



## Research article

## Altered microtubule structure, hemichannel localization and beating activity in cardiomyocytes expressing pathologic nuclear lamin A/C

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## ABSTRACT

Given the clinical effect of laminopathies, understanding lamin mechanical properties will benefit the treatment of heart failure. Here we report a mechano-dynamic study of *LMNA* mutations in neonatal rat ventricular myocytes (NRVM) using single cell spectroscopy with Atomic Force Microscopy (AFM) and measured changes in beating force, frequency and contractile amplitude of selected mutant-expressing cells within cell clusters. Furthermore, since beat-to-beat variations can provide clues on the origin of arrhythmias, we analyzed the beating rate variability using a time-domain method which provides a Poincaré plot. Data were further correlated to cell phenotypes. Immunofluorescence and calcium imaging analysis showed that mutant lamin changed NRVMs beating force and frequency. Additionally, we noted an altered microtubule network organization with shorter filament length, and defective hemichannel membrane localization (Connexin 43). These data highlight the interconnection between nucleoskeleton, cytoskeleton and sarcolemmal structures, and the transcellular consequences of mutant lamin protein in the pathogenesis of the cardiac *laminopathies*.

## 1. Introduction

In order to maintain normal heart functionality, cardiomyocytes detect and react to mechanical stresses. With each heartbeat, these cells transfer (via mechanotransduction) contractile stress mainly via cytoskeleton fiber components. These fibers (actin, microtubule, intermediate filaments) act as structural “bridges” between the cell exterior and the nucleus via the nuclear lamina. If one of these connections is missing, anomalous or unbalanced, the global mechano-dynamic behavior of the cardiomyocyte is affected. One example in which the mechanical link between the cell environment and the nucleus is perturbed is demonstrated in mutations of the *LMNA* gene encoding for lamin A and C, main components of the nuclear lamina. These mutations cause diseases called *laminopathies* [1, 2, 3]. *LMNA* mutations are responsible for progressive myocardial dysfunction leading to heart failure frequently requiring cardiac transplantation, and lethal arrhythmias [4, 5, 6]. Laminopathies show a compromised mechanical cell performance [7, 8, 9], primarily manifesting in tissues subjected to high levels of mechanical stress, such

as myocardial tissue. Although considerable progress has been made in understanding lamin structure and function, it is still not fully understood how different *LMNA* mutations affect both electrical function and contractility.

Nuclear lamins are intermediate filaments at the interface between the nuclear membrane and chromatin. They are critical for structural support of the nucleus, but evidence implies that nuclear lamins are also involved in other functions including nuclear envelope assembly, DNA synthesis, transcription, and apoptosis [10]. Furthermore, lamins, through specific proteins called “linkers of the nucleoskeleton and cytoskeleton (LINC)” directly transmit forces from the extracellular matrix into the nucleus [11]. Genetic defects in lamin A/C (*LMNA*) cause numerous *laminopathies*, disorders affecting muscle, adipose tissue, nerve, bone, and skin, and are responsible for premature-aging diseases (Progeria) [12]. *LMNA* mutations modify lamin filament organization and nuclear mechanical properties [13]. In previous work [14], we investigated three human *LMNA* mutations Glu161Lys (E161K, rs28933093), Asp192Gly (D192G, rs57045855), and Asn195Lys

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(N195K, rs28933091). We selected these three LMNA mutations because of their clinical and cellular characteristics: E161K is a recurrent mutation [15, 16, 17], D192G and N195K were associated with disruption of nuclear envelope morphology and altered internal organization of cardiomyocytes [18, 19]. We showed [14, 20, 21] that expression of  $LMNA^{E161K}$ ,  $LMNA^{N195K}$ , and  $LMNA^{D192G}$  in neonatal rat ventricular myocytes (NRVM) increased nuclear stiffness compared to wildtype  $LMNA$ . More precisely,  $LMNA^{D192G}$  expressing cells had the highest Young's modulus followed by  $LMNA^{N195K}$  and  $LMNA^{E161K}$ . These three  $LMNA$  mutations reduced the work of adhesion required to detach the spherical tip of an atomic force microscopy (AFM) cantilever from the cell membrane after indentation. In this case,  $LMNA^{D192G}$  mutation was the most disruptive, with a reduction of 45% of adhesion compared to the control cells. We confirmed that the deleterious effects of  $LMNA$  mutations extended beyond altered nuclear mechanics, to include defective cell membrane adhesion work. At the same time, cells expressing these mutations were more viscous compared to control cells, and stored less elastic energy in their cytoskeleton components when subjected to a rapid mechanical stress [20].

Because arrhythmias and poor myocardial contractility are clinical features of *laminopathies*, we hypothesized that beating frequency and force could be significantly altered at the single-cell level in mutant  $LMNA$  cardiomyocytes, and that the assessment of cardiomyocyte mechano-dynamic properties in  $LMNA$  mutant models will provide insights into the process leading to cardiomyopathy [22].

Here, we report a mechano-dynamic study of  $LMNA$  mutants in neonatal rat ventricular myocytes (NRVM) using single cell spectroscopy with AFM to measure changes in force, frequency and contractile amplitude of their beating. We investigated NRVMs carrying the three aforementioned  $LMNA$  mutation ( $LMNA^{E161K}$ ,  $LMNA^{N195K}$ , and  $LMNA^{D192G}$ ) and compared them to non-transfected (NT) and wild-type  $LMNA$  (WT) controls. We also analyzed the beating rate variability using a time-domain method and generated Poincaré plots. Data were

