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Combinatorial extracellular matrix cues with mechanical strain induce differential effects on myogenesis *in vitro*†

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Skeletal muscle regeneration remains a clinical unmet need for volumetric muscle loss and atrophy where muscle function cannot be restored to prior capacity. Current experimental approaches do not account for the complex microenvironmental factors that modulate myogenesis. In this study we developed a bio-mimetic tissue chip platform to systematically study the combined effects of the extracellular matrix (ECM) microenvironment and mechanical strain on myogenesis of murine myoblasts. Using stretchable tissue chips composed of collagen I (C), fibronectin (F) and laminin (L), as well as their combinations thereof, we tested the addition of mechanical strain regimens on myogenesis at the transcriptomic and translational levels. Our results show that ECMs have a significant effect on myotube formation in C2C12 murine myoblasts. Under static conditions, laminin substrates induced the longest myotubes, whereas fibronectin produced the widest myotubes. Combinatorial ECMs showed non-intuitive effects on myotube formation. Genome-wide analysis revealed the upregulation in actin cytoskeletal related genes that are suggestive of myogenesis. When mechanical strain was introduced to C + F + L combinatorial ECM substrates in the form of constant or intermittent uniaxial strain at low (5%) and high (15%) levels, we observed synergistic enhancements in myotube width, along with transcriptomic upregulation in myosin heavy chain genes. Together, these studies highlight the complex role of microenvironmental factors such as ECM interactions and strain on myotube formation and the underlying signaling pathways.

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1. Introduction

Skeletal muscle plays an important role in bearing weight and controlling movement of the body. Although skeletal muscle has high regenerative capacity, the ability to restore full muscle function is permanently impaired by traumatic injuries like volumetric muscle loss.¹ Clinical treatments to muscle injuries include the grafting of autologous muscle flaps or scar tissue debridement.^{2–4} However, these therapies are often associated with significant donor site morbidity due to impaired endogenous regeneration and revascularization capacity. Current experimental approaches to induce muscle

regeneration after traumatic injuries include tissue engineering, biological scaffolds, cell transplantation, rehabilitative exercise, and pro-regenerative soluble factors,^{5–12} although clinical translation is lagging.

Muscle regeneration is a complex process in which satellite cells activate into myogenic precursors that proliferate, followed by cell cycle withdrawal and myoblast fusion to form multi-nucleated myotubes.¹³ This process is orchestrated by numerous signaling pathways, including insulin-like growth factor (IGF-1), notch, prostaglandin E2, and peroxisome proliferator activated receptor gamma (PPARG).^{14–16} It is well-recognized that the skeletal muscle microenvironment plays a critical role in skeletal muscle regeneration. These microenvironmental factors include the extracellular matrix (ECM) protein composition, biophysical patterning, and substrate stiffness.¹⁷ The ECM is important to normal muscle function. Skeletal muscle ECM is composed of a combination of proteins, including laminin, collagen I, collagen I, elastin and proteoglycans.¹⁸ Therefore, combinatorial ECMs may better reflect the native muscle ECM environment. Since these factors interact directly with muscle precursor cells, which give rise to multi-nucleated myofibers, it is generally agreed that these microen-

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environmental factors can modulate myogenesis.¹⁹ Additionally, owing to the dynamic nature of skeletal muscle, studying myogenesis in the presence of strain can reflect rehabilitative strategies to promote muscle regeneration. Rehabilitative exercise has previously been shown to improve muscle regeneration in preclinical models of muscle injury.²⁰ Similarly, strain effects on myogenesis have been studied *in vitro*^{21,22} in which it was shown that strain mediated alignment of collagen I fibers enhanced the myogenesis of myoblasts, thereby recapitulating *in vivo* microenvironment.²³

To study muscle myogenesis in an improved biomimetic microenvironment, we developed a stretchable tissue chip technology platform composed of customized combinatorial ECMs and strain regimens. Using this platform, we tested the effects of both intermittent and constant strain regimens at both low and high strain magnitude, in conjunction with combinatorial ECMs, on *in vitro* myogenesis. Our *in vitro* and transcriptomic results reveal complex interactions between ECM components and strain regimens on the myogenesis process.

2. Experimental

2.1. Cell culture

Murine myoblasts (C2C12, passages 4-7, ATCC) were cultured in Dulbecco's modified Eagle's medium (DMEM) supplemented with 20% fetal bovine serum and 1% penicillin/streptomycin at 37 °C in a humidified atmosphere containing 5% CO₂. Cells were sub-cultured when they reached 80% confluency.

2.2. ECM surface functionalization of stretchable tissue chips

24-Well uniaxial stretch chambers with aligned nanopatterns in the direction of strain (Curi Bio, USA) were disinfected with 70% ethanol and then modified with poly D-lysine (0.1 mg mL⁻¹) for 16 hours at 37 °C to render them surface-reactive. Next, collagen I (C, Corning), fibronectin (F, Sigma) laminin (L, Life Technologies), and their combinations thereof were incubated at a total concentration of 10 µg mL⁻¹ for 16 hours at 37 °C at equimolar ratios.

2.3. Cell seeding

Murine myoblasts were seeded into 24-well stretch chambers having aligned nanopatterns parallel to the direction of strain (Curi Bio, USA) at density of 3×10^4 cells per cm². Cells were grown to confluency before induction of myogenesis by switching to differentiation medium (DMEM with 2% horse serum and 1% penicillin/streptomycin) and simultaneously initiating mechanical stimulation.

2.4. Mechanical stimulation

Uniaxial strain stimulation was applied to the tissue chips using the Cytostretcher (Curi Bio) as shown in Fig. 1A. The constant strain regime consisted of linearly applied low (5%) or high (15%) strain at 0.1 Hz. The intermittent strain regime

consisted of repeating cycles of stimulation (10 minutes at 0.1 Hz followed by 20 minutes of rest). As a basis for comparison, the control samples were cultured under normal static conditions. Mechanical stimulation was applied for a total of 4 days, in which the differentiation media was changed on day 2 of stimulation.

2.5. Immunofluorescence staining of myotube formation

After 4 days of differentiation, the cells were fixed with formalin (10%) for 15 minutes at room temperature and immunofluorescently stained for myosin heavy chain (Sigma), followed by incubation with secondary antibody conjugated with Alexa Fluor 594 (Fisher Scientific) and counter staining for nuclei with Hoechst 33342 (Fisher Scientific). Fluorescence images were captured at 10× magnification with an inverted confocal microscope (Zeiss LSM 710). For each independent experiment ($n = 3$), 3 images were acquired for image analysis of myotube formation.

2.6. Image analysis

For *in vitro* studies, the lengths and widths of myotubes were measured from 3 images acquired for each independent experiment in each treatment group. Myotube length and widths were measured manually using ImageJ. Only myotube fully visible within the images were quantified.

2.7. RNA isolation

A subset of samples was processed for RNA isolation to study the effects of combinatorial ECMs and strain on transcriptional modification. Medium-sized stretch chambers (12 × 12 mm, Curi Bio) with parallel-aligned microgrooves were functionalized with poly D-lysine (0.1 mg mL⁻¹) for 16 hours before coating of collagen I, laminin, or C + L + F as described in section 2.2. Cell seeding was performed according to section 2.3. Upon reaching 90% confluency, these cell-seeded chambers underwent constant high strain (15%) in differentiation media on the Cytostretcher. After 4 days of strain stimulation, the cells were lysed and the RNA was extracted using GeneJET RNA isolation kit (Fisher Scientific).

