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## Biomimetic tissue models reveal the role of hyaluronan in melanoma proliferation and invasion†

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Interactions of hyaluronan (HA) and tumor and stromal cells are highly discussed as one of the major contributors in tumor progression and metastasis. The balance of HA in the tissue is highly regulated by two key enzyme classes; hyaluronan synthases (HAS) and hyaluronidases (HYAL). Current reports hint that the HA amount in the tissue is correlated with poor prognosis in melanoma, the most life-threatening skin tumor. In this work, we generated *in vivo* mouse models with low and high expression of Has2 and used the models for studying melanoma proliferation of the B78D14 melanoma cell line. We found that a strong reduction of HA amount in the skin was correlated to decreased tissue stiffness and a reduction in tumor weight. Since tumor cells have a direct contact to the HA in the tumor and at the stroma interface, we reconstituted different biomimetic *in vitro* models using fibroblasts derived from a mouse model to recapitulate melanoma cell behavior at the tumor boundary, namely, (i) decellularized fibroblast matrix (FbECM), (ii) fibroblast embedded into 3D collagen matrices (FbColl), and (iii) well-defined HA-functionalized 3D collagen matrices (HAColl). We found no considerable effect of high and low amounts of fibroblast-derived HA in the matrices on melanoma proliferation and invasion. However, HYAL1-treated FbECM and FbColl, as well as HAColl functionalized with low molecular weight HA (34 kDa) promoted proliferation and invasion of melanoma cells in a concentration dependent manner. Our results emphasize the molecular weight specific effects of HA in regulation of melanoma behavior and provide an alternative explanation for the *in vivo* observation of HA dependent tumor growth.

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## 1. Introduction

Tumor progression relies on a close interaction of tumor cells, different cell types in the tumor, and the surrounding stromal matrix. The stroma of solid tumors represents the benign vicinity of tumor cells, as also in malignant melanoma. However, compared to normal tissue, tumor stroma is different and originates from cells which are altered by paracrine interactions with tumor cells. It consists of the stromal matrix with (micro-

vessels and embedded stromal cells (immune cells, macrophages, fibroblasts, myofibroblasts) being controlled by paracrine mediators as well as deposited matrix components. Bidirectional interactions of tumor cells and the stroma decisively influence growth and metastatic behavior of many solid tumors. Cells or matrix components of the tumor stroma contribute to important aspects of tumor progression including proliferation, migration, and induction of angiogenesis.<sup>1–4</sup>

Hyaluronan (HA) is a major glycosaminoglycan of the extracellular matrix (ECM) fulfilling essential structural and functional roles during development,<sup>5</sup> inflammation,<sup>3,6</sup> repair<sup>7,8</sup> and tumor progression.<sup>4,9–11</sup> HA accumulation in tumors is often associated with poor prognosis. It is synthesized by three HA synthases (HAS1–3), where HAS2 is most strongly expressed in many tissues.<sup>12,13</sup> Many effects of HA in tumor biology rely on the interaction with its main cellular receptors CD44 (and tumor specific splice variants) and HA-mediated motility receptor (RHAMM).<sup>14,15</sup>

HA degradation is mediated either by free radical-related depolymerization occurring in the presence of reactive oxygen species (ROS)<sup>16</sup> and Maillard products<sup>17</sup> or enzymatically by hyaluronidases.<sup>4,18</sup> Several enzymes are described (HYAL1–3,

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PH20, CEMIP, TMEM2),<sup>19–23</sup> and the impact of tumor-derived or stroma-associated hyaluronidases can be both: pro- and anti-tumorigenic.<sup>24</sup> HYAL1 on one hand hydrolyses pro-tumorigenic HA, but as a result small HA-oligosaccharides may activate cells in the tumor and the stroma, thus supporting tumor progression and angiogenesis<sup>25</sup> by induction of matrix metalloproteinases,<sup>26,27</sup> as well as lymph node metastasis.<sup>28</sup> HA-oligosaccharides may act *via* toll like receptor signaling<sup>27,29,30</sup> or by interfering with the interaction of high molecular weight HA (HMW-HA) and its receptors CD44 and RHAMM.<sup>31,32</sup> Moreover, the relevance of small HA-oligosaccharide products was suggested when Schmaus *et al.*<sup>33</sup> quantified these in tumor interstitial fluids and found a correlation with invasion of tumor cells into lymphatic vessels and the formation of lymph node metastases in colorectal cancer patients. However, recent data suggest that HYAL1 may support tumor metastasis independently of the generation of small HA-oligosaccharide products thus leaving alternative ways of action conceivable.<sup>34</sup> The prevalence of other HA degrading enzymes, like CEMIP and TMEM2, obviously also correlates with progression or metastasis in tumor patients,<sup>22,35</sup> whereas the small HA-oligosaccharide molecules were not quantified in these studies. Thus, today it remains to prove that really the HA-degradation products are the tumor promoting molecules or whether other mechanisms involving these proteins lead to cancer progression.

Due to its high water-binding capacity, HA deposition strongly contributes to specific physico-chemical properties of the surrounding stroma sensed by the tumor cells. Besides paracrine signals, the behavior of tumor cells is strongly affected by changes of mechanics and network microstructure of the fibrillar collagen matrix, as well as its compositions.<sup>36</sup> We have previously demonstrated that changing physical properties of a three-dimensional (3D) collagen biomimetic matrix can influence tumor cells' behavior, while the chemical composition of the model matrix remained unchanged.<sup>37,38</sup> Such findings resulted in the awareness that more complex 3D culture models are essential in order to receive data better resembling the *in vivo* situation than the commonly used 2D cultures on tissue culture plastics.<sup>36,39,40</sup>

In order to investigate the impact of stromal HA on the physico-chemical properties of the stroma and on tumor cells facing this defined matrix, we combined model systems of graded complexity and *in vitro* experiments using stromal mouse fibroblasts with strongly decreased HA-synthesis due to genetic knockdown of the main HA synthase 2 (*Has2*). The impact of stromal HA on melanoma cells was characterized in respect to invasive behavior, proliferation and gene expression of the tumor cells and compared to *in vivo* findings in a *Has2* knockdown mouse model on melanoma tumor growth.

## 2. Experimental

### 2.1. Generation of inducible *Has2* knockout mice

All animal procedures were performed in accordance with the Guidelines for Care and Use of Laboratory Animals of Leipzig

University and approved by the Animal Ethics Committee of Saxony/Germany (TVV 57/14, T26/16). *Has2*-Exon2-floxed, C57BL/6N-embryonic-cell-line JM8A3.N1 was purchased from KOMP Repository, University of California Davis and Children's Hospital Oakland, Research Institute. Chimeric C57BL/6 mice were generated by Konstantinos Anastassiadis (TU Dresden, BIOTEC, Genetic Engineering of Stem Cells Group) and crossed with Flp-deleter C57BL/6 mice to generate offspring with deleted neomycin resistance and deleted lacZ reporter gene but carrying heterozygous floxed *Has2*-Exon2. Exon 2 was chosen, as it contains the start of translation and a 209 amino acid sequence which is the major part of the HAS2 proteins, see also Ensembl (ensembl.org). This region contains domains with the core enzymatic activity (nucleotide-diphospho-sugar transferases). Backcrossed homozygous mice carrying the *Has2*-Exon2-floxed alleles develop and reproduce without any obvious phenotype (ESI Fig. S1†). *Has2*-Exon2-floxed mice were backcrossed with ubiquitous Cre-deleter C57BL/6 mice (human ubiquitin C (UBC) promoter sequence upstream of a Cre-ERT2 fusion gene; Stock no.: 008085; Jackson Laboratory, USA) to *Has2*<sup>flox/flox</sup>-Cre-ERT2<sup>+/-</sup> strain, ready to induce *Has2*-Exon2 deletion upon tamoxifen application (mice and fibroblasts were named '*Has2*-KD' after 4-hydroxy-tamoxifen treatment). Because characterization of the intended *Has2*-knockout mice and fibroblasts thereof showed remaining *Has2* expression, they were regarded as a *Has2*-knockdown and hence named '*Has2*-KD' throughout this work. Control mice were only *Has2*<sup>flox/flox</sup> without inducible Cre expression and mice and fibroblasts were named '*Ctrl*' after 4-hydroxy-tamoxifen induction.