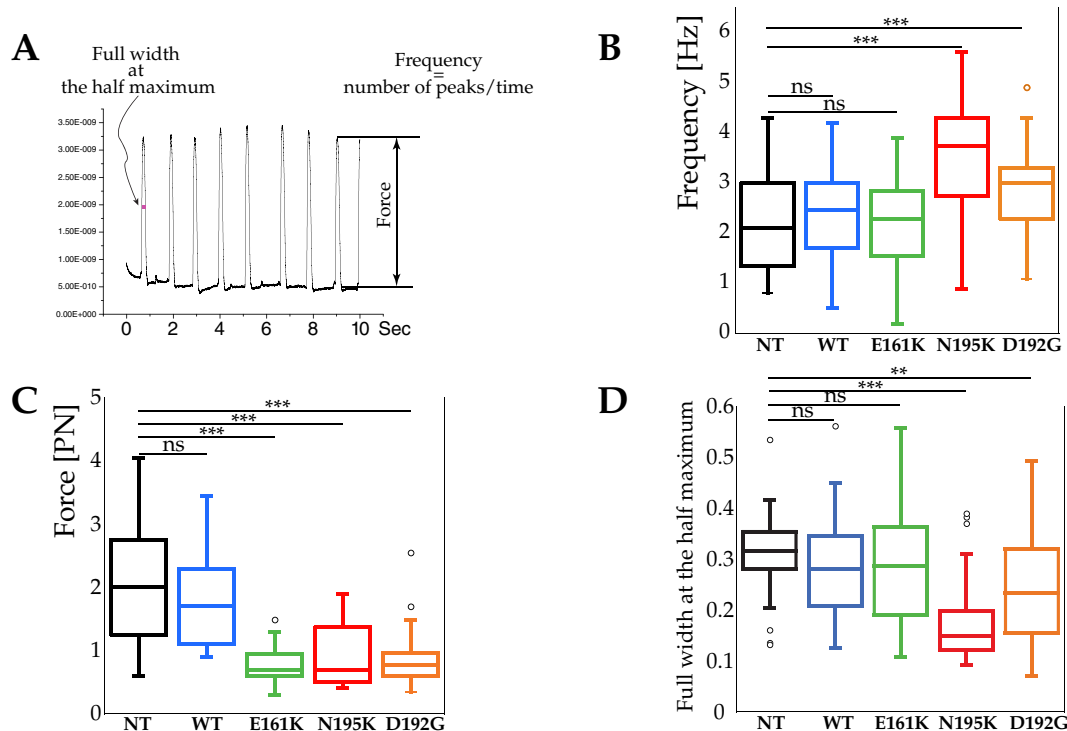
further correlated to cell phenotypes using immunofluorescence and calcium imaging analysis, showing that mutant lamin changes NRVMs beating force and frequency, and lead to altered cytoskeletal structure and hemichannel localization.

## 2. Results

### 2.1. Beating behavior in $LMNA$ mutant cardiomyocytes

To control our experimental conditions, such as temperature, pH and duration, the same protocol/methodology and the same model was used for all cell and AFM tests. Initially we checked how cell clustering affected cardiomyocyte behavior. Isolated beating cells and small clusters of beating cells were tested to assess variation in beating behavior. We found that isolated cells had significant mechano-dynamic variability during the acquisition time. Instead, control and mutant cell clusters showed a regular and stable beating pattern, providing more reliable data. Beating forces were obtained from AFM cantilever deflection data multiplied by the cantilever spring constant as detailed in the Method section. For instance, single NT cells beat with a force of 1.56 nanoNewtons (nN) (s.d. = 0.71, n = 26) while clusters beat with a force of 2.06 nN (s.d. = 0.93, n = 45,  $p = 0.01$ ); similarly, single WT cells beat with a force of 1.47 nN (s.d. = 0.54, n = 26) while cluster WT beat with a force of 1.84 nN (s.d. = 0.76, n = 45,  $p = 0.02$ ). Therefore, all further data reported in this study are related to selected cells within (in the center of) clusters obtained from NRVMs seeded at density of 20,000 cells/plate. To assess how the applied load influenced the beating property of our cells, we varied the applied AFM cantilever load. The signal remained constant for an applied force up to 4 nN; however, a 1 nN force was maintained on cells throughout all experiments.

Figure 1A shows raw AFM data. Figure 1B shows the beating frequency of NT, WT and cells expressing the  $LMNA$  mutations.



**Figure 1.** Mechanobiological cardiomyocyte properties: frequency, contraction force and peak duration (A) An example of the AFM trace during NT cardiomyocyte beating, (B) Box and whisker plots showing beating frequency of NT, WT and  $LMNA^{E161K}$ ,  $LMNA^{N195K}$ , and  $LMNA^{D192G}$  mutations. \*\*\*Represents  $p < 0.0001$ , (t-test) (C) Beating force. \*\*\*Represents  $p < 0.0001$ , (t-test) (D) Peak duration of  $LMNA$  mutant cardiomyocytes described as full width at the half-maximum (FWHM, namely the width of the beating pulse measured between those points on the y-axis which are half the maximum amplitude). \*\*\*Represents  $p < 0.0001$ , \*\* $p < 0.001$ , (t-test).

While there were no statistical differences between NT (2.23 beats/s, s.d. = 0.93, n = 45), WT (2.40 beats/s, s.d. = 0.89, n = 45) and *LMNA*<sup>E161K</sup> (2.21 beats/s, s.d. = 0.88, n = 54), both *LMNA*<sup>N195K</sup> (3.55 beats/s, s.d. = 1.14, n = 39) and *LMNA*<sup>D192G</sup> (2.87 beats/s, s.d. = 0.91, n = 36) mutations exhibited higher beating frequency as compared to NT.

Figure 1C shows the beating force for NT, WT and the three *LMNA* mutations. There were no statistical differences between NT (2.06 nN, s.d. = 0.93, n = 45) and WT (1.84 nN, s.d. = 0.76, n = 45) while *LMNA*<sup>E161K</sup> (0.76 n, s.d. = 0.27, n = 54), *LMNA*<sup>N195K</sup> (0.94 nN, s.d. = 0.49, n = 39) and *LMNA*<sup>D192G</sup> (0.88 nN, s.d. = 0.42, n = 36) had a lower beating force.

Figure 1D indicates the beat duration in terms of the full-width at half-maximum (FWHM). Values for NT, WT and *LMNA*<sup>E161K</sup> did not show any statistical differences ( $p = 0.20$ ) while *LMNA*<sup>N195K</sup> and *LMNA*<sup>D192G</sup> had small FWHM measurements than NT ( $p < 0.0001$  and  $p = 0.00094$ , respectively).

Investigations into beating rate variability (BRV) [23] utilized the Poincaré' geometric nonlinear analysis. Figure 2A shows a typical Poincaré plot. A quantitative analysis of the Poincaré plot can be created by approximating it as an ellipse, where (i) SD1 (Standard Deviation 1), is the standard deviation of the short-term beat-to-beat interval variability (minor axis of the ellipse), (ii) SD2 (Standard Deviation 2) is the standard deviation of the long term interval variability (major axis of the ellipse) [24]. Figure 2B, C show the SD1 and SD2 value for the *LMNA* mutations. In this case, only *LMNA*<sup>N195K</sup> and *LMNA*<sup>D192G</sup> displayed a statistical difference for both SD1 and SD2 compared with the NT cells, suggesting an irregular, arrhythmic beat of the mutant cardiomyocytes.

