Enhanced Bioactivity of Titanium by Laser-generated Lotus-Topographies: Molecular Insights in Osteogenic Signaling Pathways of hASCs

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Abstract
Laser-generated lotus-topographies were shown to fulfill important requirements of demanded bioactivity: inhibition of biofilms and fibrotic capsule formation and enhanced regeneration. Here, we figure out how the demonstrated marker-specific improved osteogenic differentiation of human adipose-derived stem cells (hASCs) is accomplished. Comparing flat and lotus-structured titanium, the role of the osteogenic signaling pathways via extracellular-signal-related kinase (ERK), c-Jun-N-terminal kinase (JNK), and p38 mitogen-activated protein kinases (MAPKs), and interplay with ERK/p-ERK and bone morphogenetic protein (BMP2)/p-Smad signaling are determined. To our knowledge, this interaction has not been analyzed before. Blocking of ERK and JNK suppresses osteogenic markers like alkaline phosphatase (ALP) activity and calcium mineralization independently from the surface. Their functionality requires both transcription factors osteirx and Runx2/p-Runx2, and interplay with ERK/p-ERK and BMP2/p-Smad. P38 plays a regulatory role enabling osteogenic differentiation, which depends on the surface design. Its inhibition generally increases ERK/p-ERK cascade resulting in a high extent of Runx2/p-Runx2. Surface dependent BMP2/p-Smad and osterix are activated. The chief cause in enhanced osteogenic differentiation correlates with the improved attachment on the lotus-topography. Therefore, laser-generated lotus-structures present an attractive surface functionalization, finding their application in diverse biomedical applications.

Keywords: Bioactivity; Titanium; Surface topography; Adipose-derived stem cells; Osteoinduction

Introduction
Bioactivity can be seen as biological properties of implantable materials that go far beyond the well-known biocompatibility. Last includes non-toxic effects to the surrounding tissue and patient generally, which are evaluated following ISO 10993 before approving biomedical devices [1]. Unfortunately, long-time experiences in implantation medicine have demonstrated that biocompatibility is not sufficient to guaranty longevity and functionality of implants - causing harm to the patients and re-implantations with increasing health care costs. Implant failure can be caused by technical aspects such as fatigue, corrosion, and others [2]. But in the past it became apparent that biological problems as infections caused by biofilms and fibrotic capsule formation related to immune intractable. Up to now these biological problems could not be getting under control – and there is a long-life risk for the patients to suffer from them.

Coming back to term ‘bioactivity’, it means that the biomaterial is not passive; tissue-implant interactions do not occur randomly; the biomaterial can directly interfere with the tissue; it shall actively control tissue responses in the way it is needed. In the first place this control deals with an enhanced regeneration and the surrounding tissue [3]. This is classified by cell anchorage and differentiation. In the second place biofilm and fibrotic capsule formation shall be inhibited from the very beginning [4,5]. Therefore, a selective cell control is required [6]. Many efforts have been undertaken to test which material properties can enhance the bioactivity. The current strategy is based on changing the material surface [2,6-8]. Beside the technical challenge to modify it in a reproducible manner, the big question is what kind of surface characteristics are demanded for the certain application. For good outcome, the natural environment of cells should be mimicked. To form a concept, properties of the cell-surrounding extracellular matrix (ECM) should be copied. This ECM represents defined mechanical features, certain geometry, specific anchorage points for cell-binding and further storages signaling molecules like growth factors cells require for living [9]. These so-called functionalization approaches have a physical or biochemical nature. First is characterized by changes in mechanics, wettability, and charges as well as topological features. Last incorporates a surface design with cell anchorage points such as adhesion ligands or adhesive peptides and a controlled delivery of signaling molecules. In the past many studies have demonstrated that functionalization of biomaterial surfaces enhances the bioactivity [6,10-12]. Looking at the literature it can be criticized that the studies are difficult to be compared due to the non-standardized applied protocols and investigated cell type diversity. Most of the work was not concentrated on all important aspects of bioactivity – regeneration, infection plus immune response, and in vitro and in vivo assays were rarely performed in parallel. About 10 years of study on the functionality of laser-generated surface topographies, an enhanced bioactivity could widely be demonstrated by us: (a) influence on biofilm formation, (b) inhibition of fibrotic cells in vitro and in vivo, (c) improved cell anchorage and differentiation for neuronal and orthopedic applications [6,13-17]. Attractive technical aspects of laser-based biomaterial functionalization refer to the high reproducibility of defined surface designs in micro- and nano-scale without negative influence on the biocompatibility of the used material.