### 2.2. Isolation of primary fibroblast from mouse skin tissue

Primary fibroblasts were isolated from full thickness skin from the back of the mice, from both *Has2*<sup>flox/flox</sup>-Cre-ERT2<sup>+/-</sup> and *Has2*<sup>flox/flox</sup>. Shaved back skin was disinfected with 70% isopropanol before approximately 4 cm<sup>2</sup> of the skin was excised and cut into small pieces. To isolate fibroblasts from the tissue, 26 U of Liberase™ DL Research Grade (Sigma-Aldrich, Steinheim, Germany) were added to 3 mL serum-free DMEM (Biochrom, Berlin, Germany) and incubated over 2 h at 37 °C while stirring. Afterwards, the cell suspension was filtered through a 70 µm cell strainer (Becton Dickinson, Heidelberg, Germany) and the filtrate centrifuged at 300g for 4 minutes at room temperature. Collected cells were plated on cell culture Petri dishes and cultivated in DMEM (Biochrom) supplemented with 10% fetal calf serum (Biochrom) and 1% ZellShield (Biochrom) at 37 °C, 5% CO<sub>2</sub> and 95% humidity for 24 h. Then, cells were rinsed with PBS to remove any dead and unattached cells. After another 24 h of cultivation, cells were used for *in vitro* experiments up to the 4th passage.

### 2.3. Induction and characterization of *Has2*-KD fibroblasts

Isolated fibroblasts from *Has2*<sup>flox/flox</sup>-Cre-ERT2<sup>+/-</sup> and *Has2*<sup>flox/flox</sup> were treated with 2 µM 4-hydroxy-tamoxifen (4-OHT, Sigma-Aldrich) in DMEM (Biochrom) supplemented with 10% fetal calf serum (Biochrom) and 1% ZellShield (Biochrom) for 48 h



at 37 °C, 5% CO<sub>2</sub> and 95% humidity to induce Cre-recombinase translocation into the nucleus in *Has2*<sup>flox/flox</sup>-Cre-ERT2<sup>+/-</sup> fibroblasts for *Has2*-Exon2 deletion (*Has2*-KD fibroblasts). To verify the deletion, DNA and RNA were analyzed after 48 h culture without 4-OHT. Presence of *Has2*-Exon2 was investigated by amplification of *Has2*-Exon2 gene locus and agarose-gel-electrophoresis. *Has2* mRNA expression was quantified using RT-qPCR and. Sequences of primers using for the PCR are listed in the ESI Table 1.†

#### 2.4. Tumor injection in mouse models

*Has2*<sup>flox/flox</sup>-Cre-ERT2<sup>+/-</sup> and *Has2*<sup>flox/flox</sup> mice at the age of 8–12 weeks were intraperitoneally injected with 1 mg 4-OHT dissolved in 100 µL 10% absolute ethanol (vwr, Dresden, Germany) in sunflower seed oil from *Helianthus annuus* (Sigma-Aldrich) per day for 8 days. At the end of 4-OHT treatment, the mice recovered for 5 days before  $1.0 \times 10^6$  B78D14 cells (resuspended in 50 µL PBS (Biochrom)) were injected intradermally into the lower back. B78D14 tumors grew for 16 days, until tumor diameter reached 10 mm. Healthy skin from the back and tumor samples were taken for further analysis.

#### 2.5. Reconstitution of fibroblast-derived matrix (FbECM)

Fibroblast-derived matrix (FbECM) were reconstituted according to an established protocol.<sup>41</sup> Briefly, cover slips (13 mm in diameter, VWR) were coated with cross-linked gelatin before seeding  $1.6 \times 10^5$  *Has2*-KD or *Ctrl* fibroblasts on top at 100% confluency. After 24 h, cell culture medium was replaced with DMEM media supplemented with 50 µg mL<sup>-1</sup> ascorbic acid for 9 days. Cell culture media was changed every 3 day of culture to enhance ECM production. Afterwards, fibroblasts were lysed using ammonium hydroxide (20 mM) and Triton X-100 (0.5% v/v) in PBS. Extracellular DNA was digested with 100 U mL<sup>-1</sup> DNase (Roche, Mannheim, Germany) in DNase buffer (50 mM Tris, 10 mM MnCl<sub>2</sub>, 1 mM CaCl<sub>2</sub>, at pH 7.5) for 15 min at RT. After carefully washing with PBS, intact FbECMs were kept in PBS containing penicillin/streptomycin (100 U mL<sup>-1</sup>, 100 µg mL<sup>-1</sup> respectively, Biochrom) prior to use for cell experiments. As HA-negative controls, FbECMs were incubated with HYAL1 (2 U mL<sup>-1</sup>, Hyaluronidase-1 from bovine testis, Sigma-Aldrich) in 20 mM sodium acetate buffer for 2 h at 37 °C after DNase treatment.

#### 2.6. Reconstitution of fibroblast-embedded 3D collagen matrices (FbColl)

Fibroblast-embedded 3D collagen matrices were reconstituted on 13 mm poly(styrene-*alt*-maleic anhydride) (PSMA; MW 20 000–30 000 g mol<sup>-1</sup>, Sigma-Aldrich) coated coverslip by embedding  $1 \times 10^4$  *Has2*-KD and *Ctrl* fibroblasts into 3D collagen matrices. To reconstitution of collagen matrices, rat-tail type I collagen (Corning, NY, USA) was mixed with 250 mM phosphate buffer at pH 7.5, as previously reported.<sup>37,38</sup> The collagen fibrillation was introduced by incubation at 37 °C and 95% humidity. After 5 days of cultivation, cells were lysed with double-distilled water for 1 h at room temperature. As HA-negative controls, FbColl were incubated with HYAL1

(Hyaluronidase from bovine testis, Sigma-Aldrich) for 2 h at 37 °C after DNase treatment.

#### 2.7. Reconstitution of defined HA-immobilized collagen matrices (HAColl)

Firstly, 3D collagen matrices were reconstituted on 13 mm poly(styrene-*alt*-maleic anhydride) (PSMA; MW 20 000–30 000 g mol<sup>-1</sup>, Sigma-Aldrich) coated coverslip. Rat-tail type I collagen (Corning, USA) was mixed with 250 mM phosphate buffer at pH 7.5 to archive collagen concentration of 2 mg mL<sup>-1</sup>, as previously reported.<sup>37,38</sup> Matrices were fibrillated by incubation at 37 °C and 95% humidity. HA with a molecular weight of 34 kDa (low molecular weight HA, LMW-HA) and 1170 kDa (HMW-HA) were prepared in 100 mM MES buffer at pH 5. To immobilize defined HA amounts of 1.3 µg per matrix (0.02 µg HA per µg Coll) and 3.3 µg per matrix (0.41 µg HA per µg Coll), HA solutions in MES were prepared at 20 and 50 µg mL<sup>-1</sup> (LMW-HA) and 10 and 30 µg mL<sup>-1</sup> (HMW-HA), respectively. Reconstituted collagen matrices were incubated with 500 µL of HA solution for 2 h at room temperature. Afterwards, HA solution was discarded and 500 µL of 4 mg mL<sup>-1</sup> of *N*-(3-dimethylaminopropyl)-*N'*-ethylcarbodiimide hydrochloride (EDC; Sigma-Aldrich) dissolved in 100 mM in 2-(*N*-morpholino)ethanesulfonic acid (MES; Sigma-Aldrich) at pH 5 was directly added without washing. Matrices were incubated with EDC crosslinking solution for 2 h at room temperature and followed by washing 3 times each 5 min with PBS. Prior to cell experiments, matrices were equilibrated in cell culture medium for 1 h.

#### 2.8. Topological characterization of reconstituted matrices

To visualize and analyze topological parameters, cell-free FbColl and HAColl matrices were stained with 50 µM 5-(and 6)-carboxytetramethylrhodamine succinimidyl ester (5(6)-TAMRA-SE) (Invitrogen, Carlsbad, USA) at room temperature for 1 h and rinsed 3 times with PBS (Biochrom). Matrices were imaged using a confocal laser scanning microscope (cLSM, LSM700, Zeiss, Jena, Germany) using 40×/NA 1.3 oil immersion objective (Zeiss). Acquired images were 1024 × 1024 pixels in resolution (*xyz*-voxel size: 0.13 × 0.13 × 0.5 µm) and a vertical stack size of 200 images (equivalent to 100 µm). Pore size and fibril diameter were determined as previously described in ref. 38 and 42 using a home-built image analysis tool using an erosion algorithm and autocorrelation analysis, respectively.