## 2.2. Altered connexin 43 localization in mutant *LMNA* cardiomyocytes

Since intercellular communication is a vital organizational feature of the heart [20, 25, 26] and disrupted ion channel function could affect contractile rates and forces, we examined the levels and localizations of gap-junctions through immunostaining for Connexin 43 (Cx43). Numerous studies have indicated that Cx43 organization is crucial for normal ventricular function and cellular impulse propagation in the healthy human heart [25, 27]. On the other hand, variation of the Cx43 organization might be one of the leading mechanisms for arrhythmias

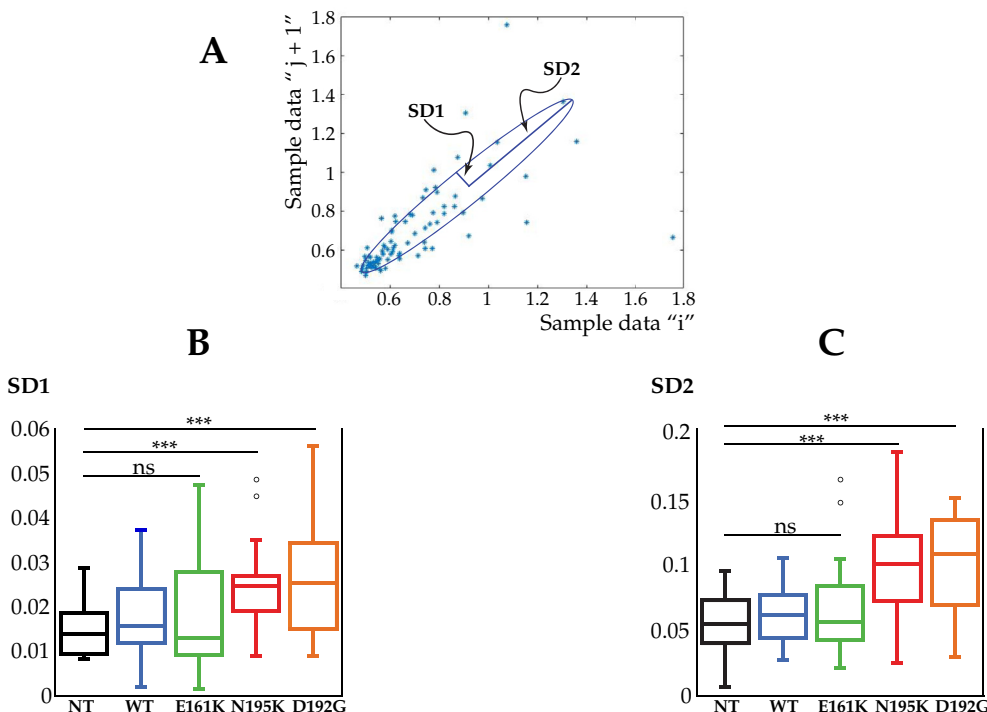
[28]. Figure 3A shows western blot results for Cx43; the results from four western blots indicate that there is no significant Cx43 difference between the groups. Figure 3B shows representative images of NRVMs stained for the cardiac marker  $\alpha$ -actinin, Cx43, GFP and nuclei; data are represented as fold change over NT. Figure 3B reveals that mutant NRVMs presented significantly smaller with punctate staining for Cx43 (161:  $0.24 \pm 0.07$ ; 192:  $0.44 \pm 0.11$ ; 195:  $0.21 \pm 0.17$  fold over NT) compared to controls (NT  $1 \pm 0.17$ -fold; WT  $0.95 \pm 0.22$ -fold over NT). The altered localization of Cx43 on the sarcolemma, in spite of preserved Cx43 protein expression, could be due to defective cytoskeletal structures anchoring to membrane proteins.