2.8. RNA sequencing

RNA sequencing was performed by Novogene Corporation. In brief, messenger RNA was purified from total RNA using poly-T oligo-attached magnetic beads. After fragmentation, the first strand cDNA was synthesized using random hexamer primers, followed by the second strand cDNA synthesis using either dUTP for directional library or dTTP for non-directional library. The library was checked with Qubit and real-time PCR for quantification and bioanalyzer for size distribution detection. Quantified libraries will be pooled and sequenced on Illumina platforms, according to effective library concentration and data amount.

Original image data file from high-throughput Illumina sequencing platform was transformed to sequenced reads (Raw Data) by CASAVA base recognition (Base Calling). Raw data are stored in FASTQ (fq) format files which contain



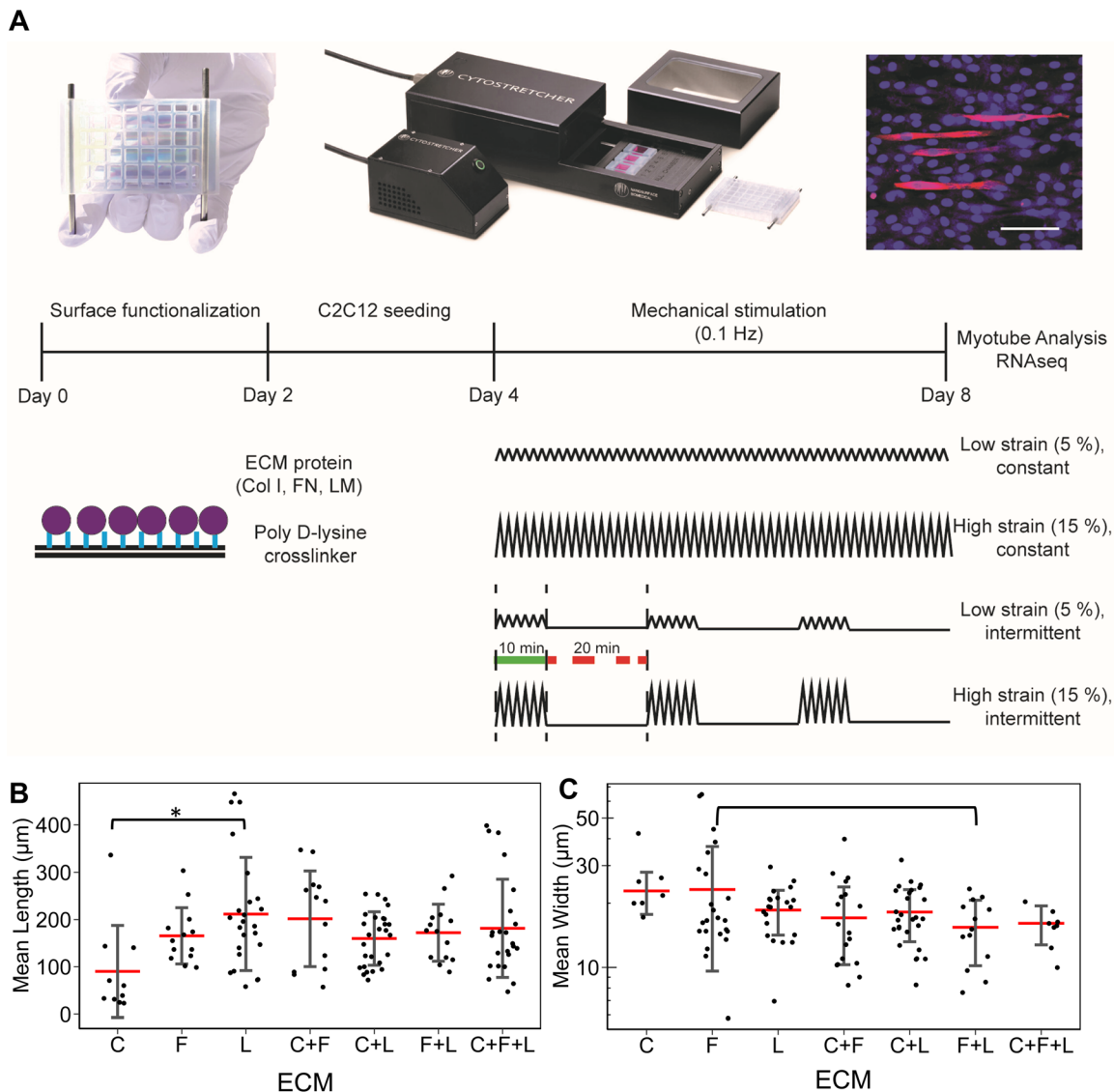


Fig. 1 Schematic and quantification of myotube length and width on ECM combinations under static conditions. **A**. Schematic of the experimental setup and representative fluorescent image of myosin heavy chain (red) immunostaining in C2C12 myoblasts cultured on different ECM combinations. Total nuclei were counterstained using Hoechst 3342. Scale bar: 100 μm . **B** and **C**. Quantification of mean myotube length and width respectively. Each dot represents individual myotube values from 3 independent experiments. Mean values are represented by the red line. Standard deviation is represented using the error bars. Statistical analysis was performed using ANOVA with *post hoc* Tukey test for individual comparisons. * denotes statistically significant comparisons ($P < 0.05$).

sequences of reads and corresponding base quality. Raw data was filtered by removing reads with adapter contamination, with uncertain nucleotides constituting more than 10 per cent of either read ($n > 10\%$) and with low-quality nucleotides (base quality less than 5) constituting more than 50 per cent of the read. This was followed by alignment of reads using HISAT2 to the reference with the following considerations: (1) the whole sequence was aligned to a single exon, (2) the sequencing sequence was piecewise aligned to two exons of the genome, and (3) the sequencing sequence was segmented and aligned to more than three (including three) exons of the genome, and (4) the fragments per kilobase of transcript sequence per

millions base pairs sequenced (FPKM) was quantified for each aligned read to estimate gene expression levels, which takes the effects into consideration of both sequencing depth and gene length on counting of fragments.

For RNAseq data analysis, the FPKM counts were normalized, and log transformed. Surrogate variable analysis was done using RUVseq package in R, and removeBatchEffect function was used to remove the surrogate variables found. Differential accessibility analysis was done using lmFit function of package limma in R. Specific comparisons (strain/ECM) were statistically tested in the model using contrasts.fit function. Adjusted P value (P_{adj}) and fold change was calcu-



lated for each region and comparison. Specific differential gene expression lists were made for specific comparisons to enable enrichment analysis on EnrichR website. For gene set enrichment analysis (GSEA) analysis, the java desktop application was used, with instructions from the Wiki page was utilized. The bar plots were made using ggplot package and pheatmap package was used to create all the heatmaps in the figures.

2.9 Quantitative PCR (qPCR)

First, reverse transcription on isolated RNA sample was performed using the Superscript II reverse transcriptase (Invitrogen) and cDNA was synthesized according to the manufacturer's instruction and previous publications.²⁴ Primers used for Taqman qPCR consisted of MyoD, Myh1, Myh3 and GAPDH (all from Applied Biosystems, Foster City, CA). The qPCR was performed on a 7900 real-time PCR system (Applied Biosystems) for 40 cycles. The data were analyzed and processed by the $\Delta\Delta C_t$ method, normalized to GAPDH housekeeping gene, and then quantified as normalized relative fold changes with respect to collagen I under static conditions ($n = 2-3$).