#### 2.9. Mechanical characterization of full skin tissue and reconstituted matrices

Mechanical properties of full skin sample cross sections and reconstituted cell-free matrices were quantified by colloidal probe force spectroscopy using a scanning force microscope (NanoWizard 3, JPK Instruments, Berlin, Germany), as previously reported.<sup>37,38</sup> Briefly, a 50 µm glass microbead (Polyscience Europe GmbH, Eppelheim, Germany) was attached to a tipless HQ-CSC38 cantilever (NanoAndMore, Wetzlar, Germany) with a spring constant of approximately 0.1 N. At least 50 force–distance curves were mapped at an area of



100 × 100 μm at 3 positions of each sample from 3 independent tumor samples with indentation rate of 5 μm s<sup>-1</sup> in PBS buffer (Biochrom) at room temperature. An effective Young's modulus was determined as elastic modulus by fitting the retract part of force distance curves (typical indentation depth 5 μm) using the Hertz model.

### 2.10. Quantification and visualization of HA

For quantitative analysis, full-thickness skin samples, FbECM, and FbColl were lysed with 20 U mL<sup>-1</sup> protease from *Streptomyces griseus* (Sigma-Aldrich) and 5 mM deferoxamine mesylate salt (Sigma-Aldrich) solved in HA-prep buffer (150 mM Tris, 150 mM NaCl, 10 mM CaCl<sub>2</sub>, at pH 8.3). After shaking, overnight incubation at 55 °C, protease was inactivated by heating at 95 °C for 10 min. HA amount was quantified using commercial HA-ELISA kit (TECOMedical Group, Switzerland).

For visualization of HA, full-thickness skin samples, FbECM, and FbColl were stained with biotinylated hyaluronan binding protein (HABP, Sigma-Aldrich) followed by streptavidin-conjugated Alexa Fluor 488 (Invitrogen). HABP-stained matrices were imaged using cLSM (LSM700, Zeiss) using 40×/NA 1.3 oil immersion objective (Zeiss). Staining specificity was controlled both by pre-treatment with hyaluronidase or by omitting HABP incubation during staining procedure.

For HA optical quantification of HAColl, ATTO565- labelled HA was used to functionalize reconstituted matrix, as previously described.<sup>43</sup> Briefly, HMW-HA or LMW-HA derived from ozonolysis of high molecular weight HA (from *Streptococcus*, Aqua Biochem, Dessau, Germany) was conjugated with Atto565-NH<sub>2</sub> (ATTO-TEC, Siegen, Germany) at carboxylic groups along the HA chain. HAColl were digested with papain and fluorescence signal were obtained at excitation wavelength of 565 nm and emission wavelength of 590 nm using fluorescence spectrometer (TECAN Infinite 200 PRO, TECAN, Grödig, Austria). The fluorescence value was related to HA amount using a standard curve. Immobilized HA were visualized using cLSM (LSM700 (Zeiss)) using 40×/NA 1.3 oil immersion objective.

### 2.11. Melanoma cell proliferation and invasion in animal models, FbECM, FbColl, HAColl matrices

B78D14 murine malignant melanoma (MM) cell line was kindly provided by P. Holst (University Copenhagen, Denmark) (Becker *et al.* 1996).<sup>67</sup> B78D14 cell line is a derivative of the model cell line B16F10 characterized by slower growth behavior *in vivo* and *in vitro*. Cells were grown in DMEM supplemented with FCS (10%) (Biochrom). For experiments, cells were seeded on top of reconstituted matrices and cultivated in DMEM (Biochrom) supplemented with 10% fetal calf serum (Biochrom) and 1% ZellShield (Biochrom) at 37 °C, 5% CO<sub>2</sub> and 95% humidity for 3 days. Cell proliferation was determined either by *Ki67* mRNA expression (primer sequence is listed in ESI Table 1†) or by using commercially available EdU cell proliferation kits (Baseclick GmbH, Neuried, Germany).

*Ki67* antigen was detected in cryostat sections of experimental tumors that were fixed with ice-cold acetone (Carl-Roth, Germany). Anti-*Ki67* antibody (Bethyl Laboratories Inc., USA, IHC-00375) was added overnight (1:150). Bound antibody was detected with Alexa546 labelled goat-anti-rabbit secondary antibody (ThermoFisherScientific). Nuclei were stained with 4',6-diamidino-2-phenyl-indoldihydrochlorid (DAPI, Merck) and sections were mounted with Prolong Gold (ThermoFisherScientific). Microscopy was performed with Keyence BZ-9000E and corresponding software BZ-II Viewer, BZ-II Analyzer (Keyence, Neu Isenburg, Germany).

For cell invasion analysis, GFP-expressing B78D14 cells (transfected with pGFPmax (amaxa, Cologne, Germany) and FACS-sorted for strong expressing cells) or DAPI-stained cells were imaged by an epifluorescence AxioObserver Z1 microscope with scanning stage (Zeiss, Jena, Germany) using a 10× objective (Zeiss, Germany) in bright field and fluorescent mode (DAPI and Alexa Fluor 488). Cell invasion parameters, namely percentage of invaded cells and maximal invasion distance, were quantified using home-built MATLAB script (MATLAB 2019a, MathWorks, USA), as previously described in ref. 37 and 44. At least 600 cells from 3 positions per matrix conditions were analyzed.

### 2.12. Data analysis and statistics

Experiments were performed at least in triplicate, if not otherwise stated. Levels of statistical significance were determined by one-way ANOVA followed by Tukey's *post hoc* analysis using GraphPad Prism 7 (GraphPad Software, Inc., La Jolla, CA, USA). Significance level was set at 0.05.

## 3. Results and discussion

In this work, we aimed to decipher the impact of HA in melanoma progression by means of both *in vivo* mouse model and *in vitro* models. *Ctrl* and *Has2-KD* mice with injected B78D14 mouse melanoma cells and analyzed concerning tumor growth. These results were compared to three different biomimetic *in vitro* ECM models, namely (i) decellularized fibroblast matrix (FbECM), (ii) fibroblast embedded into 3D collagen matrices (FbColl), and (iii) well-defined HA-functionalized 3D collagen matrices (HAColl). Used fibroblasts were derived from the mouse models to recapitulate *Has2-KD* in ECM production and modification. B78D14 mouse melanoma cells were used to demonstrate the impact of HA in terms of cell proliferation and invasive behavior.

### 3.1. Animal model: *in vivo* mouse model with designated *Has2*-knockout

The *Has2-knockout* mouse was generated and characterized regarding HA production and mechanical properties of full-thickness skin. A significant, but incomplete knockout was achievable in these mice (hence, termed *Has2-knockdown* (*Has2-KD*) in the following sections). B78D14 mouse melanoma cells were injected into the lower back of *Ctrl* and *Has2-*