## 2.3. Altered calcium transients in mutant *LMNA* cardiomyocytes

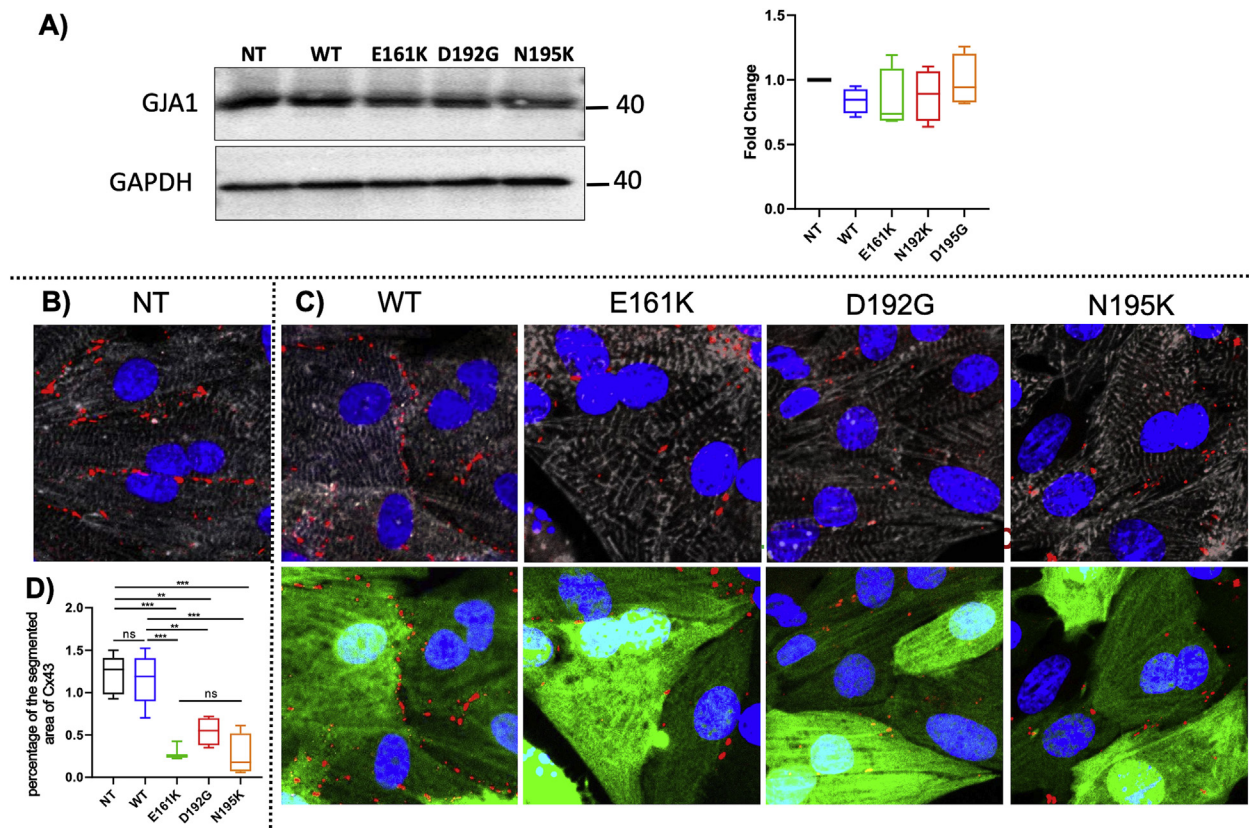
Calcium imaging was used to recapitulate the mechano-dynamic behavior assessed with AFM. Calcium transients were measured by the change in Fura Red fluorescence signal ( $\Delta F$ )/F0 according to Neher et al. [29] It is well known that calcium flux and contractile activity are related [30]. Wild-type cells exhibited rhythmic frequency and timing comparable to those of NT cardiomyocytes (Figure 4A). However, *LMNA*<sup>N195K</sup> and *LMNA*<sup>D192G</sup> cardiomyocytes exhibited a noisier spectrum for calcium spikes, suggestive of arrhythmic behavior (Figure 4A), and had significantly smaller intracellular  $Ca^{2+}$  transient amplitudes compared to those of the controls ( $p = 0.0001$ , Figure 4B). This finding was less pronounced in the case of *LMNA*<sup>E161K</sup> ( $p < 0.005$ ). Likewise, the frequency of calcium transients was significantly higher in *LMNA*<sup>N195K</sup> and *LMNA*<sup>D192G</sup> ( $p = 0.00091$  for *LMNA*<sup>N195K</sup> and  $p < 0.0001$  for *LMNA*<sup>D192G</sup> respectively) compared to controls, while in the case of *LMNA*<sup>E161K</sup> it was not statistically significant (Figure 4C). This impaired ability to handle  $Ca^{2+}$  agrees with the finding of a weaker force production and therefore a lower contractility.

## 2.4. Microtubule organization

Figure 5 shows disarray of the tubulin network in mutant *LMNA* NRVMs compared to controls. To quantify the tubulin network organization in the different mutations (Figure 5A), we used the



**Figure 2.** Beating rate variability by Poincaré plot of *LMNA* mutant cardiomyocytes. (A) A typical Poincaré Plot of RR intervals for NT cardiomyocyte. The peak interval (RR n) is displayed on the x-axis and the extent of the following interval (RR n+1) on the y-axis. All points described by successive beats of equal duration (RR n = RR n+1) are situated on the identity line. The points below this line represent all shortenings of the interval between 2 consecutive beats (RR n > RR n+1) while the points above the identity line correspond to all prolongations (RR n < RR n+1). (B) Analysis of the beating rate variability of *LMNA* mutant cardiomyocytes. SD1 (the standard deviation of the instantaneous - short term - beat-to-beat variability) and (C) SD2 (the standard deviation of the long-term interval variability) for NT, WT and *LMNA*<sup>E161K</sup>, *LMNA*<sup>N195K</sup>, and *LMNA*<sup>D192G</sup> mutations. \*\*\*Represents  $p < 0.0001$ , (t-test).



**Figure 3.** Cx43 protein expression in LMNA mutant cardiomyocytes. (A) Western blot showing Cx43 expression (left) and the results from western blots ( $n = 4$ ) indicating no significant difference between the groups (right),  $p = 0.5038$ . (B) Cx43 localization in control (NT) cardiomyocytes. Fluorescence images of Connexin 43 (red dots), sarcomeric  $\alpha$ -actinin (gray), LMNA GFP (green) and Hoechst (blue) staining of NRVMs (C) Cx43 localization in (from left to right): wild-type LMNA, LMNA<sup>E161K</sup>, LMNA<sup>D192G</sup>, LMNA<sup>N195K</sup>. Upper panels: Sarcomeric  $\alpha$ -actinin and connexin 43. Lower panels: LMNA GFP expression and connexin 43. (D) Quantification of Cx43 gap junction area: significant differences on Cx43 gap-junction were observed between LMNA<sup>E161K</sup>, LMNA<sup>D192G</sup> and LMNA<sup>N195K</sup> and the non-transfected NT cells and WT. Data are presented as mean  $\pm$  S.D. ( $n = 5$ ). 161 vs non-treatment: \*\*\* $p$  value:  $<0.0005$ ; 192 vs non treatment: \*\* $p$  value:  $<0.0045$ ; 195 vs non treatment: \*\*\* $p$  value:  $<0.0001$ ; 161 vs WT: \*\*\* $p$  value:  $<0.0009$ ; 192 vs WT: \*\* $p$  value:  $<0.0086$ ; 195 vs WT: \*\*\* $p$  value:  $<0.0002$ . Non-significant differences were observed between the mutant cells and between WT and NT. A full, non-adjusted image is provided in supplementary material (Figure S1).

optical Haralick correlation metric. This optical method computes texture features at multiple offset distances and angles. As shown in Table 1 and Figure 5B, the three mutant LMNA cell models showed contrast, uniformity (homogeneity or angular second moment), local homogeneity (Inverse Difference Moment (IDM)), entropy and correlation significantly different from the control LMNA models, indicating less organized microtubule networks with shorter microtubule filaments length.

### 3. Discussion

In this study, we found that mutant lamin (LMNA<sup>D192G</sup> and LMNA<sup>N195K</sup>) produced abnormal contractile and beating behavior of NRVMs, as indicated by decreased beating force, increased and irregular beating frequency, decreased and irregular calcium transients and abnormal sarcolemma localization of Cx43 hemichannels. These data all point to a strong correlation between the nuclear envelope, global cardiomyocyte mechanotransduction and electrical function in LMNA mutant cardiomyocytes, highlighting that lamin A/C function and dysfunction extend beyond the nuclear membrane. Not surprisingly, LMNA<sup>E161K</sup> caused more subtle changes, which aligns with the phenotype of LMNA<sup>E161K</sup> patients found to be less severe than in LMNA<sup>D192G</sup> and LMNA<sup>N195K</sup> patients [14].