2.10 Statistical analysis

For *in vitro* studies, the lengths and widths of myotubes from 3 independent experiments were shown as mean \pm standard deviation. Statistical analysis was performed by analysis of variance with Tukey *post hoc* test, where statistical significance is accepted at $P < 0.05$. For RNA sequencing, a linear model was computed for the gene expression as function of the ECM and mechanical stimulus (3 biological replicate each). The differences between each treatment group were calculated using established methods in the contrasts.fit function in limma package R. The statistics for the differential gene expression is calculated using eBayes functions in which analysis of variance (ANOVA) with Tukey's *post hoc* test was performed for all comparisons. Statistical significance was accepted at $P_{adj} < 0.05$.

3. Results

3.1 Combinatorial ECMs under static conditions differentially modulate myotube length and width

To study the effects of combinatorial ECMs on myogenesis, we first examined myotube formation over 4 days on stretch chambers that were surface-modified by collagen I (C), fibronectin (F), laminin (L); their two-factor combinations (C + F, C + L, F + L); and their three-factor combination (C + F + L) (Fig. 1A and ESI Fig. 1†). The average myotube length and width using myosin heavy chain immunostaining was quantified. These morphological parameters were selected based on prior publications that establish a correlation between morphological features and myotube formation.^{25,26} We first characterized the myogenesis of myoblasts in the presence of combinatorial microenvironments. Laminin exhibited the

highest average myotube length ($211 \pm 119 \mu\text{m}$) among all ECM compositions, reflecting >2 -fold higher length compared to collagen I ($90 \pm 97 \mu\text{m}$, $P = 0.008$) as represented in bar plot in Fig. 1B. However, myoblasts cultured on laminin showed no significant difference in myotube width compared to other ECMs (Fig. 1C). Interestingly, fibronectin + laminin (F + L) resulted in lowest average myotube width in myoblasts ($15 \pm 5.2 \mu\text{m}$), especially compared to fibronectin alone, which supported the highest myotube width compared to all other ECM combinations ($23 \pm 13 \mu\text{m}$, $P = 0.06$). This non-intuitive finding may be a result of inhibitory effects, as has been observed in combinatorial ECM interactions in other cellular contexts.^{27,28} It was also observed that different ECMs resulted in highest myotube length and width, pointing towards different molecular mechanisms being affected by different ECM combinations.

3.2 RNA sequencing analysis reveals combinatorial ECM effects on myogenic differentiation of under static condition

To understand regulatory mechanisms underlying differential effect of ECM combinations on average myotube length and width, we performed bulk RNA sequencing of myoblasts that were cultured on a subset of the ECM conditions. To study the molecular underpinnings in the presence of combinatorial vs. single-factor ECMs, we selected the three-factor ECM (C + F + L) combination, compared to the single-factor collagen I (C) and laminin (L) constituents that have distinct effects on myogenesis. Collagen I significantly exhibited upregulation of ECM related genes, specifically those relating to collagens, fibronectins, laminins, and integrins, including *Col1a1*, *Col1a2*, *Itgb1*, *Itgb5* and *(Fig. 2A). This result suggests a higher ECM production in myoblasts on collagen I, compared to that on laminin or C + F + L under static conditions. Furthermore, gene enrichment analysis revealed the upregulation in the categories associated with ECM organization (*Col1a1*, *Mfap5* (a glycoprotein in myofibrils in ECM), *Itga11*) and muscle contraction (*Myl1*, *Tnnt1*, *Tcap* (sarcomere assembly)) gene sets on collagen I compared to laminin (Fig. 2C). In agreement with higher myogenesis on collagen I, the gene set enrichment analysis showed downregulation of categories associated with cell cycle, compared to laminin, as withdrawal from cell cycle is required for myogenesis (Fig. 2D).²⁹ Since prostaglandin E2, notch, and PPARG (peroxisome proliferator activated receptor gamma) are also important myogenesis pathway genes, we observed that collagen I shows significant upregulation in a subset of genes from these pathways such as *Pparg*, *Ppargc1a*, *Igf1*, *Ptgis*, *Jag1* compared to C + F + L under static conditions.³⁰⁻³³*

To substantiate the *in vitro* finding that laminin supported the formation of the longer myotubes, compared to collagen (Fig. 1B), we show that laminin substrates upregulated multiple genes in the prostaglandin E2, notch signaling and *Pparg* signaling pathways, signifying a unique muscle regeneration signature associated with laminin's role in modulating myogenesis (Fig. 2B). However, it is important to note that a different subset of myogenesis-related genes were upregulated