Concerning osteogenic differentiation, we observed that not all
typical osteogenic markers were increased by a lotus-topography in
titanium (Ti) [17]. This indicates that the osteogenic signaling pathways
differ for each marker and/or the topography can selectively interfere
on the molecular level. Both theories are not far-fetched looking at
the multiple, interconnected and still unknown possibilities how the
osteogenic transcription factors being responsible for the osteogenic
markers can become activated [18]. Since the role of signaling cascades
cannot be generalized due to cell specific differences, results can further
 correlate with the applied method [19], research is required to identify
the role of the surface functionalization. This research is absolutely
necessary to guaranty the effectiveness of the bioactivity. Above that
knowledge on the cell biological details of biomaterial-cell interactions
could facilitate future functionalization approaches.

Here, we are interested in the osteogenic molecular signaling
pathways in dependence of extracellular-signal related kinase (ERK),
c-Jun-N-terminal kinase (JNK), and p38 mitogen-activated protein
kinases (MAPKs). Additionally, the interplay with ERK/p-ERK and
bone morphogenetic protein (BMP2)/pSmad pathway is determined.

All of them were shown to be involved in osteogenic differentiation
[18]. Their impact on osteogenic differentiation of human adipose
derived stem cells (hASCs) is estimated via the osteogenic transcription
factors osterix, Runx2/p-Runx2 following the activation of osteogenic
markers such as alkaline phosphatase (ALP) activity and calcium
mineralization; and the impact on proliferation and actin cytoskeleton
with focal adhesion complex formation (via phosphorylated focal
adhesion kinase (p-FAK Tyr397)). The main focus is finding out the
influence on these cascades by the Ti lotus-topography.

Materials and Methods

Surface structuring

Lotus-topographies were generated on disk shaped titanium samples
(Goodfellow GmbH) with 6 mm in diameter and 0.25 mm in
thickness. For surface structuring, a commercially available amplified
Ti-Sapphire femtosecond laser system (Femtopower Compact Pro,
Femtolasers Produktions GmbH, Vienna, Austria) was used. It delivers
sub-30-fs pulses at 800 nm wavelength with energy of up to 1 mJ and a
repetition rate of 1 kHz. An achromatic lens with a focal distance of 200
mm was used for focusing the laser beam. The laser beam was scanned
along its linear polarization direction (x-direction). The scanning
speed in the x-direction was 800 µm/s and the translation step along
the y-direction was 15 µm. Lotus-topographies were generated by laser
ablation at a fluence of 8 J/cm² by using of circularly polarized laser
light [17]. For the cell studies, both Ti flat and lotus-topographies were
sterilized under UV light for 30 min.