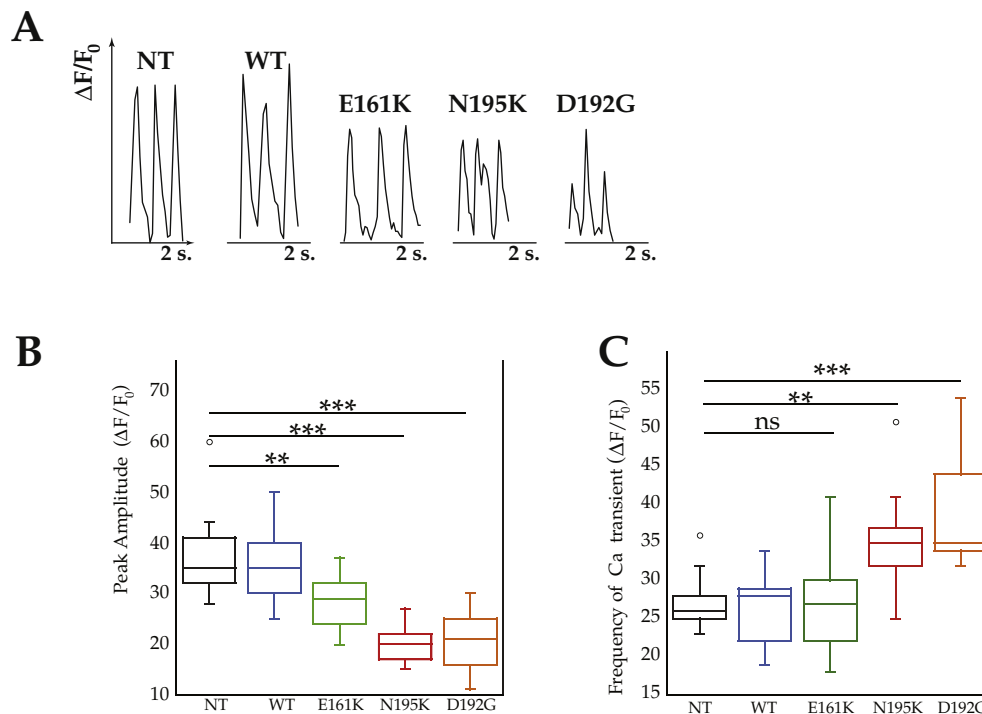
#### 3.1. Single cell spectroscopy unravels mechanical and electrical defects in LMNA mutant cardiomyocytes

We posited that valuation of the beating properties of mutant cardiomyocytes at the single-cell scale would help clarify the pathophysiology of heart failure and arrhythmias in laminopathies, and provide novel insights for the future development of new therapies. Moreover, our study at the single-cellular level offered unique advantages, such as the analysis of the inotropic/chronotropic behavior of cardiomyocytes with LMNA mutations without influences such as neuro-hormonal activities, or pre- and after-loads. Our findings are in line with single cell spectroscopy studies of other inherited cardiomyopathies, which likewise found altered beating frequency and force of mutant induced pluripotent stem cell (iPSC)-derived cardiomyocytes [31, 32, 33, 34].

As shown in Figure 1C, all three mutations showed a decreased force of beating suggesting a negative inotropic effect of the LMNA mutations. Furthermore, LMNA<sup>D192G</sup> and LMNA<sup>N195K</sup> mutations had a positive chronotropic effect, since the frequency of beating increased (Figure 1B). Finally, both the Poincaré plot, which analyzes beat-to-beat variability, and the Ca<sup>2+</sup> transients data revealed an irregular beating compared to controls, indicating an arrhythmic behavior at the cellular level.

It should be noted that for our single-cell spectroscopy analysis, we used the spherical AFM probe, which is ideal for soft biological samples since the force is applied to a broader cell area rather than with a sharp





**Figure 4.** Spontaneous calcium transients of NRVMs. (A) representative spontaneous calcium transient phenotype. From left to right: Control (NT), WT, *LMNA*<sup>E161K</sup>, *LMNA*<sup>N195K</sup> and *LMNA*<sup>D192G</sup>. (B) quantification of peak amplitude. (C) Quantification of Ca<sup>++</sup> transient frequency. \*\*\*Represents  $p < 0.0001$ , \*\* $p < 0.001$ , (t-test).

tip, resulting in a lower pressure and less cell damage. Due to the fact that cells or tissues are very inhomogeneous and consist of different components (nucleus, cytoskeletal components, etc.) a spherical probe will return more reliable data than a sharp tip.

### 3.2. *LMNA* mutations alter hemichannel localization in cardiomyocytes

For correct and efficient cardiac function, inter-cellular communications are crucial. Electrical and mechanical activities need to be synchronized so that the work of individual cardiomyocytes collectively organizes into normal cardiac contraction. Electrical current propagation throughout the heart is mediated through precisely regulated ion movements coordinated by various gap junction proteins [35]. Connexins assemble in hemichannels to form gap junctions which facilitate communication and signaling between adjacent cells. Cx43 is an important protein primarily expressed in the heart and in functioning myocytes, which ensures coordinated contractile action [36]. Our study found altered Cx43 localization with unchanged overall cellular levels of Cx43, which aligns with prior published data on the *LMNA*<sup>N195K</sup> mutation [37]. Little is known concerning the molecular mechanisms of remodeling of Cx43 in pathological context. Trafficking of Cx43 is regulated in part by the microtubule network [38, 39] and in fact, we observed a partial variation of the microtubule organization compared to our control cells as indicated by the disarray of the microtubule network (Figure 5, Table 1), with less organized microtubule networks and shorter microtubule filaments length. This result is in agreement with data from Macquart et al. [40] who showed that microtubule instability leads to an aberrant localization of Cx43 in cells expressing a lamin A mutation. Another study [41] suggested that appropriate nuclear lamina organization and a correct microtubule network is necessary for a collaborative effort to maintain a correct nuclear morphology and function. Indeed, we have already reported that the three mutations studied show changes in nuclear morphology [14]. Moreover, several studies showed that microtubules adversely affect the beating rate of neonatal

cardiomyocytes, in fact when microtubules in neonatal heart cells are depolymerized the beating rate increases [42, 43, 44, 45]. Using the Haralick optical technique of texture analysis (Table 1), we found that the microtubule network was compromised by the *LMNA* mutations, and at the same time, the beating rate was increased, providing a link between the defective nuclear envelope and the abnormal hemichannels localization.