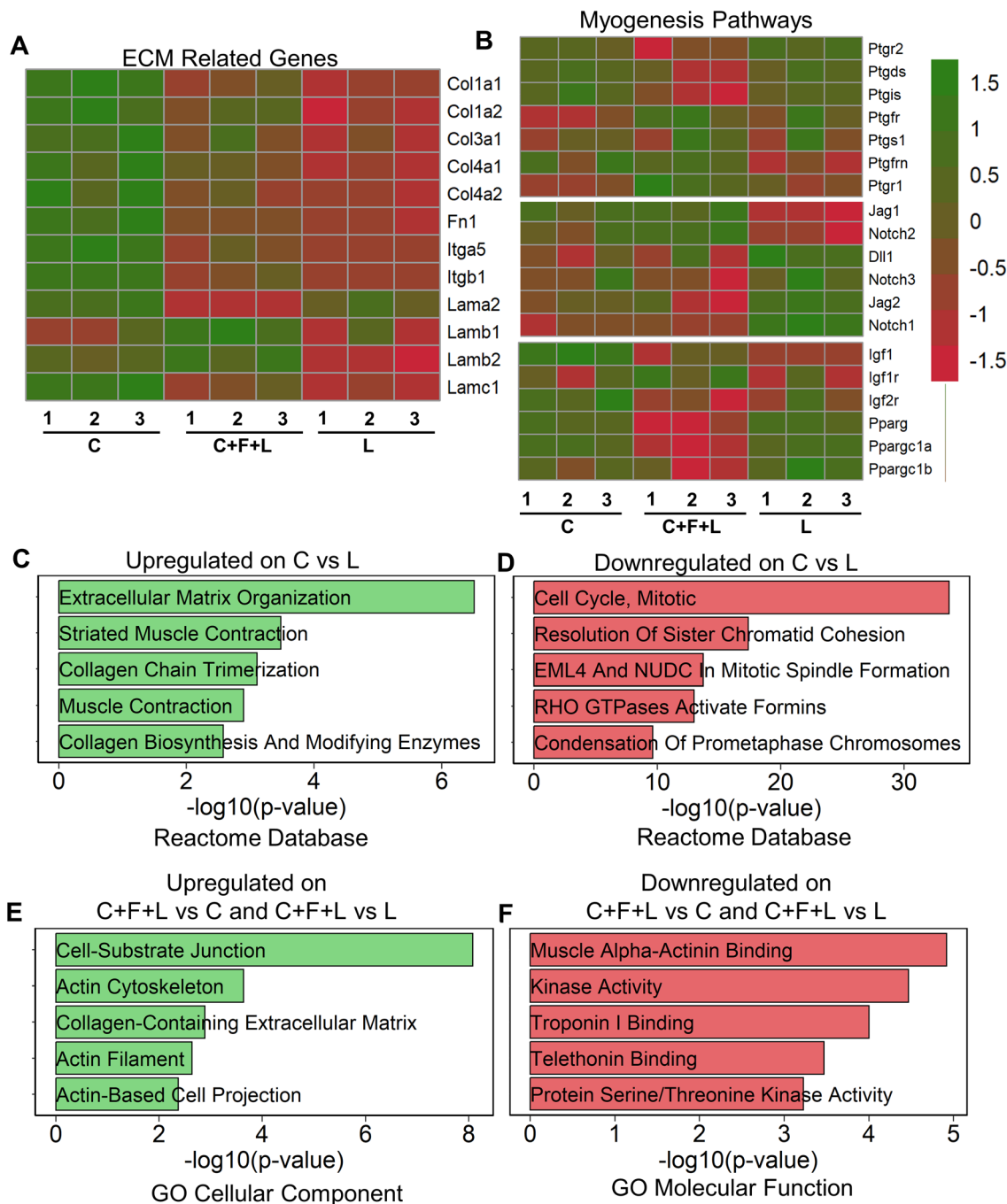


Fig. 2 Transcriptomic analysis of myoblasts cultured on ECM combinations under static conditions. A and B. Heatmap of differentially expressed ECM-related genes and myogenesis pathway genes, respectively. C. Enriched gene set categories upregulated in myoblasts cultured on collagen I (C), compared to laminin (L), based on Reactome database of genes. D. Enriched gene set categories downregulated in myoblasts cultured on collagen I, compared to laminin, based on Reactome database of genes. E. Enriched gene set categories upregulated in myoblasts cultured on C + F + L, compared to collagen I or laminin, based on GO cellular component database. F. Enriched gene set categories downregulated in myoblasts cultured on C + F + L, compared to collagen I or laminin, based on GO molecular function database.

on laminin substrates compared to both collagen I and C + F + L under static conditions. The C + F + L ECM combination specifically exhibited significantly higher expression of laminin b1 and b2 compared to both collagen I and laminin, while also demonstrated an overall higher expression of ECM

related genes compared to only laminin, including *Col1a1*, *Col1a2*, and *Fn1* (Fig. 2A).

Upregulated genes on C + F + L ECM combination, compared to either laminin or collagen I, included the categories of actin cytoskeleton and cell-substrate junction (*Actn1*, *Fnlb*,



Vcam1) gene sets from the GO cellular component database (Fig. 2E). This finding suggests that the C + F + L ECM combination supported more interaction with the underlying substrate, compared to laminin and collagen I alone. Downregulated genes on C + F + L, compared to collagen I or laminin corresponded to muscle alpha-actinin binding, tropomyosin binding pointing towards decreased expression of muscle contraction and assembly genes (Fig. 2F). Additionally, myogenesis pathway genes such as *Ptgs1*, *Igfbp1* and notch signaling receptor *Notch2* were specifically upregulated on C + F + L (Fig. 2B). Together, these data suggested that single-component and combinatorial ECMs had a unique molecular signature and effect in modulating myogenesis.

Since sarcomeric structure is an important functional characteristic of myotube function, we quantified genes from sarcomere geneset in the gene ontology database in our transcriptomic analysis (ESI Fig. 2†). We discovered three distinct gene profiles, each enriched in a specific ECM combination. Specifically, genes in cluster A were upregulated in C + F + L combination under static conditions and corresponded to gene such as *Calml1*, *Calml3* and *Actn1*, which are an integral part of the protein subunit for calcium signaling in sarcomeres.³⁴ Additionally, cluster B corresponded to genes upregulated on either single-factor ECMs under static condition, containing genes such *Ttn* and *Tnnt1*, which are responsible for muscle contraction in sarcomeres.³⁵ These findings suggest that sarcomeric genes were distinctively different on the C + F + L combination, compared to the single factor components.

3.3 Combined effects of strain with C + F + L on myogenesis

Having studied the effect of ECMs on the myogenic differentiation of C2C12 cells under static conditions, we next sought to understand the combined role of ECM composition and strain on the myogenic differentiation. Two levels of uniaxial strain (high or low, Fig. 1A) were applied to the cells cultured on all ECM combinations. Furthermore, these two levels of uniaxial strain were applied either constantly or intermittently. It was observed that high constant strain to myoblasts on C + F + L ECM combination resulted in the highest average myotube length, which was significantly higher compared to other strain regimens (Fig. 3A). Specifically, high constant strain on C2C12 myoblasts on C + F + L ECM exhibited a 50% increase compared to both high intermittent strain ($P = 0.003$) and low constant strain ($P = 0.0005$), and a 34% increase compared to low intermittent strain ($P = 0.03$; Fig. 3A). On the other hand, the highest myotube width was observed with high intermittent strain for C2C12 cells cultured on C + F + L. Specifically, an increase of 56% in myotube width was observed with high intermittent strain, compared to the static condition ($P = 0.006$; Fig. 3B). These findings show that high constant strain increased the average myotube length whereas high intermittent strain significantly increased average myotube width.

Compared to the combinatorial environment of C + F + L, the individual components alone did not have a significant difference in both the myotube length and width with the high constant strain application, with the exception of a signifi-

cantly lower myotube length on collagen I compared to C + F + L ECM (Fig. 3C and D). Genome-wide analysis of myoblasts cultured on C + F + L ECM with and without high constant strain demonstrated higher expression of multiple myosin heavy chains with the application of strain. In particular, *Myh4* (slow), *Myh1* (fast), *Myh2* (fast) and *Myh8* (fetal muscle) were found to be significantly upregulated with the strain on the C + F + L ECM (Fig. 3F). Additionally, upregulated genes C + F + L ECM with high constant strain compared to static conditions were enriched in myosin related protein domains (*Myo1e*, *Myo5a*) in the Pfam InterPro domains database (Fig. 3E). It was also observed that multiple muscle regeneration related growth factors and pathways such *Igfbp1*, *Pparg*, notch signaling was also upregulated with the application of strain on C + F + L (Fig. 3G). Overall, strain application to the differentiating myoblasts cultured on the C + F + L ECM combination enhanced both myotube width and length by upregulating certain myosin heavy chain expression and muscle-regeneration related pathways such as IGF-1 and notch.

3.4 Combined effects of strain and laminin on myogenesis

Next, we evaluated the effects of strain on laminin substrates. On laminin alone, a significantly higher myotube length was observed under high constant strain and low intermittent strain ($P < 0.01$), but not under low constant strain or high intermittent strain (Fig. 4A and ESI Fig. 3†). We also observed that all strain regimens increased the average width of myotubes, compared to on static conditions ($P < 0.05$, Fig. 4B). Interestingly, combination of C + L did not significantly affect the myotube width and length with application of high constant strain (Fig. 4C), even though under static conditions a significant decrease was observed in the myotube length with on C + L ($P = 0.03$) (Fig. 1A). A significant decrease in both myotube length and width was observed with pairing F + L under high constant strain condition, which was also observed under static conditions (Fig. 1A). Interestingly, this decrease in myotube length and width is not observed with the C + F + L ECM combinations, suggesting a non-intuitive influence of F + L compared to C + F + L on myogenic differentiation of myoblasts (Fig. 4C and D).

At the transcriptional level, the expression of skeletal myosin heavy chain-associated genes was upregulated with the application of high constant strain on laminin. These genes included isoforms *Myh3* (embryonic, associated with regenerating muscle), *Myh1* (type IIa, fast) and *Myh2* (type IIb, fast), signifying increased muscle functionality of differentiating myoblasts cultured on laminin with high constant strain, compared to on static conditions (Fig. 4E). Furthermore, gene set enrichment analysis revealed significantly higher enrichment in the category of skeletal myofibril assembly in myoblasts on laminin with high constant strain, compared to static conditions (enrichment score 0.7; $P = 0.048$) (Fig. 4F). Some of the upregulated genes in this category included cytoskeletal markers such as *Tcap*, *Ttn*, and *Tmod1*.