Cell culture and experimental setup

All chemicals were purchased from Sigma-Aldrich (München,
Germany) unless otherwise noted. Human adipose-derived stem
cells (hASCs), used as passage 4-5, were purchased from Lonza
(Basel, Switzerland; Lot number 000039065). The original control
medium Dulbecco’s modified Eagles medium, including 10% foetal
bovine serum (Biochrom AG, Berlin, Germany) and antibiotics,
was additionally supplemented with 0.1 mM dexamethasone, 10
mM β-glycerolphosphate, and 50 µM ascorbate-2-phosphate
to induce osteogenic differentiation. To block the enzymes involved
in osteogenic differentiation, following inhibitors and concentration
were added to the osteogenic culture media: 20 µM ERK (U0126,
Biomol, Hamburg, Germany); 30 µM JNK (SP600125, Santa Cruz
Biotechnology, Heidelberg, Germany), 10 µM p38 (SB203580, Abcam,
Cambridge, UK). Since the inhibitors were prepared in dimethyl
sulfoxide (DMSO), a DMSO solvent control with a concentration
of 0.04% v/v in osteogenic media was additionally tested. The media
was changed every third day. For each experiment 1.8 × 10⁶ hASCs
per treatment was seeded out into 48 well plates filled with 300 µl cell
media culture. Staining analysis of the actin cytoskeleton, p-FAK Tyr397
as a component of focal adhesion complexes, the transcription factors
osterix, Runx2/p-Runx2, ERK/-p-ERK and BMP2/-p-Smad signaling was
performed after 3 days cultivation time, proliferation, ALP activity,
and calcium mineralization after 7, 14, and 21 days.

The osteogenic signaling cascades were analyzed on the control
surface (well plate bottom) first. For this purpose, undifferentiated
hASCs were seeded out in control medium, osteogenic medium, and
osteogenic medium supplemented with DMSO solvent and each
inhibitor, respectively. Since control hASCs did not undergo osteogenic
differentiation as shown for the markers ALP activity and calcium
mineralization and significantly reduced osteogenic transcription
factors (Supplementary Figure 1) [17,18], they were not used for the
further studies on the comparison of Ti flat and Ti lotus. Following the
staining protocols, the impact of p38 on osteogenic hASCs cultivated
on the control surface was additionally determined (Supplementary
Table 1), whereas the influences of ERK and JNK on the control surface
can be found in [18].

Imaging and staining protocols

Imaging was performed after 3 days cultivation time. To document
cell morphology and adhesion in dependence of the surface and
inhibitors, scanning electron microscopy (Microscope Quanta 400F
ESEM (FEI Company, Oregon, USA)) was used. After washing the
samples with phosphate buffer saline (PBS), they were fixed with 2.5%
glutaraldehyde for 15 min, followed by 2% osmium tetroxid for 30
min. After washing steps and a dehydration series with increasing ethanol
concentrations, the samples incubated in hexamethyldisilazane.
Finally, the samples were sputter-coated with gold and visualized [17].

For molecular insights, the cells were fixed in 4% paraformaldehyde
and permeabilized with 0.3% Triton X-100/PBS. To prevent nonspecific
antibody binding, a 2% bovine serum albumin/PBS solution was
incubated at 37°C for 2 h. Afterwards, the cells were kept in the primary
antibody solution, diluted in 0.3% Triton X-100/PBS, at 4°C overnight.
Following primary antibodies and concentrations were applied: p-FAK
(rabbit, 1:200; Cell Signaling, Danvers, USA), osterix (rabbit, 1:50, Santa
Cruz Biotechnology), Runx2 (goat, 1:50, Santa Cruz Biotechnology,
Heidelberg, Germany), p-Runx2 (rabbit, 1:100, Bios, Freiburg,
Germany), ERK (rabbit, 1:100, Cell Signaling), p-ERK (rabbit, 1:200,
Cell Signaling), BMP2 (goat, 1:50, Santa Cruz Biotechnology), p-Smad
1/5/8 (rabbit, 1:200, Cell Signaling). After several washing steps, the
fluorescence-conjugated secondary antibody was added at 37°C for 1 h
(Alexa Fluor® 488 conjugated Goat anti-rabbit IgG (H+L)/Alexa Fluor®
555 conjugated Goat anti-rabbit IgG (H+L), Diana, Hamburg,
Germany). Cell nuclei were stained with Hoechst 33342. For staining
of actin filaments, phallolidin-Atto 550 was added for 20 min. The cells
were kept in PBS for further imaging using a fluorescence microscope
(Nikon Eclipse, TE 2000-E, Nikon, Düsseldorf, Germany). All pictures
were taken with the same image recordings.