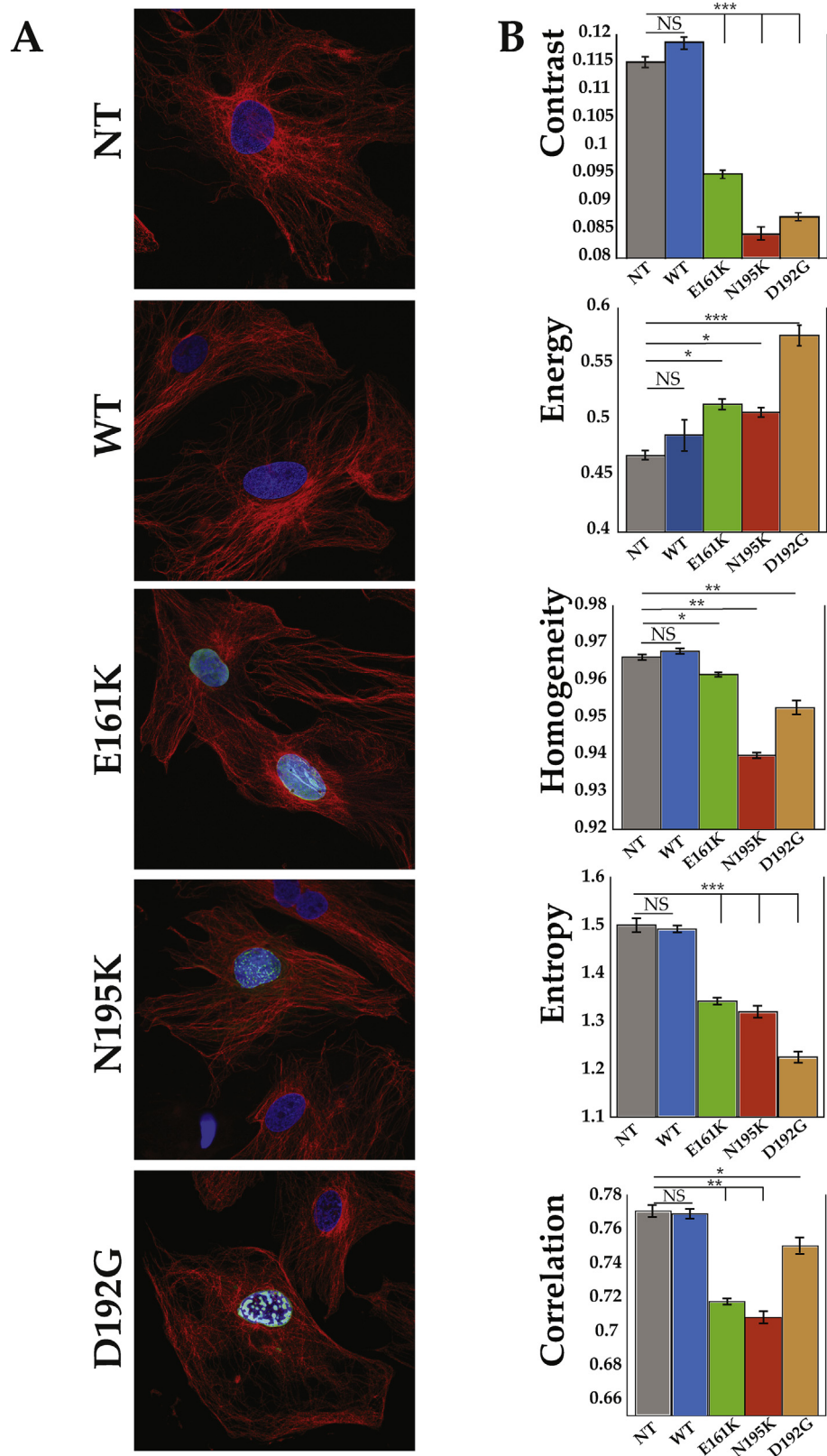
Our findings suggest that in cardiomyocytes, the chain of transfer of electrical and mechanical signaling from the cell membrane to the nucleus must be intact, and if one of the components, such as *LMNA*, is defective, the whole mechanism is compromised.

### 3.3. Limitations of the study

Even though it is well known that rodents show differences with the human biology [46], they represent an established and reproducible *in vitro* model [21], since cardiac tissue from patients is difficult to obtain and does not survive in long-term culture. Our results show that AFM can be used to assess the mechano-dynamic behavior of the chosen cells: we realize that orientation of actin-myosin filaments within a cardiomyocyte is anisotropic and therefore different parts of the cardiomyocytes may show varied amounts of movement and contractile forces with each beat. Furthermore, lateral modes of the contraction may not be accounted by the simple z-movement of the AFM cantilever. However, since in our methods of cell handling, study protocol and methodology, remained consistent throughout the entire series of experiments, we assume that our calculated beating force can be compared for all specimens.

## 4. Conclusions

Our study shows that *LMNA* gene mutations cause impairment of force production, frequency and rhythm of spontaneous beating and calcium handling in a single-cardiomyocyte cell model. Although the *in vitro* model presents limitations, it is intriguing that the changes found in



**Figure 5.** Microtubule network organization. (A) Example of microtubule organization used to calculate the Haralick features. (B) Haralick texture features: Contrast, Angular second moment (energy), Inverse difference moment (homogeneity), Entropy and Correlation are shown only for an angle of 0° (complete set of data in Table 1); Data are presented as mean ± S.D. ( $n = 4$  independent experiments). \* $p < 0.05$ ; \*\* $p < 0.005$ ; \*\*\* $p < 0.0001$ .