Additionally, we validated gene expression findings from RNAseq using quantitative PCR (ESI Fig. 4†). Similar to the



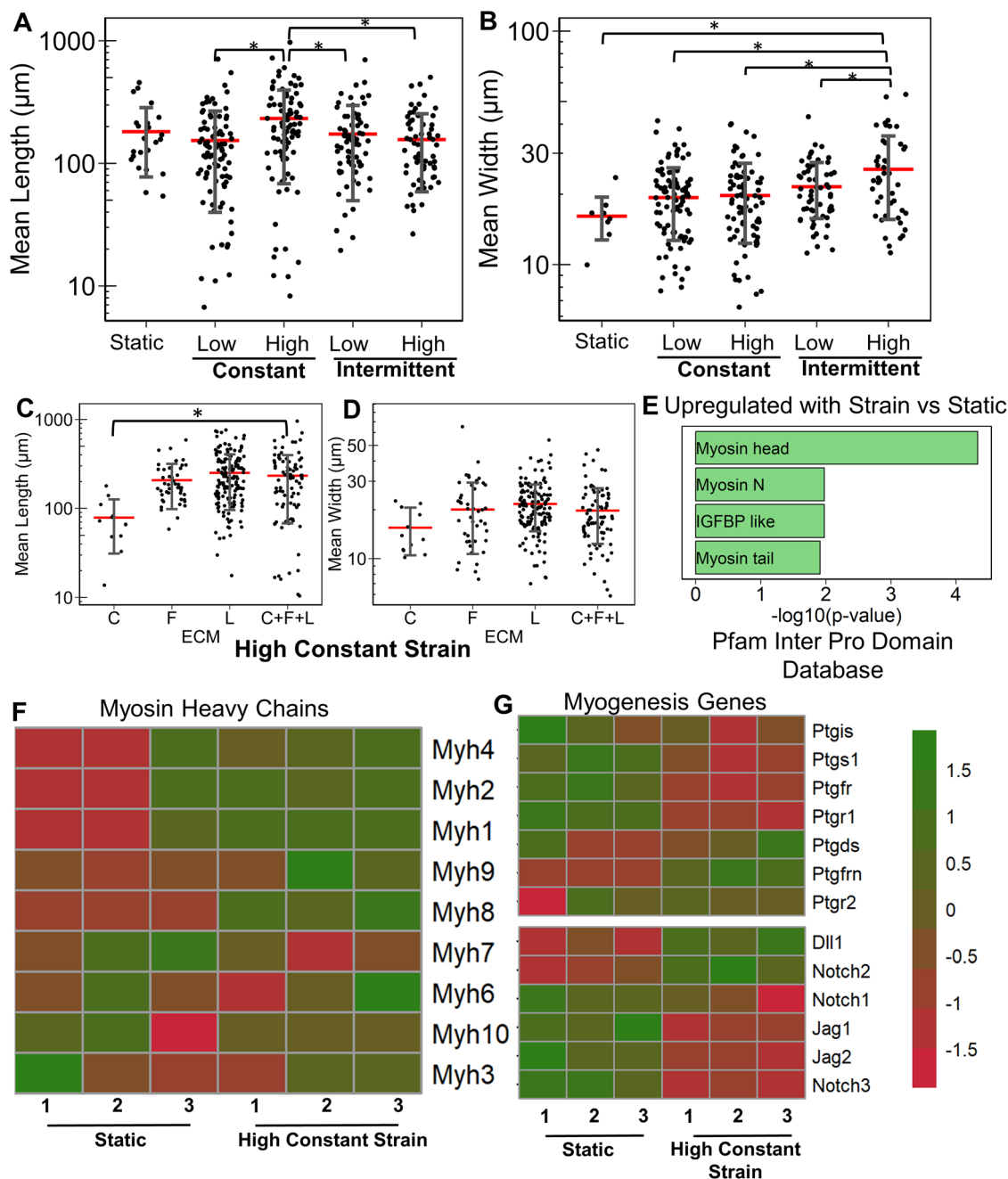


Fig. 3 Effect of uniaxial strain regimens on myogenesis cultured on C + F + L ECM substrates. A and B. Quantification of mean myotube length (A) and width (B) with the application of different strain regimens in differentiating myoblasts cultured on collagen I + fibronectin + laminin (C + F + L) substrates. C and D. Quantification of mean myotube length (C) and width (D) with the application high constant strain on C + F + L, collagen I (C), fibronectin (F), or laminin (L) substrates. Each dot represents individual myotube values from 3 independent experiments. Mean values are represented by the red line. Standard deviation is represented using the error bars. Statistical analysis was performed using ANOVA with *post hoc* Tukey test for individual comparisons. * denotes statistically significant comparisons ($P < 0.05$). E. Enriched gene set categories upregulated in differentiating myoblasts cultured on C + F + L with high constant strain, compared, to static conditions, based on the Pfam Inter Pro database. F and G. Heatmap of differentially expressed myosin heavy chain genes and myogenesis pathway genes respectively in myoblasts cultured on C + F + L with high constant strain, compared to static conditions.

RNAseq, upregulation of two myosin heavy chains, *Myh1* and *Myh3*, was observed on laminin in the presence of high constant strain. Additionally, an early myogenesis marker, *MyoD*, was found to be upregulated on laminin in the presence of

strain, signifying the synergistic effect of mechanical stimulation and combinatorial ECM combination on early myogenesis. Together, these findings suggest that C2C12 myoblasts cultured on laminin and under specific strain regime