To quantify the relative fluorescence intensities, ImageJ software
was applied (http://rsweb.nih.gov/ij). For the transcription factors and
p-Smad, a rectangle of 2.5 × 10⁴ pixels was placed on the nuclei, for ERK,
p-ERK and BMP2 staining a rectangle with 10×103 pixels was placed
on the entire cell bodies. For each image the corresponding rectangle

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size was placed aside the cells to estimate the background signal. Using the task 'measure RGB', the color intensity could automatically be quantified. Thereafter, the corresponding background was subtracted. These relative results were averaged ± SEM and obtained from 100 rectangles per treatment [18].

**Proliferation and quantification of osteogenic differentiation**

Both analyses were performed with cell lysis, which were obtained after washing the samples with PBS and treatment with 1% Triton X-100/PBS [18]. The cell lysis was centrifuged at 12,000 g for 10 min, whereupon the supernatant was used to quantify the cell density for proliferation and osteogenic marker ALP activity. The pellet and bottom of the sample were incubated with 0.5 N HCl at 4°C overnight. Afterwards, both solutions for each sample were combined and utilized to measure calcium content as an additional osteogenic marker. To quantify the absorbance for each value, a microplate reader was used (Tecan Infinite M200Pro with i-controlTM software, Crailsheim, Germany). All results were obtained of three independent measurements after subtracting the blank and given as average ± SEM.

Following the online protocol of OPS Diagnostics (Lebanon, USA), the cell density was measured via lactate dehydrogenase (LDH) assay. The LDH activity was detected at 492 nm wavelength and correlated with a standard cell curve prepared under the same conditions. The results were expressed as cell number/cm².

The ALP activity was correlated with a standard curve of p-nitrophenol (10 mmol/L). Using 405 nm wavelengths, the ALP activity results of each sample were represented as nmol of p-nitrophenol/min/cm². Measuring calcium content followed the protocol of liquicolour assay (Human Diagnostics, Wiesbaden, Germany). Here, the standard calcium curve was raised from a stocking solution of 8 mg/dL calcium. Calcium was measured at 570 nm wavelength and given as µg/cm².

**Statistical analysis**

To analyze statistical differences, an unpaired two-sample Student’s t-test was applied with significant levels of p<0.05 (*), p<0.01 (**), and p<0.001 (***). First, DMSO solvent and applied inhibitors for ERK, JNK, and p38 were compared with the osteogenic setup (*); second, Ti flat with lotus-topography (#). The activity in vitro and in vivo [18,23]. In that connection it has to be pointed out that all analyzed osteogenic markers, their extent and duration, depend on the cell type, donor, and applied method [19]. Therefore, obtained results cannot be generalized.

Osteogenic markers of interest are ALP activity, calcium mineralization and specific osteogenic proteins such as osteocalcin or osteopontin [24]. The progress of ALP activity is characterized by an increase up to a maximum followed by a decrease. For hASCs this maximum occurred after 14 days cultivation time (Figure 4 and Supplementary Figure 1B). Since hASCs cultivated with control cell medium did not present these markers on control surfaces and functionalized Ti, they were not further analyzed (Supplementary Figure 1B and 1C) [17]. Concerning the enhanced osteogenic differentiation of hASCs on laser-generated lotus-topographies in Ti, we concluded that this improvement correlated with a control of cell adhesion mechanism on the lotus-structures: a different localization of integrin receptors occurred, hASCs attached significantly stronger, which was measured via adhesion forces, with a higher degree of vinculin as part of focal adhesion complexes [17]. Here, we additionally demonstrated that the cells were elongated on the substrates, stretching over many micro-sized surface features and forming diverse cellular extensions being in contact with the nano-roughness (Figure 1). Even though the cells were less expanded and reduced stress fibers on the lotus-structures compared with Ti flat, it supports the finding that the cells attached – since p-FAK was basically expressed at the end of the stress fibers, it indicates the formation of focal adhesions (Figure 2). For osteogenic cells higher cell densities on the control surface occurred, while on Ti flat and lotus it was comparable; therefore, no clear surface structure effect occurred (Figure 3 and Supplementary Figure 1A).