**Table 1.** Haralick correlation metric to determine microtubule organization.

Haralick Texture Features (0°)	NT	WT	%	E161K	%	N195K	%	D192G	%
Contrast	0.1151	0.1187	3.13	0.0951	17.38	0.0843	26.74	0.0874	24.06
Angular second moment (energy)	0.4678	0.4864	3.98	0.514	9.88	0.5070	8.37	0.5755	23.00
Inverse difference moment (homogeneity)	0.9526	0.9399	0.29	0.9615	0.93	0.9678	1.62	0.9663	1.46
Entropy	1.5004	1.4921	0.55	1.3424	10.53	1.3199	12.03	1.2248	18.37
Correlation	0.7707	0.7689	0.23	0.7171	6.95	0.7077	9.08	0.7501	2.66
Haralick Texture Features (45°)	NT	WT	%	E161K	%	N195K	%	D192G	%
Contrast	0.1403	0.1369	2.42	0.1041	25.80	0.0897	36.08	0.1032	26.39
Angular second moment (energy)	0.4495	0.4561	1.47	0.484	7.69	0.5031	11.93	0.5626	25.16
Inverse difference moment (homogeneity)	0.9300	0.9344	0.47	0.949	2.04	0.9552	2.71	0.9484	1.97
Entropy	1.5717	1.5466	1.60	0.714	53.92	1.3369	14.94	1.2730	19.01
Correlation	0.7212	0.7258	0.64	0.709	1.69	0.7052	2.22	0.7112	1.39
Haralick Texture Features (90°)	NT	WT	%	E161K	%	N195K	%	D192G	%
Contrast	0.1929	0.1853	3.94	0.129	32.86	0.1273	34.01	0.1217	36.94
Angular second moment (energy)	0.3657	0.37	1.18	0.385	5.28	0.3819	4.42	0.4945	35.21
Inverse difference moment (homogeneity)	0.9038	0.9098	0.66	0.933	3.23	0.9364	3.60	0.9392	3.92
Entropy	1.8226	1.7961	1.45	0.877	51.88	1.6512	9.41	1.3871	23.90
Correlation	0.6388	0.6455	1.05	0.566	11.2	0.572	10.46	0.5042	21.04
Haralick Texture Features (135°)	NT	WT	%	E161K	%	N195K	%	D192G	%
Contrast	0.2109	0.2094	0.71	0.185	2.51	0.1745	17.23	0.1386	34.25
Angular second moment (energy)	0.3558	0.3646	2.47	0.372	4.55	0.3904	9.72	0.4839	36.02
Inverse difference moment (homogeneity)	0.8948	0.9031	0.93	0.9490	6.06	0.9127	2.00	0.9308	4.02
Entropy	1.8573	1.8373	1.08	1.388	25.23	1.7738	4.49	1.4246	23.30
Correlation	0.6037	0.6121	1.39	0.496	17.68	0.4791	20.62	0.4604	27.9

cardiomyocytes recapitulate at the cellular level the clinical phenotype of LMNA cardiomyopathy patients, who typically present with decreased cardiac contractile function and arrhythmias, eventually leading to heart failure and sudden cardiac death. Single-cell spectroscopy by atomic force microscopy represents a unique tool to complement cardiac cell biology investigations and elucidates the complex mechanisms leading to cardiac dysfunction in *laminopathies*.

## 5. Materials and methods

### 5.1. Isolation and culture of ventricular cardiomyocytes from neonatal rat (NRVMS)

The University of Colorado Denver institutional ethical committee gave approval for these experiments. All animal studies were performed according to the guidelines Animal Care and Use Committee. All animal experiments have been done using all possible methods to alleviate or minimize potential pain, suffering or distress, and enhance animal welfare. Animals have been provided with housing in an enriched environment, with at least some freedom of movement, food, water and daily care and cleaning. The well-being and state of health of experimental animals have been observed by competent persons, dedicated to the management of the Animal House, able to prevent pain or avoidable suffering, distress or lasting harm. Experiments have been performed solely by competent authorized persons. Primary cultures of neonatal rat ventricular cardiomyocytes (NRVMS) were isolated and cultured from 1-3 days old Sprague Dawley rat pups (Charles River), following decapitation by enzymatic digestion as previously described with minor modifications [28, 47, 48, 49]. Briefly, ventricles were separated from the atria using scissors and then dissociated in CBFHH (calcium and bicarbonate-free Hanks with Hepes) buffer containing 0.5 mg/ml of Collagenase type 2 (Worthington, Biochemical Corporation), and 1 mg/ml of Pancreatine (Sigma). Cardiomyocytes were enriched (>90% purity) over non-myocytes by a pre-plating step on 100-mm dishes in Minimum Essential Media (MEM), supplemented with 5% bovine calf serum and 2 mg/ml vitamin B12 (Sigma). Myocytes that were either in solution or lightly attached were then separated from the adherent

stromal cells by gentle mechanical disaggregation and subsequently plated at a density of  $2 \times 10^5$  cells/ml in primary petri dishes (Falcon) or in multi-chambered slides coated with 0.1% gelatin (Sigma) and cultured as previously described [28, 49]. After 18 h, the culture medium was changed and cells were subjected to infection with the relevant adenoviral-LMNA constructs.

### 5.2. Isolation adenoviral constructs and infection

Methods for adenoviral infection have been previously reported [20, 21]. In brief, shuttle constructs were generated in Dual CCM plasmid DNA containing GFP gene and human LMNA cDNA. Constructs were bicistronic with the two inserts (LMNA and GFP) driven by two different CMV promoters to identify cells expressing LMNA protein using GFP as a marker of cellular infection. NRVMS were infected by adenoviruses at a multiplicity of infection (MOI) of 50 in serum free medium; 6 h post-infection, complete medium was replaced and the cells were incubated at 37 °C and 5% CO<sub>2</sub>. Previous control experiments have shown that GFP transfection and expression did not affect endpoints of interest in NRVMS in our experimental conditions [21]. Five experimental conditions were present: NT-non-transfected; WT-wild-type LMNA construct and the three LMNA mutant conditions (*LMNA*<sup>E161K</sup>, *LMNA*<sup>N195K</sup>, and *LMNA*<sup>D192G</sup>).

### 5.3. Western blotting

Proteins were extracted from transduced NRVMS as published [50]. Briefly, transduced cells were lysed in a RIPA Buffer containing protease inhibitors and phosphatase inhibitor. Protein concentrations were determined by colorimetry RC-DC protein assay (Bio-Rad Laboratories). Equal amount of protein extracts was denatured and resolved on SDS-PAGE gels, and were then transferred to nitrocellulose membranes. Target proteins were detected using primary antibodies by overnight incubation at 4 °C. The respective horseradish linked secondary antibodies were incubated at room temperature for 1 h at concentration recommended by the manufacturer. The signal was amplified by an ECL chemiluminescence assay (GE Healthcare) and was detected using gel

imager from Biorad. The primary antibodies used were as follows: rabbit polyclonal CX43 (Sigma, 1:1000) and mouse monoclonal GAPDH (ThermoFisher, 1:10000) as loading control.