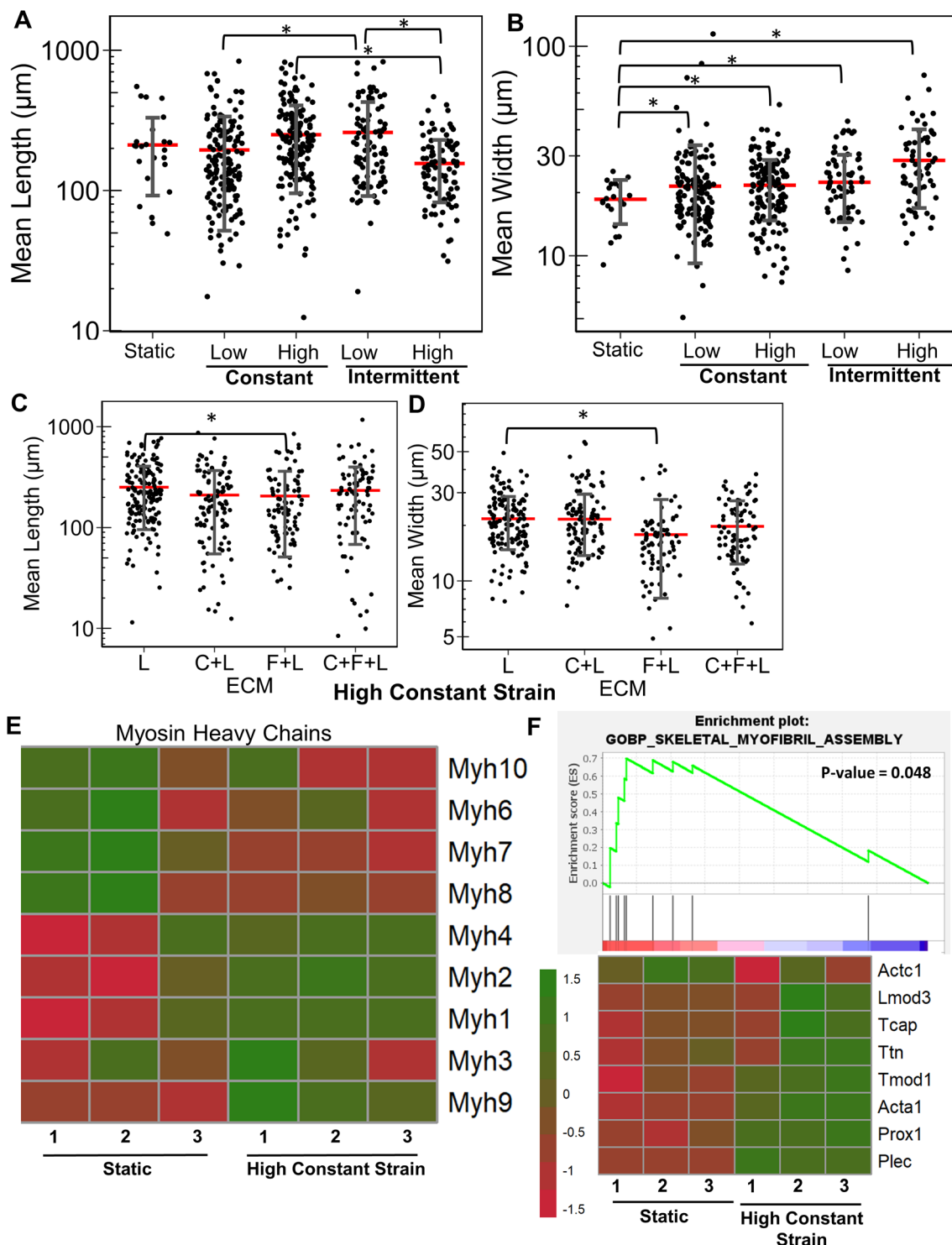


Fig. 4 Effect of uniaxial strain regimens on myogenesis cultured on laminin. A and B. Quantification of mean myotube length (A) and width (B) with the application of different strain regimens in C2C12 myoblasts cultured on laminin (L) substrates. C and D. Quantification of mean myotube length (C) and width (D) with the application high constant strain in myoblasts cultured on C + F + L, C + L, F + L, and L. Each dot represents individual myotube values from 3 independent experiments. Mean values are represented by the red line. Standard deviation is represented using the error bars. Statistical analysis was performed using ANOVA with *post hoc* Tukey test for individual comparisons. * denotes statistically significant comparisons ($P < 0.05$). E. Heatmap of differentially expressed myosin heavy chain genes in differentiating myoblasts cultured on laminin with high constant strain and static conditions. F. GSEA enrichment plot comparing differentiating myoblast gene expression when cultured on laminin with high constant strain (left-red) or on static condition (right-blue) for enrichment to skeletal myofibril assembly gene set in the GO biological process database. The associated genes and their expression in the database are represented in the heatmap below.



increased myotube width and length as well as resulted in overexpression of multiple myogenesis related genes.

3.5 Combined effects of strain and collagen I on myogenesis

Similarly, we examined the strain effects on collagen I. A significant reduction in myotube length and width was observed with application of specific strain regimens (ESI Fig. 1†). With the application of high constant strain, a significant 26%

increase in average myotube length was observed with C + F + L ECM compared to collagen I alone (Fig. 5A). However, it was C + L ECM combinations that had a significant increase in myotube width compared to collagen I alone (Fig. 5B). At the genome-wide level, upregulated genes on collagen I with high constant strain were enriched in categories related to ECM degradation and collagen degradation, compared to on static conditions (Fig. 5C). Furthermore, contrary to the strain effects

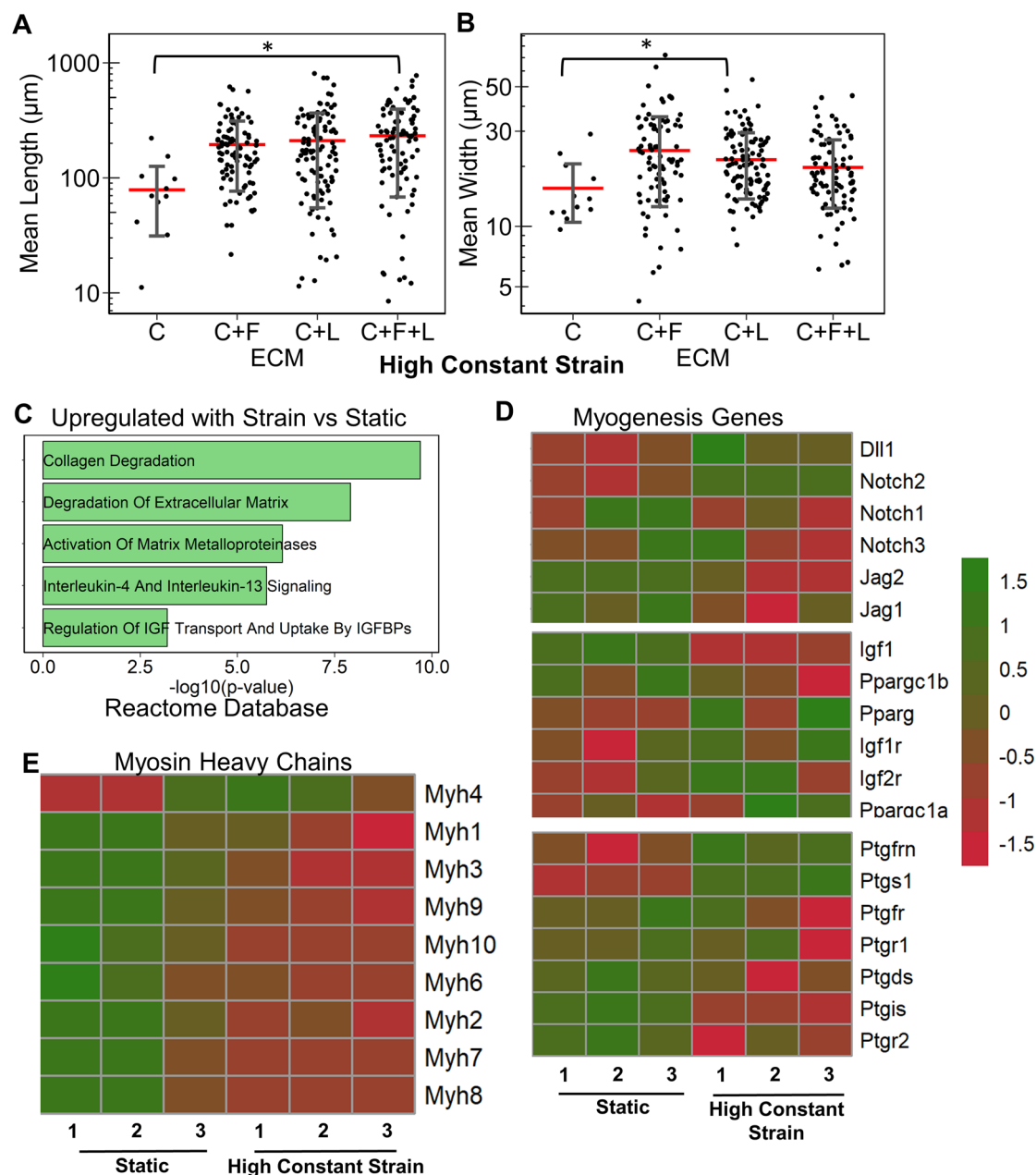


Fig. 5 Effect of uniaxial strain regimens on myogenesis cultured on collagen I. A and B. Quantification of mean myotube length (A) and width (B) with the application high constant strain in C2C12 myoblasts cultured on C + F + L, C + F, C + L, and C substrates. Each dot represents individual myotube values from 3 independent experiments. Mean values are represented by the red line. Standard deviation is represented using the error bars. Statistical analysis was performed using ANOVA with *post hoc* Tukey test for individual comparisons. * denotes statistically significant comparisons ($P < 0.05$). C. Enriched gene set categories upregulated in differentiating myoblasts cultured C with high constant strain, compared to static conditions L, based on the Reactome database. D and E. Heatmap of differentially expressed myogenesis pathway (D) and myosin heavy chain genes (E) in differentiating myoblasts cultured on laminin with high constant strain and static conditions.