On the lotus-topography marker specific effects were estimated. For osteogenic hASCs ALP activity was significantly increased; but calcium values were similar (Figure 4). This is in contradiction to our first study, where the results were opposite [17]. Since different analyses were applied, it supports the fact that discrepancies can be raised from the method [18,19].

Generally, the mentioned osteogenic differentiation markers are activated by the transcription factors osterix and Runx2 [18]. Their transcriptional activation requires cofactors and phosphorylation, which in turn is initiated by diverse molecular signaling cascades and interplays [25]. In this connection a regulation via MAPK cascades (via JNK, ERK, and p38) and BMP-Smad pathway are suggested [18]. Comparing it with the control surface, it turned out that Ti flat and lotus significantly increased Runx2 and p-Ruxn2; osterix was also significantly increased on Ti flat, but similar values on Ti lotus were obtained (Figure 5 and Table 1B). Thereby, osterix and Runx2 were basically located within the nuclei, while p-Ruxn2 was further found

**Results and Discussion**

Many studies have demonstrated that a biochemical and physical functionalization of biomaterial surfaces enhances differentiation of stem cells [6,20-22]. This observation can be transferred to biomedicine in a way that such surface designs might improve tissue regeneration after implantation – one criterion of enhanced material bioactivity. An attractive surface modification refers to laser-structuring. It enables after implantation – one criterion of enhanced material bioactivity. An attractive surface modification refers to laser-structuring. It enables at least a good cell source in the past, due to the abundant supply after mesenchymal stem cells derived from adipose tissue (hASCs) became taken into account to classify osteogenic differentiation. Human biomaterial-cell interactions. signaling pathways, requires further cell biological analysis in terms of material information are transferred to cells and stimulate molecular material properties improve differentiation generally, how the specific of biofilm and fibrotic capsule formation, improvement of neuronal and nano-scale and enhances the demanded bioactivity: inhibition

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in the entire cell bodies after three days (Figure 5A and 5C). It can be concluded that both transcription factors are involved in osteogenic differentiation, which supports previous studies [18]. The higher extent might indicate an improvement of differentiation, whereas Runx2 signaling was more dominant on the lotus-topography. An increase of Runx2 expression was also documented for MSCs cultivated on surface-structured Ti and zirconia [26]. However, the activation mechanism of the transcription factor remained unclear.

Focusing on the interplay with ERK/p-ERK and BMP2/p-Smad signaling, surface specific influences for hASCs were estimated (Figures 6, 7 and Table 1B). ERK, p-ERK, BMP2, and p-Smad were located over the entire cell bodies, whereas p-Smad was dominantly presented within the nuclei after 3 days in culture. Control hASCs significantly reduced the amount for ERK, p-ERK, and p-Smad on the control surface, while the BMP2 value was comparable with the osteogenic cells (Table 1) [18]. Osteogenic hASCs significantly increased ERK/p-ERK on Ti flat; the activated form p-ERK on Ti lotus as well. This indicates that osteogenic differentiation Ti flat and lotus was dependent on interplay with this pathway. BMP2 values were either similar on Ti flat or significantly reduced on Ti lotus, while p-Smad was significantly increased on Ti flat or comparable with lotus topography. This can be interpreted in a way that the BMP2/p-Smad pathway is involved in osteogenic differentiation on Ti flat – but less important on Ti lotus. Generally, the role of BMP2/p-Smad is not solved for hASCs [18]. These findings indicate that this pathway correlates with the external cell environment, but further research is necessary to classify the interplay.

To get more insights in the MAPK cascade, ERK, JNK, and p38 were specifically blocked. Additionally, a DMSO solvent control was tested, which caused no remarkable impacts. JNK inhibition resulted in a changed cell morphology and poor focal adhesion formation; additionally cell proliferation was significantly reduced. ERK and p38 not clearly affected adhesion and morphology, while p38 significantly increased cell growth and ERK had no impact (Figures 1-3 and Supplementary Figure 1A) [18].

