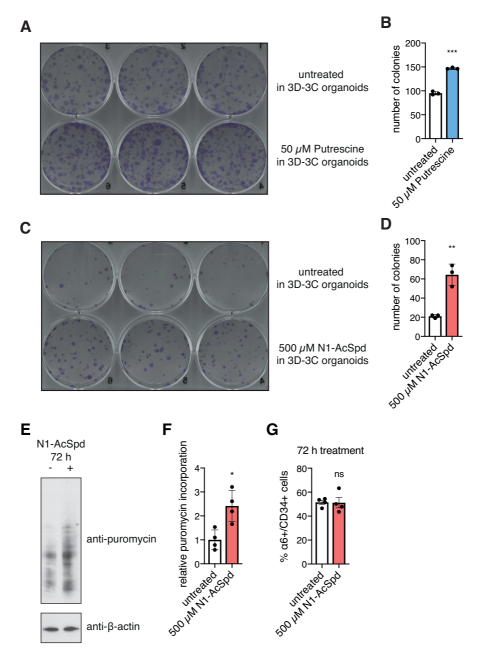


Figure S3: N1-acetylspermidine treatment increases stem cell potency without reducing translation. (A) Representative image of tissue culture plate after colony formation assay using cells with or without putrescine treatment in 3D-3C organoids (n=2). (B) Quantification of colony number of plate in (A). Mean \pm SD (n=3). (C) Representative image of tissue culture plate after colony formation assay using cells with or without N1-AcSpd treatment in 3D-3C organoids (n=3). (D) Quantification of colony number of plate in (C). Mean \pm SD (n=3). (E) Representative Western blot analysis after puromycin incorporation in 3D-3C cultured cells. N1-AcSpd treatment (500 µM) was performed for the last 72 h of the culture. (F) Quantification of the Western blot in (E). Mean \pm SD (n=4). (G) Ratio of α 6+/CD34+ cells after two weeks of 3D-3C culture with or without N1-AcSpd treatment for the last 72 h. Mean \pm SEM (n=4). (A-G) Statistical significance was calculated by t test. *** p<0.001, ** p<0.05, ns: not significant.

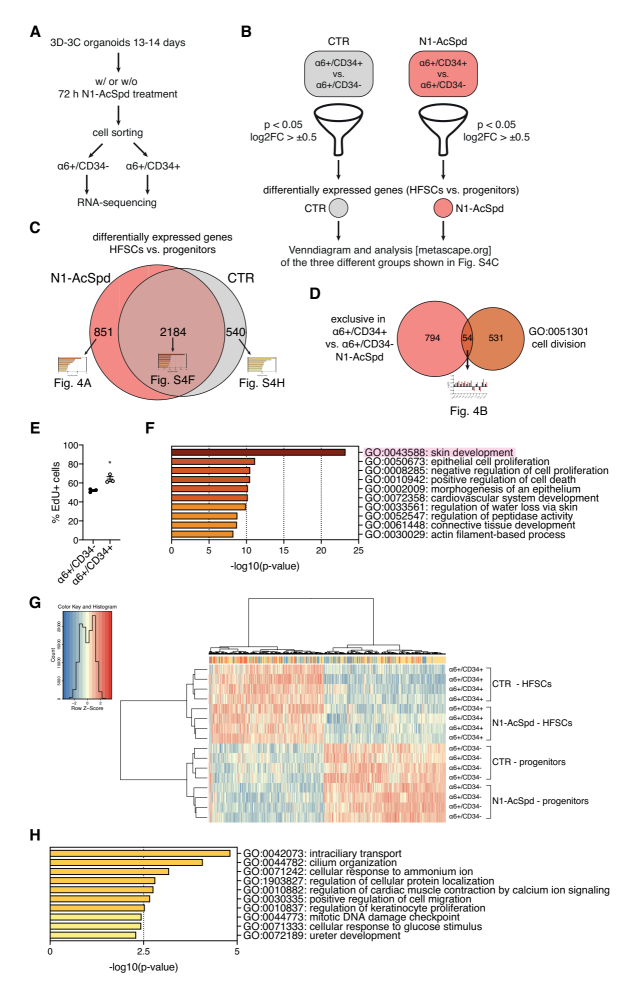


Figure S4: Schematic representation of the workflow of RNA sequencing analysis, which revealed maintained cell identity upon N1-acetylspermidine treatment. (A) Schematic representation of the workflow for RNA-seq sample collection. (B) Schematic representation of the bioinformatic workflow. α 6+/CD34- cells and α 6+/CD34+ cells were compared for untreated and treated conditions. Genes were filtered (p-value < 0.05, log2FC > \pm 0.5) and the resulting lists were used for further analysis. Control (CTR) is shown in gray, N1-AcSpd is depicted in red. (C) Venn diagram of the two groups from (B). CTR is shown in gray, N1-AcSpd is depicted in red. (D) Venn diagram of differentially expressed genes upon N1-AcSpd treatment (red) and genes covered by GO term cell division (brown). (E) Ratio of EdU+ cells in α 6+/CD34- progenitors and α 6+/CD34+ HFSCs. Mean \pm SEM (n=4). * p<0.05 (t test). (F) GO term analysis of differentially expressed genes between cell types (p-value < 0.05, log2FC > \pm 0.5) from RNA-seq experiment common in untreated and treated cells (overlap shown in (C), biological process, metascape.org). (G) Heat map showing differentially expressed genes with a p-value \leq 0.05 (α 6+/CD34- vs. α 6+/CD34+; treated or untreated). (H) GO term analysis of differentially expressed genes between cell types (p-value < 0.05, log2FC > \pm 0.5) from RNA-seq experiment only in untreated cells (gray in (C), biological process, metascape.org).