on C + F + L vs. laminin (Fig. 4), strain on collagen I substrates led to a significant decrease in expression multiple myosin heavy chains (Fig. 5E). Additionally, upregulation of *Ptgs1*, *Notch2* on strain was observed, concomitant with a downregulation of *Igf1*, *Jag1*, *Jag2*, *Ppargc1b* was also quantified (Fig. 5D). Overall, application of different strains to differentiating myoblasts on collagen I did not improve myogenic potential, and instead multiple myogenic genes were downregulated.

3.6 Combinatorial ECMs under high constant strain conditions differentially regulate myogenesis

Having established the effects of different strain regimens, compared to static conditions, we then performed a comparative analysis of myogenesis in ECM compositions under high constant strain. Comparatively, the myotube length was significantly higher on C + F and C + F + L, compared to collagen I alone ($P < 0.03$, Fig. 6A). The myotube width was significantly higher on C + F, when compared to C + F + L ($P = 0.02$), F + L ($P = 0.00009$), or collagen I alone ($P = 0.02$) (Fig. 6B). A notable finding was that the ECM effect on myotube length was amplified by the application of high constant strain. For example, laminin exhibited a 3.2-fold increase in myotube length compared to collagen I under high constant strain (Fig. 6A), compared to only a 2.3-fold increase with static conditions (Fig. 1A). Similarly, under high constant strain, C + F + L also exhibited a significant 2.9-fold increase in myotube length, compared to collagen I (Fig. 6A), whereas there was no significant difference between the two ECM compositions under static conditions (Fig. 1A).

Transcriptome analysis revealed various molecular signatures unique to each ECM composition in response to the high constant strain application. Three gene clusters were found in myogenesis pathway genes, whereas two distinct clusters were found in ECM genes (Fig. 6C). Specifically, cluster A in myogenesis pathways genes corresponded to upregulation of *Igf1*, *Jag2*, *Dll1*, *Ppargc1b* on laminin, along with a significant downregulation on collagen I. In contrast, cluster B corresponded to downregulation *Notch2*, *Ptgs1*, *Igf1r* on laminin and upregulation on collagen I. Cluster C in the myogenesis pathways genes corresponded to specific downregulation of genes on C + F + L ECM combination (Fig. 6C). With respect to ECM-related genes, the effects of high constant strain could be distinguished by three distinct clusters corresponding to genes downregulated on C + F + L (*Fln1*, *Lamb1*, *Lama1*), genes upregulated on both C + F + L and L (multiple isoforms of *Col5* and *Col6*) and also the genes upregulated on C + F + L only (*Col4a5*, *Lamb2*, *Itgb1*) (Fig. 6D). Quantification of myosin heavy chains genes revealed upregulation of *Myh1*, *Myh2* and *Myh3* on laminin compared to collagen I (cluster A), whereas C + F + L led to the upregulation of *Myh8* and *Myh10* compared to both collagen I alone and laminin alone (cluster B) (Fig. 6E). These transcriptional signatures suggest that high constant strain effects on myogenesis show partial benefits in the setting of combinatorial ECMs, compared to single-component ECMs. Importantly, these findings also highlight the

complexity of cell–ECM interactions that often lead to non-intuitive findings in combinatorial ECMs, compared to single ECMs.

3. Discussion

In this study, we investigated the effects of combinatorial ECM proteins and different regimens of uniaxial stretching on myogenesis. The salient findings are that: (1) ECM compositions had a significant effect on myotube formation. Under static conditions, laminin was observed to produce the longest myotubes, whereas fibronectin produced the widest myotubes; (2) ECM combinations often produced non-intuitive effects compared to that of the single component ECMs that comprise them; (3) the benefits of combined strain and ECM cues are complex. Myotube formation findings showed that different strain regimes on laminin and C + F + L substrates led to enhanced myotube length and width, whereas no improvement on collagen I was observed with the addition of strain; and (4) genome-wide analysis of strain and ECM composition demonstrated potential benefits of combinatorial ECMs over single-component ECMs in the modulating of myogenesis.

We did not observe synergistic effects of combinatorial ECMs in relations to *in vitro* myotube morphology. However, when comparing transcriptional expression, we observed differential effects on the expression of combinatorial ECMs such as C + F + L, compared to collagen I alone. For example, cells cultured on collagen I alone in static conditions showed a significant downregulation of cell cycle related genes consistent with cell cycle withdrawal that precedes myotube formation (Fig. 2D), whereas cells on C + F + L substrates were enriched in actin cytoskeletal-related categories (Fig. 2E). These synergistic effects are compounded with mechanical stimulation, where myosin heavy chain gene expression levels were upregulated, compared to static culture alone (Fig. 3E). Together, our data suggests that combinatorial ECM microenvironment in concert with mechanical stimulation provide synergistic benefits to the myogenic process.

It is established that ECM proteins provide important biochemical cues for myoblast proliferation, attachment and differentiation.³⁶ In our assessment of ECM effects, we studied three ECM components of skeletal muscle tissue, which included collagen I, laminin and fibronectin. Our results showed that laminin produced the longest myotube and this agreed with Duffy *et al.*,³⁷ whereas Ding *et al.* observed fibronectin to be superior.³⁸ Ito *et al.* observed that collagen IV resulted in the longest myotube.³⁹ There are some conflicting data pertaining to the most myogenic ECM within the context of a singular component. In this study we have also included combination of the three ECM to investigate how multiple ECM interact to produce synergistic effects on myogenesis. We did not observe the additive or synergistic effect when combining two single-component ECMs. This could be due to a range of reasons such as interference of molecular binding sites of