Starting with the osteogenic markers, specific effects occurred (Figure 4 and Table 1). Inhibition of JNK and ERK significantly reduced ALP activity (day 14) and calcium production (day 21) on all surfaces when compared with osteogenic hASCs, which supports the findings in [18]. The maximum of ALP activity was delayed to day 21, but still lower rates than for osteogenic cells were measured. The ALP values were lower than on the control surface, whereas the decrease was less on Ti lotus. On Ti flat calcium mineralization was higher for both inhibitors than on the control; on Ti lotus ERK blocking reduced it and JNK increased it. Thereby, the lotus-topography slightly supported ALP activity, but not calcium mineralization. It can be concluded that JNK and ERK are the main candidates to promote osteogenic differentiation independently from the surface. With respect to osteogenic hASCs, inhibition of p38 was more dependent on the surface: ALP activity was increased on Ti flat or comparable with lotus topography. This can be interpreted in a way that the BMP2/p-Smad pathway is involved in osteogenic differentiation on Ti flat – but less important on Ti lotus. Generally, the role of BMP2/p-Smad is not solved for hASCs [18].

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A

B

C

D

Figure 4: Quantification of the osteogenic markers at different time points of cultivation, comparing Ti flat (A, C) and lotus topography (B, D). (A, B) ALP activity via p-nitrophenol [nmol/min/cm²]; (C, D) calcium mineralization [µg/cm²]. The averaged results ± SEM were obtained of three independent measurements. An average cell density of 1.8 \times 10^4 cells per treatment was seeded out. Student’s-t-test was applied (p<0.05, p<0.01, p<0.001) to estimate significant differences between osteogenic and DMSO setup, and effects of the inhibitors of ERK, JNK, and p38 (*); and Ti flat with lotus topography (#).

component does not play a role in osteogenic differentiation – or its inhibition activates other signaling pathways that are linked to it in order to sustain lineage commitment. Other studies also assumed that p38 plays a negative role in osteoblast differentiation [27]. However, the significant differences between the surfaces indicate that the functionality of p38 strongly correlates with the external cell environment.

On the control surface, blocking of ERK and JNK significantly reduced osterix and Runx2/p-Runx2, when compared with osteogenic hASCs. This means that they require both transcription factors for osteogenic differentiation [18]. Inhibition of p38 also blocked osterix, but Runx2 values were comparable and p-Runx2 was even significantly increased. The impact on osterix was not a novelty, since p38 directly activates osterix [25]. It can be concluded that the sustained differentiation after p38 blocking only correlated with Runx2/p-Runx2 pathway. The presence of ERK and JNK inhibitors further suppressed osterix on Ti flat and lotus. On Ti flat Runx2 was not affected by ERK and JNK, but p-Runx2 was significantly increased after ERK and reduced after JNK treatment. On Ti lotus ERK reduced Runx2/p-Runx2, while p-Runx2 was significantly increased. Even though both inhibitors reduced the transcription factors on Ti less than on the control surface – except for osterix on Ti lotus, the suppression correlated with the observed inhibition of osteogenic markers. The sustained p-Runx2 expression after ERK treatment was not sufficient for osteogenic differentiation, indicating that osteogenic lineage commitment of these MAPKs requires both transcription factors. Similar to the osteogenic markers, the impact of p38 on transcription factors was very dependent on the external cell environment. On the control surface, osterix was significantly reduced, Runx2 values were similar and p-Runx2 was significantly increased in comparison to osteogenic hASCs.

On Ti flat osterix values were comparable, while Runx2/p-Runx2 was significantly increased. This positive outcome is in accordance with the enhanced extent of osteogenic markers. On Ti lotus osterix was significantly increased, Runx2/p-Runx2 was significantly increased. This positive outcome is in accordance with the enhanced extent of osteogenic markers. The improved osteogenic differentiation after p38 inhibition is rather dependent on Runx2/p-Runx2 pathway (Figure 5 and Table 1).