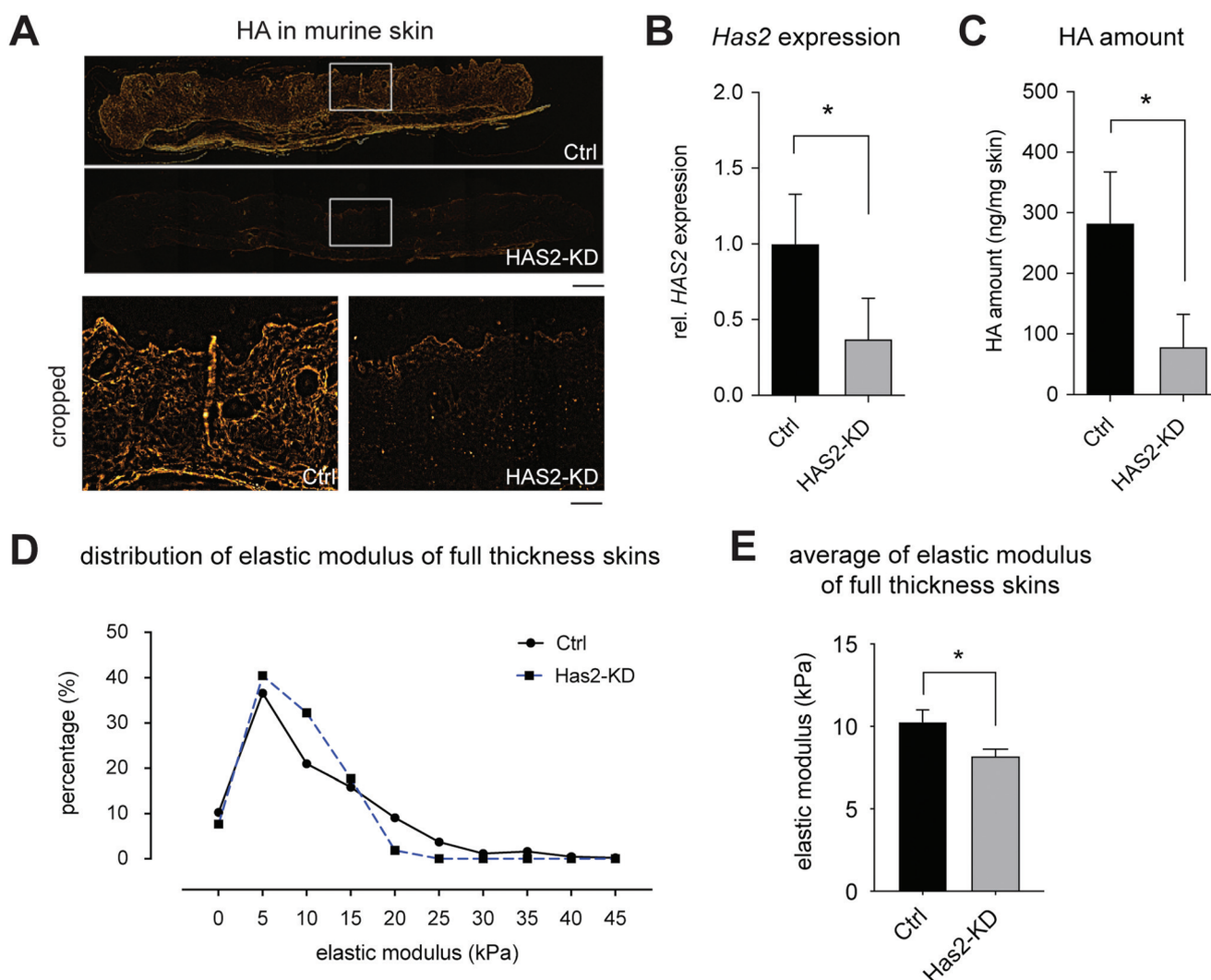


*KD* mice and were analyzed regarding proliferation by means of tumor weight and Ki67 staining.

**3.1.1. *Has2*-knockout induction led to *Has2*-knockdown (*Has2-KD*) and reduced HA deposition in skin.** *Has2-KD* as well as *Ctrl* mice showed no visual change in resting phenotype, as demonstrated by the animal weight (ESI Fig. S1†). To visualize the reduction of HA deposition in *Has2-KD* mouse, full-thickness skin cross-sections were stained with HA-binding protein (HABP) and imaged using cLSM (Fig. 1A). We found a strongly diminished HA signal in *Has2-KD* mice. Gene expression analysis of *Has2-Exon2* confirmed a significant decrease by 67% compared to *Ctrl* (Fig. 1B). PCR product at 257 bp confirmed the *Has2-Exon2* deletion on DNA level (ESI Fig. S2†). However, undeleted PCR products at 2440 bp within

the same sample could be observed, pointing out the partially remaining undeleted *Has2-Exon2* gene. This was also true in explanted fibroblasts with subsequent 4-OHT treatment (ESI Fig. S2†). Gene expression of isoenzymes *Has1* and *Has3* were found unchanged low in both groups (ESI Fig. S3†). Besides the *Has2* gene expression analysis, HA amount in skin was quantified (Fig. 1C). Skin HA concentration in *Has2-KD* mice was decreased by 72%, compared to *Ctrl*, which confirmed fluorescence microscopy analysis using HABP staining.

These results led to the conclusion of a significant but incomplete knockout (*Has2-KD*), where at least a fraction of fibroblasts was not affected by the 4-OHT induction. Although, HAS isoenzymes may still contribute to a small amount of HA production, there was no compensatory upregulation of either



**Fig. 1** *Has2*-knockdown led to depletion of hyaluronan and reduction of tissue stiffness. (A) Representative image of HA distribution in murine skin. HA was stained using HABP (scale bar = 1 mm and 100  $\mu$ m for upper and cropped images, respectively). (B) Relative expression of *Has2* gene in murine skin using RT-qPCR. (C) Quantification of HA amount in murine skin using HA-ELISA. (D) Distribution of elastic modulus of representative single murine skin samples determined at 50 different positions. (E) Elastic modulus of full thickness murine skin. The experiment was performed from 3 murine skins from *Ctrl* and *Has2-KD* mice, with a mean elastic modulus evaluated for each skin sample from force mapping as shown in (D) (data are represented as mean  $\pm$  SD; \* – significance level of  $p < 0.05$ ).



*Has1* or *Has3* mRNA. Also, remaining HA in *Has2-KD* skin might be a tightly integrated portion and/or enzymatically modified type which was not affected by HA turnover after *Has2-KD*. The fact that this model resulted in a variety of reduced HA concentrations in skin opened the opportunity to investigate HA impact on tumor growth *in vivo* in dependence on HA concentration.

### 3.1.2. *Has2-KD* mice exhibit reduced skin tissue stiffness.

The deposition of HA has been reported to alter tissue stiffness due to its ability to retain water, and in turn, regulate tumor cell behavior both in *in vitro* and *in vivo* models.<sup>36,45,46</sup> The elastic modulus of the skin tissue was quantified by colloidal probe force spectroscopy in a hydrated state. As shown in Fig. 1D and E, *Has2-KD* skin exhibited a reduction in mean tissue stiffness, in which especially sampling areas with higher stiffness were found less frequent. The result is well correlated to other reports, showing an increase in tissue stiffness with increasing HA amounts, both in *in vivo* models<sup>47</sup> and HA-functionalized 3D collagen matrices.<sup>11,48</sup>

### 3.1.3. Tumor progression was reduced in *Has2-KD* mice.

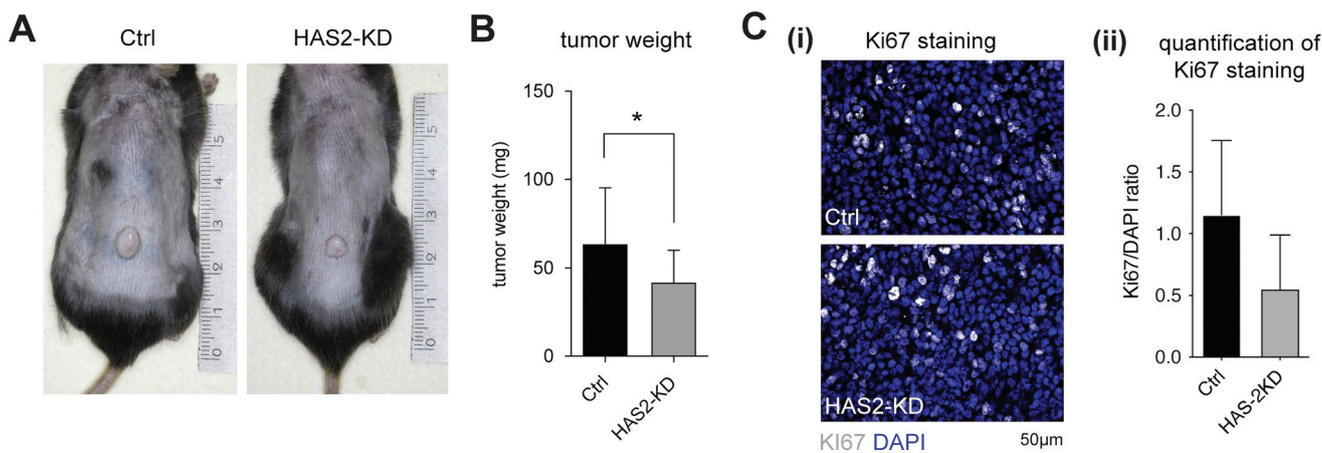
To study the melanoma progression as a function of HA depletion, B78D14 melanoma cells were intradermally injected into the lower back of *Ctrl* and *Has2-KD* mice. Tumor growth was observed for 16 days (Fig. 2A) and the solid tumor was dissected from the skin and weighted. As shown in Fig. 2B, *Has2-KD* mice showed a significant reduction in tumor weight compared to *Ctrl*. The decreased primary tumor weight in the knockdown condition supports the evidence that the high amount of HA in tumor microenvironment triggered tumor progression, as previously reported.<sup>10,49–51</sup> To analyze cell proliferation in the tumor, we stained the tumor with Ki67 antibodies and found a trend to increased Ki67 staining in the samples of *Ctrl* mice. Additionally, we found that cells at the tumor boundary exhibited higher Ki67 expression (ESI Fig. S4†), suggesting that cell-HA interactions at the tumor