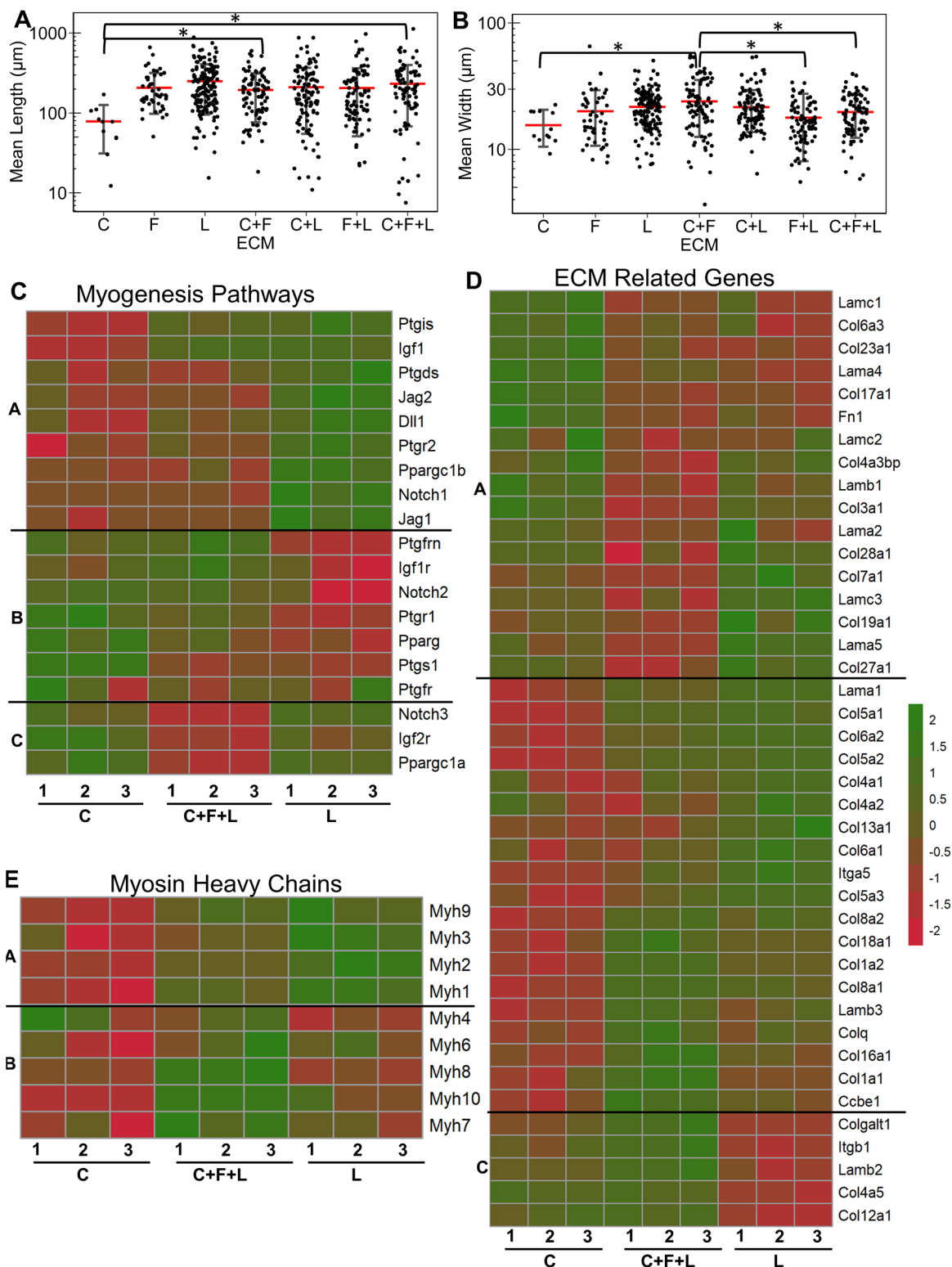


Fig. 6 Myogenesis of C2C12 myoblasts as a function of ECM combinations with high uniaxial constant strain. A and B. Quantification of mean myotube length (A) and width (B) in differentiating myoblasts cultured on ECM combinations at high constant strain. Each dot represents individual myotube values from 3 independent experiments. Mean values are represented by the red line. Standard deviation is represented using the error bars. Statistical analysis was performed using ANOVA with *post hoc* Tukey test for individual comparisons. * denotes statistically significant comparisons ($P < 0.05$). C–E. Heatmap of differentially expressed myogenesis pathway genes (C) ECM related genes (D), and myosin heavy chain genes (E) in differentiating myoblasts cultured on C, L and C + F + L with high constant strain.



the ECM competing for the cellular receptors or steric interactions within ECMs resulting in a net negative number of total binding sites for cell-binding receptors. Investigating the molecular level using RNAseq, we were able to discern differences between singular component ECM (collagen and laminin) with the combination of three ECMs (C + F + L). This highlights the nuances in early myogenesis that are not observed in mature myotube that express myosin heavy chain by immunofluorescence staining.

Mechanical stimulation also promotes differentiation and maturation of myoblast into myotubes. Uniaxial stretching was achieved using the Cytostretcher system from Curi Bio. This system has been used for studies involving chondrocytes and peripapillary scleral fibroblasts.^{40–43} Strain and frequency were chosen to corroborate with previous studies as strains of more than 20% decreases myogenesis.⁴⁴ Multiple studies have shown that mechanical stimulation increases myoblast differentiation and myotube formation.^{22,45–47} These tend to employ general molecular techniques to observe the regulation of known signaling pathways associated with muscle regeneration.⁴⁸ Here, we used RNAseq to determine transcriptomic changes with respect to combinatorial ECMs in conjunction with strain regimens.

One interesting finding is the upregulation of growth factor-associated pathways was only in the C + F + L group while undergoing constant high straining. This suggests that combinatorial ECMs with strain provide beneficial effects to the myogenesis process. This data falls in line with other studies using decellularized ECM from muscle related tissue or Matrigel basement membrane matrix, which was shown to improve myogenesis.⁴⁹ The synergistic effects of combinatorial ECM with mechanical stimulation could be potentially explained by mechanotransduction related pathways. Uniaxial stretching may also increase the presentation of ECM binding sites for cell interactions, as this phenomenon has been observed in fibronectin and collagen I.^{50,51}

A limitation of this study includes the small number of ECMs studied, in order to study the interaction of ECM composition with strain regimens. We selected collagen I in this study as it is the most abundant collagen I isoform, although collagen IV is the second most abundant isoform and may also exert important signaling function. Future studies should expand on the number of ECM proteins to be investigated.³⁹ Additionally, topography is also an important consideration. In this study, the cells were cultured on micropatterned substrates that confine the cells along the direction of strain. We and others have shown that micropatterning enables myoblasts to organize their cytoskeleton and form myotubes along the direction of the microgrooves.^{26,52,53} There have also been significant developments in incorporating spatio-temporal control of multiple biomechanical factors in directing cellular fate.^{54–57} Additional parameters such as three-dimensionality, substrate stiffness and viscoelasticity could be incorporated to develop a more biomimetic system that better reflects the complexity of the muscle microenvironment.^{58,59}

In conclusion, we have developed a stretchable tissue chip system to probe the combined effects of ECM composition and strain on myogenesis. Using this platform, we could systematically study the effects of ECM combinations along with strain regimes on myotube formation and the associated transcriptional signatures. Our findings illustrate the complexity by which ECM and strain cues modulate myogenesis in myoblasts. This stretchable tissue chip system has broad applications towards other cell types and for modeling *in vitro* diseases.

Data availability

All data from this project are available from the corresponding author upon reasonable request.

Author contributions

Conceptualization, A. H. P. C. and N. F. H.; methodology, A. H. P. C., I. J., B. N., N. G., and N. F. H.; investigation, A. H. P. C., I. J.; B. P. O., T. Z., resources, N. F. H.; data curation, A. H. P. C., I. J., and N. H.; writing – original draft preparation, A. H. P. C., I. J., and N. H.; writing – review and editing, A. H. P. C., I. J., B. N., N. G., B. P. O., and N. H.; visualization, A. H. P. C., I. J., and N. H.; supervision, N. F. H.; project administration, N. F. H.; funding acquisition, A. H. P. C., B. P. O., and N. F. H. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

B. D. and N. G. are employees of Curi Bio that manufactures the Cytostretcher equipment. Curi Bio, Inc. had no role in the design of *in vitro* or RNA sequencing studies.

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