On Ti flat and lotus higher rates of p-ERK and p-Smad and lower rates for ERK and BMP2 were obtained after ERK and JNK treatment when compared with the control surface. But comparing it with osteogenic hASCs, blocking of ERK and JNK caused an inhibition of ERK/p-ERK and BMP2/p-Smad pathway independently from the
The presence of p38 inhibitors also suppressed BMP2/p-Smad in an analogous manner, but the extent of suppression was less on Ti flat compared to Ti lotus. In all cases, blocking of p38 caused a significant increase of ERK/p-ERK (Figures 6, 7 and Table 1)

This means that all three MAPKs crosstalk with these signaling cascades. Inhibition of ERK and JNK caused an inhibition of ERK/p-ERK and BMP2/p-Smad; inhibition of p38 inhibits BMP2/p-Smad, but activates ERK/p-ERK. This in turn has to correlate with the followed high extent of Runx2/p-Runx2. Even though Smads are formerly known to serve as a coactivator for Runx2 [18], their interaction after p38 blocking can be excluded on the control surface, since p-Smad values were reduced. This was not the case on Ti flat and lotus: higher rates of p-Smad and osterix occurred (Figures 5 and 7). However, the activated ERK/p-ERK cascade can explain why hASCs still undergo osteogenic differentiation after p38 blocking. In contrast to ERK and JNK, the role of p38 was strongly dependent on the surface design. Therefore, it can be concluded that p38 plays a regulatory role in osteogenic differentiation, being dependent on the external cell environment. The strong attachment to surface, as seen for the lotus-topography, supports it the best.

**Conclusion**

The molecular pathway of hASCs includes the MAPKs ERK, JNK, and p38 to stimulate osteogenic differentiation. They interact with ERK/p-ERK and BMP2/p-Smad signaling and activate the transcription factors osterix and Runx2/p-Runx2. ERK and JNK play a dominant role; their blocking abolishes all osteogenic markers. The functionality of p38 is regulatory and depends on the external cell environment. In this connection, the improved osteogenic differentiation on lotus-structures correlates with the enhanced cell attachment. This knowledge is essential to explain the improved bioactivity of laser-generated lotus structures – making this biomaterial functionalization very attractive for biomedical applications.

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Figure 5: Analysis of the transcription factors after 3 days cultivation time. Fluorescence images of (A) osterix (green), and (C) Runx2 (red) and p-Runx2 (green). Nuclei (blue) are stained via Hoechst. Scale bars= 50 µm. Quantification of the relative fluorescence intensities obtained from 100 cells per treatment, given as average ± SEM of (B) osterix, (D) Runx2, and (E) p-Runx2. Student’s-t-test was applied (p<0.05, p<0.01, p<0.001) to estimate significant differences between osteogenic and DMSO setup, and effects of the inhibitors of ERK, JNK, and p38 (*); and Ti flat with lotus topography (#).
Figure 6: Analysis of ERK/p-ERK signaling cascade. Fluorescence images of (A) ERK (green), (C) p-ERK (green). Nuclei (blue) are stained via Hoechst. Scale bars=50 µm. Quantification of the relative fluorescence intensities obtained from 100 cells per treatment, given as average ± SEM of (B) ERK, and (D) p-ERK. Student’s-t-test was applied (p<0.05, p<0.01, p<0.001) to estimate significant differences between osteogenic and DMSO setup, and effects of the inhibitors of ERK, JNK, and p38 (*); and Ti flat with lotus topography (#).

Figure 7: Analysis of BMP2/p-Smad signaling cascade. Fluorescence images of (A) BMP2 (red), (C) p-Smad 1/5/8 (green). Nuclei (blue) are stained via Hoechst. Scale bars= 50 µm. Quantification of the relative fluorescence intensities obtained from 100 cells per treatment, given as average ± SEM of (C) BMP2, and (D) p-Smad 1/5/8. Student’s-t-test was applied (p<0.05, p<0.01, p<0.001) to estimate significant differences between osteogenic and DMSO setup, and effects of the inhibitors of ERK, JNK, and p38 (*); and Ti flat with lotus topography (#).
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