boundary might drive the proliferation of melanoma cells. However, underlying mechanisms, on how different amounts of HA affect melanoma progression, remain unclear, since the mechanical properties of the tissue and HA availability are interlinked. Therefore, biomimetic tissue models which allow to decipher those parameters would help to understand the complexity of *in vivo* models. To address this underlying problem, we reversely engineered HA incorporated tissues using three different approaches to construct biomimetic matrices for *in vitro* analysis: (i) decellularized fibroblast matrix (FbECM), (ii) fibroblasts embedded into 3D collagen matrices (FbColl), (iii) and HA-functionalized 3D collagen matrices (HAColl), with degrading complexity in that order. Fibroblasts were derived from the mouse models to recapitulate *Has2-KD* in ECM production and modification. These models allowed us to mimic cell-HA interactions of cells at the tumor boundary in a well-defined manner.

## 3.2. Biomimetic tissue models *in vitro*

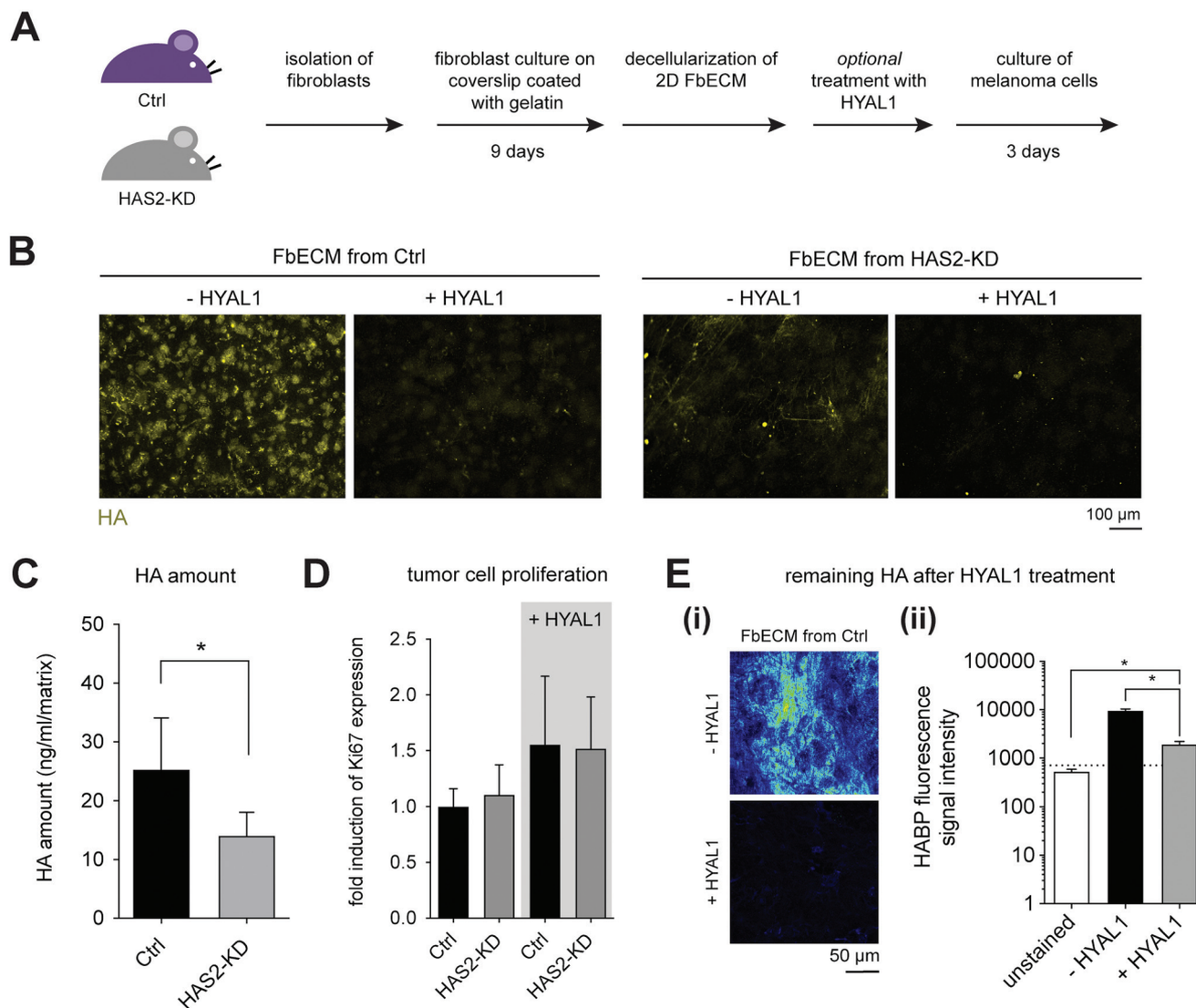
### 3.2.1. Decellularized fibroblast matrix (FbECM): HYAL1 treated FbECM enhanced proliferation of melanoma cells.

The process and timeline of the reconstitution of FbECM is shown in Fig. 3A. Briefly, fibroblasts from *Has2<sup>flx/flx</sup>-CreERT2* and control mice without CreERT2 were isolated from skin and cultured. To induce Cre translocation into nucleus with subsequent *Has2*-Exon2 knockdown, 4-OHT was added to cell culture medium. Afterwards, *Has2-KD* and *Ctrl* fibroblasts were stimulated to produce a fibroblast-derived ECM (FbECM). After FbECMs were decellularized, HA was visualized using HABP and characterized regarding HA amount using HA-ELISA kit. The efficiency of decellularization was checked by staining with DAPI and fluorescence microscopy (ESI Fig. S5†). As shown in Fig. 3B and C, average HA deposition in *Has2-KD* FbECM was reduced to 56% compared to control. The results revealed decreased HA amounts by both fluorescence



**Fig. 2** Suppressed growth of melanoma cells in *Has2-KD* mice. (A) Representative image of melanoma tumor *in vivo*. (B) Quantitative analysis of tumor weight from *Ctrl* and *Has2-KD* mouse after 16 days of injection. (C) Proliferation activity in the tumor was analyzed by fluorescence microscopy using DAPI (blue) and Ki67 antibody (white) staining. (C(i)) Representative image of the staining (scale bar: 50 μm) and (C(ii)) quantitative analysis of the Ki67 staining normalized against DAPI staining (data are represented as mean ± SD; \* – significance level of  $p < 0.05$ ).





**Fig. 3** HYAL1 digested FbECM matrices promote proliferation of melanoma cells. (A) Schematic illustration of the experimental workflow. (B) Representative images of FbECMs reconstituted using *Ctrl* and *Has2-KD* fibroblasts. Matrices were treated with or without HYAL1 and analyzed by cLSM using HABP staining. (C) Quantitative analysis of HA amount in FbECM using commercial HA-ELISA kit. (D) Proliferation analysis of melanoma cells by means of *Ki67* expression. (E) Analysis of remaining HA after HYAL1 treatment by cLSM using HABP staining. (E(i)) Representative image of FbECM from *Ctrl*. Color of the images is shown as heatmap. (E(ii)) Quantitative analysis of HABP fluorescence signal intensity. The dot line represents the fluorescence signal of FbECM stained with secondary antibody alone (data are represented as mean  $\pm$  SD; \* – significance level of  $p < 0.05$ ).

microscopy and quantitative analysis of HA amount using ELISA.

To study the impact of HA on melanoma cell proliferation, B78D14 cells were cultivated on decellularized FbECM for 5 days. The gene expression of *Ki67* was quantified to compare proliferative capability of B78D14 cells grown on different FbECM. Additionally, we used HYAL1 treated FbECM samples, as HA negative control due to the enzymatic degradation of existing HA by HYAL1. As shown in Fig. 3D, *Ctrl* and *Has2-KD* FbECM demonstrated no differences in *Ki67* expression. No remaining HA could be measured using HA-ELISA kit after HYAL1 digestion. However, low signal intensity was still detectable with HAPB using cLSM (Fig. 3E-ii). This finding suggests

remaining HA residues after treatment with HYAL1. Interestingly, cells cultivated on HYAL1-treated FbECMs of both *Ctrl* and *Has2-KD* fibroblasts showed an increase in proliferation (Fig. 3D). Hence, it is suggested that the residuals of low molecular HA might cause an increased proliferation of melanoma cells. It is interesting to note, that HYAL1 is known to be highly expressed in tumor microenvironments and it is suspected to contribute to tumor aggressiveness.<sup>34,52</sup>

Decreased HA deposition in *Has2-KD* FbECM was shown to have no impact on melanoma cell proliferation, but HYAL1 treatment of both FbECM types led to upregulated proliferation. However, the matrix organization and HA deposition by fibroblasts are not homogenous, as depicted in Fig. 3B, which

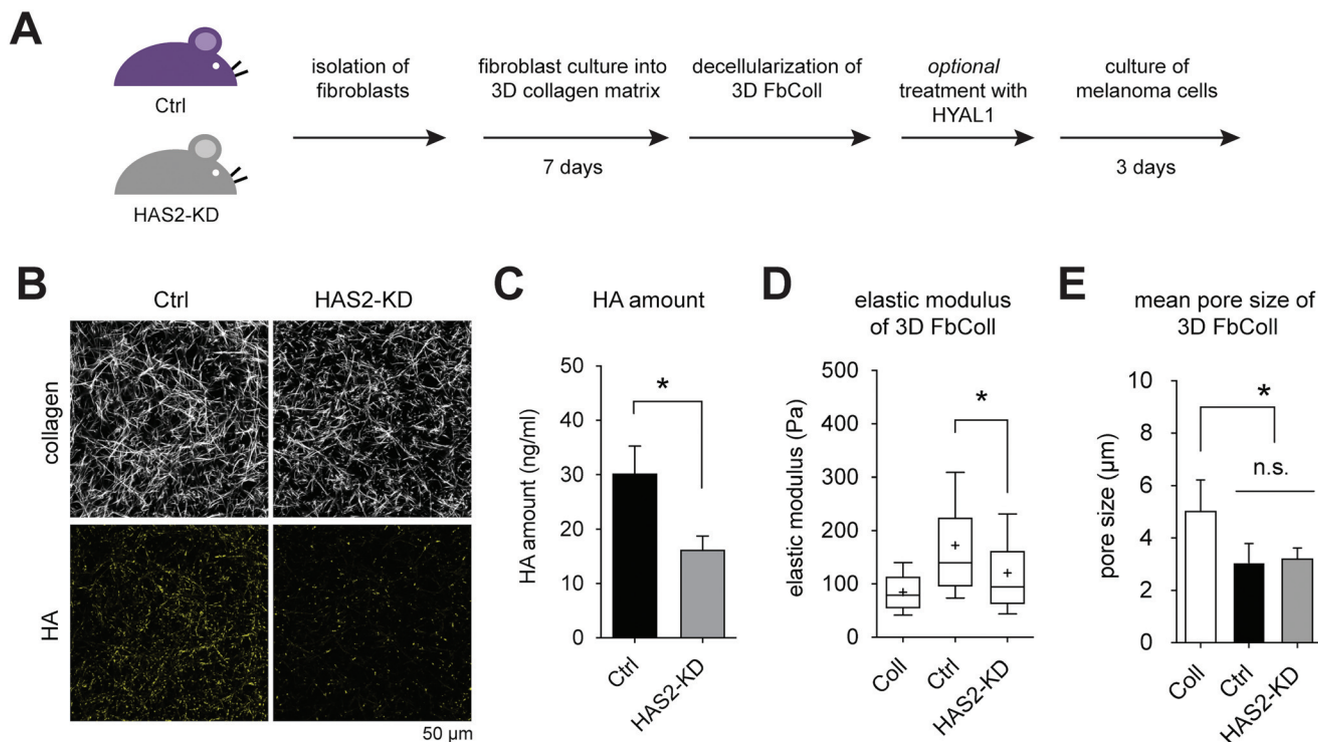


limits the comparison of melanoma behavior on FbECMs. Enhanced proteolysis of the FbECM by melanoma cells could be excluded by analyzing B78D14 melanoma cells growing on FbECM by mRNA expression microarray: none of the major MMPs was significantly regulated in these melanoma cells (data not shown). Additionally, we may not exclude that decrease of HA content in FbECM might lead also to changes in the deposition of collagens or fibronectin. However, microarray analysis of skin from *Ctrl* vs. *Has2-KD* mice did not reveal significant changes of mRNA levels for collagens I and III and fibronectin (data not shown). Furthermore, FbECM is a very thin ECM of approximately 3 to 7  $\mu\text{m}$ , which did not allow cell invasion studies and measurement of mechanical properties.

**3.2.2. Fibroblast embedded into 3D collagen matrices (FbColl): HYAL1 treated FbColl induced proliferation and invasiveness of melanoma cells.** Due to the limitations of FbECM, FbColl were utilized as an alternative 3D cell culture model to elucidate the impact of HA on melanoma cells behavior. To produce FbColl matrices, fibroblasts were embedded into 3D collagen matrices and cultured for 5 days, as shown in Fig. 4A. Analogous to FbECM, FbColl was decellularized, analyzed using HABP staining and cLSM microscopy (Fig. 4B) and characterized regarding HA amount using HA-ELISA kit. It was found that *Has2-KD* fibroblasts deposited 55% less HA than *Ctrl* fibroblasts (Fig. 4C), similarly to FbECM. The results con-

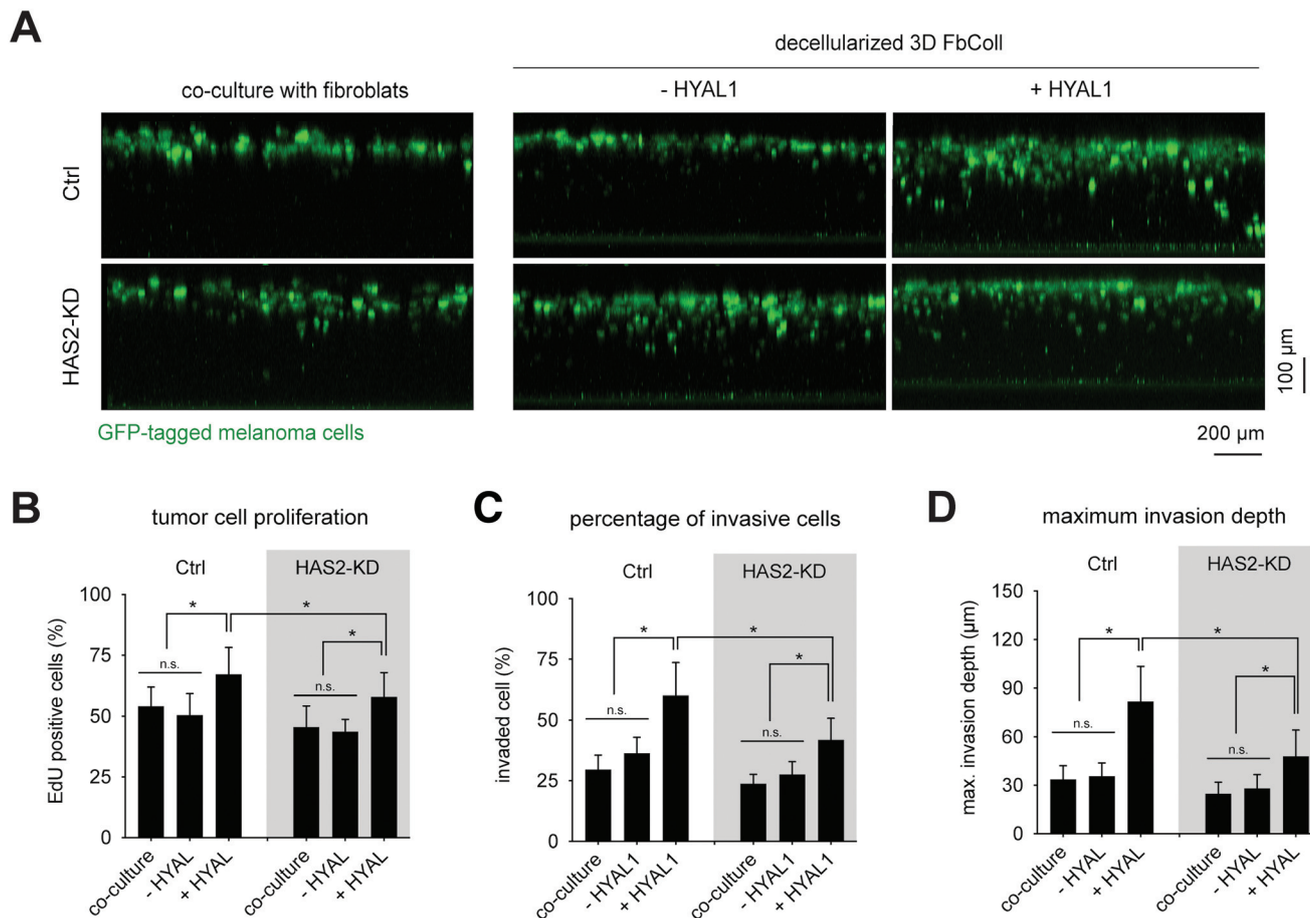
firmed the reduced HA deposition. Since the FbColl had a thickness of 350–450  $\mu\text{m}$ , it allowed quantifying the mechanical properties of FbColl in dependence of HA amount. As shown in Fig. 4D, FbColl from *Ctrl* fibroblasts (high HA deposition) exhibited a significantly higher matrix stiffness (170 Pa), comparing to collagen alone (80 Pa) and FbColl from *Has2-KD* (120 Pa). The stiffening of the matrix can be caused by HA deposition or cell-mediated matrix remodeling. To clarify this issue, an image-based quantitative porosity analysis was performed. As shown in Fig. 4E, the results revealed a reduction of matrix porosity by fibroblasts from 5  $\mu\text{m}$  (collagen alone) to approximately 3  $\mu\text{m}$  (FbColl). No significant differences in porosity were found between FbColl from *Ctrl* and from *Has2-KD*. The result points out that an increase of HA deposition by fibroblasts enhanced matrix stiffness, while it did not change the mean matrix porosity. This finding compared well with the *in vivo* results of increased skin stiffness at higher HA amounts.

Analogous to the FbECM, GFP-expressing B78D14 melanoma cells were cultured for 5 days on decellularized FbColl matrices with and without HYAL1 treatment. Additionally, non-decellularized matrices were used for co-culture with fibroblasts. Cells were analyzed regarding their invasiveness and proliferative capacity. Fig. 5A shows an *xz*-plot of B78D14 cells in FbColl matrices. Proliferating cells were quantified by counting EdU positive cells (Fig. 5B). Representative images of EdU assay are



**Fig. 4** Characterization of FbColl matrices. (A) Schematic illustration of the experimental workflow. (B) Representative images of FbColl reconstituted using *Ctrl* and *Has2-KD* fibroblasts. Collagen and HA were analyzed by cLSM using TAMRA-SE staining and HABP staining, respectively. (C) Quantitative analysis of HA amount in FbColl using commercial HA-ELISA kit. Quantitative analysis of (D) elastic modulus and (E) pore size of FbColl matrices (data are represented as mean  $\pm$  SD; \* – significance level of  $p < 0.05$ ).





**Fig. 5** HYAL1 treated FbECM matrices enhance proliferation and invasion of melanoma cells. (A) Representative images of *xz*-view of GFP expressing melanoma cells invasion into 3D FbColl matrices (B) and proliferation analysis of melanoma cells by means of EdU incorporation. EdU positive cells were analyzed throughout the matrix thickness by counting cells using home-built MATLAB script. (C) Quantitative analysis of percentage of invasive cells and (D) maximum invasive distance of melanoma cells. Cells found  $>20 \mu\text{m}$  beneath the matrix surface were counted as invasive cells. The maximum invasive distance was defined as the distance crossed by 10% of all cells (data are represented as mean  $\pm$  SD; \* – significance level of  $p < 0.05$ ).

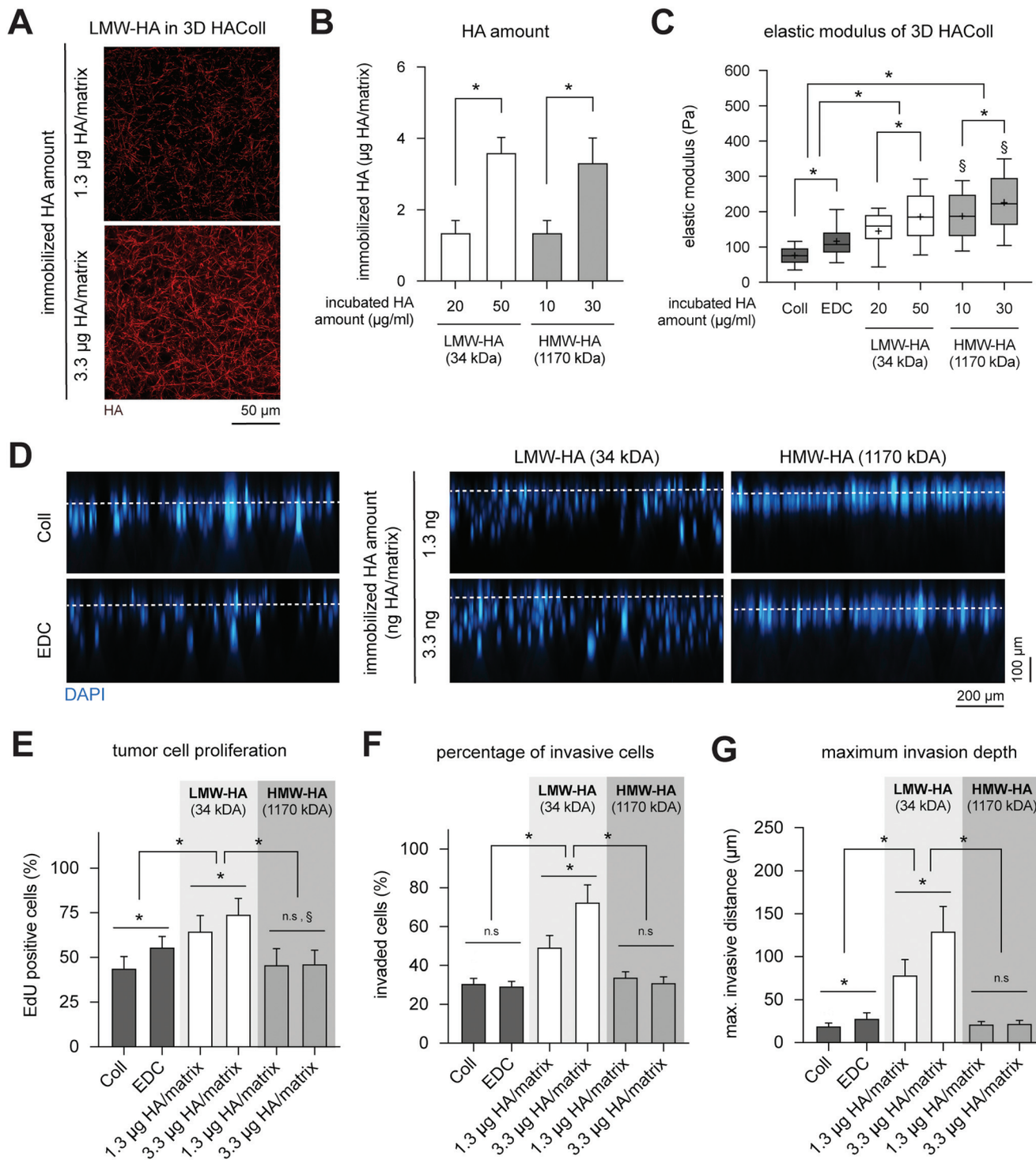
shown in ESI Fig. S6.† There was no difference between *Has2-KD* and *Ctrl* FbColl matrices, both in co-culture as well as in decellularized matrices. The only significant difference was an increased proliferation of melanoma cells due to HYAL1 treatment of matrices. This happened in both *Ctrl* and *Has2-KD* matrices. However, for *Ctrl*-HYAL1-treated matrices the effect was stronger than for *Has2-KD*-HYAL1-treated matrices. Again, these findings showed that reduced deposition of fibroblast-derived HA did not play a crucial role in melanoma cell proliferation. However, HA degradation did, even in a concentration dependent manner, since *Has2-KD* matrices had less HA incorporated and therefore less degraded HA. A similar influence was observed on the percentage of invaded cells and maximum invasion depth, where co-culture and/or reduced HA-deposition did not change melanoma cell behavior (30% invaded cells and maximum invasion distance of  $30 \mu\text{m}$ ), but HYAL1 treatment did (Fig. 5C and D). Here, an even higher impact of total amount of HA before HYAL-treatment was observed, as *Has2-KD*-HYAL1-treated matrices had both less

invading cells and smaller invasion distance (65% and  $90 \mu\text{m}$  for *Ctrl* compared to 45% and  $65 \mu\text{m}$  for *Has2-KD*).

In sum, HYAL1-treatment of matrices led to increased proliferation in both, FbECM and FbColl, accompanied by a decrease in cell invasion in the 3D FbColl matrices. Based on these findings we hypothesized that digested, small fragments of HA (in the range of  $<30 \text{ kDa}$ ) still remained in FbECM and FbColl, since HYAL1 degrade HA into HA fragments in the range of  $20 \text{ kDa}^{53}$  to  $66 \text{ kDa}^{54}$ . Furthermore, it has to be recognized, that the small HA fragments cannot be detected with the HA-ELISA kit used here or with HA-binding proteins.<sup>55</sup> Since *Ctrl* matrices contained higher amounts of HA than *Has2-KD* matrices, it can be assumed that more LMW-HA fragments were present after HYAL treatment, and in turn, could have led to an increased response of melanoma cells.

**3.2.3. HA-functionalized 3D collagen matrices (HAColl).** In order to investigate the hypothesis that the increased melanoma proliferation might be caused by remaining HYAL1-digested HA fragments, defined HA-functionalized 3D collagen





**Fig. 6** LMW-HA immobilized 3D collagen matrices modulate proliferation and invasion of melanoma cell in HA concentration dependent manner. (A) Representative image of LMW-HA immobilized 3D collagen matrices and the quantitative analysis of (B) HA immobilized amount (C) elastic modulus. (D) Representative images of xz-view of DAPI-stained melanoma cells invasion into 3D HAColl matrices. (E) Proliferation analysis of melanoma cells by means of EdU incorporation. EdU positive cells were analyzed throughout the full matrix thickness by counting cells using home-built MATLAB script. (F) Quantitative analysis of percentage of invasive cells and (G) maximum invasive distance of melanoma cells. Cells found >20 µm beneath the matrix surface were counted as invasive cells. The maximum invasive distance was defined as the distance crossed by 10% of all cells (data are represented as mean ± SD; \* – significance level of  $p < 0.05$ ).



matrices (Coll) were reconstituted. As discussed above, small HA fragments could not be detected with a conventional HA-ELISA kit. We therefore used ATTO565-labelled HMW-HA (1170 kDa) and LMW-HA (34 kDa) in the following experiment. Collagen matrices were post-functionalized by varying concentrations of HA, allowing adjustment of immobilized HA amount. Pore size of the matrices were in the range of 5  $\mu\text{m}$  and comparable to FbColl matrices. As shown in Fig. 6A, HA was analyzed using cLSM. The amount of HA was quantified using fluorimetry. Immobilized HA concentration of 1.3  $\mu\text{g}$  HA per matrix (0.02  $\mu\text{g}$  HA per  $\mu\text{g}$  Coll) and 3.3  $\mu\text{g}$  HA per matrix (0.41  $\mu\text{g}$  HA per  $\mu\text{g}$  Coll) for both LMW-HA and HMW-HA were generated (Fig. 6B) and used in these experiments. Even lower amounts of ATTO565-labeled HA could not be reproducibly bound to the collagen matrices and were therefore omitted. Matrix stiffness was analyzed and showed that an increase of HA amount enhanced matrix stiffness of both LMW-HA and HMW-HA (Fig. 6C). HA functionalized matrices exhibit higher stiffness than EDC-crosslinked matrices, similar to our previous report,<sup>11</sup> pointing out that HA contributes to matrix stiffening.

Culturing of B78D14 cells on HAColl matrices showed that melanoma cell proliferation and invasion is dependent on molecular weight of presented HA (Fig. 6E). Cell proliferation was found to be enhanced in a concentration-dependent manner for LMW-HA, but not for HMW-HA-functionalized matrices (Fig. 6B). Similar to cell proliferation, a higher invasion, both percentage of invaded cells and maximum invasion distance, was observed in a concentration-dependent manner for LMW-HA functionalized matrices (Fig. 6D, F and G). Again, HMW-HA did not affect invasion of melanoma cells, as previously reported for other human melanoma cell lines and *in vitro* models.<sup>11</sup> It has to be noted that controls of EDC cross-linking without HA immobilization showed a slight increase in proliferation and invasion in comparison to pure Coll, however, to a much lower extent as LMW-HA immobilization. This is in line with previous reports that matrix stiffness and LMW-HA, but not HMW-HA, co-jointly regulate proliferation of melanoma cells, as also reported in an animal model.<sup>56</sup>

In sum, only LMW-HA was able to increase proliferation and invasion of melanoma cells in a concentration-dependent manner, whereas HMW-HA did not affect both cell functions. This result, together with the effects seen in FbECM and FbColl matrices caused by HYAL1 treatment suggests that HMW-HA might have a homeostatic effect on the surrounding cells, whereas LMW-HA tend to trigger signal transduction in cells in terms of proliferation and invasion.<sup>57,58</sup>

### 3.3. Discussion: biomimetic models reveal alternative explanation for *in vivo* observation

An increase of HA, especially *via* overexpression of HAS2, showed tumor driving properties *via* different mechanisms.<sup>49,59–61</sup> Our findings fall in line with those reports, since the knockdown of *Has2* led to decreased primary tumor mass. It can be assumed that knockdown of the main HMW-HA producing enzyme *Has2* and only the smaller

amount of existing HMW-HA would result in smaller tumors. But taken the *in vitro* models into account (FbColl and HAColl), we could show that exposure to matrix-incorporated HMW-HA did not lead to increased proliferation or migration of B78D14 melanoma cells, even when increasing HMW-HA amounts.

This led to the question why decreased expression of HMW-HA *in vivo* would lead to reduced tumor weight. An alternative explanation could be that lower expression of HMW-HA is accompanied by less enzymatic catabolism to LMW-HA. Since some tumors are known to have high expression of HYALs<sup>62–64</sup> and B78D14 cells also produces a moderate amount of hyaluronidase (see ESI Fig. S7†), constant HYAL expression can lead to digestion of HMW-HA in the tumor microenvironment accompanied by activating effects on tumor cells. A reduction of HMW-HA amount, *e.g.* by *Has2* knockdown, would also decrease those activating effects, resulting in reduced tumor progression. However, this hypothesis could not be easily proved in an *in vivo* model because of its complexity. Furthermore, knocking down of genes in animal models can cause changes in expression of other ECM components *e.g.* versican and fibronectin,<sup>65</sup> which might affect tumor progression and development.<sup>66</sup>

Using biomimetic matrices, we finally could show that such an alternative explanation of an HA-dependent melanoma activation could be in place. We showed that both, HYAL1-treated matrices and LMW-HA functionalized matrices, enhanced proliferative and invasive capabilities of melanoma cells, which fit into the hypothesis of LMW-HA-induced tumor progression. Moreover, the results is well correlated to our previous report demonstrated that LMW-HA enhanced proliferation and invasion of BRO human melanoma cells, which was attributed to strong CD44-LMW-HA interactions.<sup>11</sup>

## 4. Conclusion

Overall our study found a correlation of *Has2*-knockdown dependent melanoma tumor growth *in vivo* and melanoma cell proliferation and invasion in three biomimetic matrix models of HA presentation of varied complexity. We demonstrated that the usage of engineered and well-defined cell culture models can help to better understand the complex behavior of melanoma growth *in vivo*. Moreover, our findings suggest that HA expression and its degradation by hyaluronidases in the tumor microenvironment trigger tumor progression.

## Conflicts of interest

The authors declare that there is no conflict of interest.

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