#### 5.4. Immunofluorescence

NRVMs were fixed in PBS containing 4% PFA for 15 min at room temperature. Cells were permeabilized at room temperature, with 1% Triton X-100 for 90 min, blocked with 2% BSA in PBS for 45 min and incubated overnight with alpha actinin 1:500 (Sigma) and Connexin 43 1:200 (Sigma) prepared in 2% BSA. Goat anti-mouse antibody conjugated to Cy5 (abcam) and Goat anti-rabbit antibody conjugated to Alexa Fluor 594 were used as secondary antibody 1:300 (Invitrogen). Each sample was stained with Hoechst 1:2000 (Thermo Fisher Scientific) to counter-staining the nuclei. Representative immunofluorescence images were acquired using Zeiss LSM780. Expression of human wild-type and mutant LMNA proteins were verified by western blot to confirm exogenous human LMNA in NRVMs together with the endogenous rat level expression.

#### 5.5. Calcium imaging

To assess the electrical activity of cardiomyocytes with different mutations and controls, 2 days after infection, we recorded intracellular spontaneous calcium signaling. Cell-permeant Fura Red AM (Thermo Fisher Scientific) was added to each sample according to manufacturer instructions and incubated for 15 min. Samples were washed 3 times with warm media before imaging. Calcium transients were measured by the change in fluorescence divided by the resting fluorescence ( $\Delta F/F_0$ )<sup>29</sup>. Calcium transients during spontaneous cardiomyocytes beating were recorded using a Zeiss LSM780 confocal microscope and were measured for 21s (150 frames). Experiments were performed in triplicate from 3 independent experiments and averaged. Data was corrected for background epifluorescence.

#### 5.6. Atomic force microscopy beating frequency and force

AFM is one of the methods for measuring cardiomyocyte mechanodynamics properties [51]. In AFM, a small cantilever is brought in contact with the cell being measured. While the cell beats, the vertical movement of the cell is tracked producing the beating signal [52]. For instance, hPSC derived cardiomyocyte beating characteristics have been measured with AFM using this kind of protocol by Liu and colleagues [32].

In this work, a JPK NanoWizard 4a BioScience AFM was used to acquire the beating data. In AFM testing, it is very important to bear in mind that cells can be in different states and thus show distinct properties that can be difficult to compare. Therefore (i) AFM experiments were always performed on the same day after the adenoviral infection, (ii) multiple measurements from different cells have been collected to control for variability and 'average' data determined, and (iii) cells were monitored and their morphological details observed (an optical light microscope was used for cell selection throughout the tests). The AFM was equipped with PetriDishHeater™ tool to operate in liquid at controlled temperature (37 °C). A cantilever modified with a spherical gold probe (diameter of ~5.5 μm, CP-PNPL-Au-C, NanoAndMore GmbH) was used to precisely apply a compression force to the nucleus.

AFM probes were cleaned, prior to the indentation experiments, by embedding them in Tween (2% for 30 min) and then rinsed in milliQ water and ethanol, in order to remove contaminant molecules adsorbed on the probe surface.

All studies were performed on living, intact cells in cell culture medium. Only well-spread cells were investigated, those with a round shape and a dark edge were rejected. Furthermore, cardiomyocytes were

selected based on the morphological aspect compared to the possible presence of fibroblasts and only if they were beating. When cluster cells were selected, based on cluster size, one or more cells were tested (usually 2–3 cells in every cluster).

The AFM tip was moved toward the cell with speed of 5 μm/s. The speed range was chosen to avoid cell movement (at low compression speed) or hydrodynamic force contribution (significant at high speed). All experiments were performed at the same velocity since these AFM tests are rate-dependent. Data were taken from a single point on the cell's surface on top of the nucleus.

The distal regions, away from the nucleus, were avoided since measurements performed around the nucleus are less affected by artifacts due to the substrate stiffness. For acquiring the beating data, the AFM tip was lowered until it touched the cell and the cantilever registered a deflection corresponding to 1 nN, then the z-piezo was locked for 30 s and beating frequency and probe travels were recorded. The beating forces were found by multiplying the probe displacements due to contractile activity and the cantilever's spring constant, assessed using the thermal noise method in liquid. The beating frequency of the cardiomyocytes was computed by quantifying the number of peaks in a measurement of a sufficiently long time ( $\geq 15$  s). Peak duration was quantified in terms of full width at the half-maximum (FWHM, namely the width of the beating pulse measured between those points on the y-axis which are half the maximum amplitude). Data were processed using a self-developed MATLAB script. We used this approach either at single-cell resolution or on a cluster of cells which were beating synchronously. We assessed the beating force and frequency using at least 20 clusters for each condition (untreated/uninfected control: NT, wild type lamin: WT and three LMNA mutations: 161, 192 and 195). The test duration was never longer than 45–50 min to ensure cell viability.

#### 5.7. Beating rate variability

Beating rate variability (BRV) was evaluated using the Poincaré plot, automatically generated during MATLAB processing. The Poincaré plot is a technique suggested by Tulppo et al. [53] for studying heart rate signals, as it helps to assess components of the beat variability related with short- and long-term connections of the signal [54, 55, 56]. It is a scatter graph of the peaks interval plotted against the preceding peaks interval (RR). The first peaks interval [i] represents the x-coordinate, the second interval [i+1] represents the y-coordinate. The quantitative analysis of the plot is realized by fitting an ellipse to the plot with its center coinciding with the centroid of the ellipse. The point where both ellipse axes intersect corresponds to the total mean of the intervals. The length of minor and major axis of the ellipse are 2SD1, 2SD2, where SD1, SD2 are the dispersion perpendicular to the line of identity (minor axis) and along the line of identity (major axis) respectively as proposed by [53]. SD1 is the standard deviation of the instantaneous (short term) beat-to-beat variability while SD2 is the standard deviation of the long-term interval variability.

#### 5.8. Microtubule organization

Microtubule organization is calculated using Haralick features, which are pixel intensity-based algorithms for quantifying image texture [57, 58]. First, a gray-level co-occurrence matrix is calculated for given orientation and pixel pair offset distances. The co-occurrence matrix p calculates the frequency at which pixels within a specified intensity range are matched by spatially separated pixels of the same intensity. The result is a  $i \times i$  matrix, where  $i$  is the number of gray-levels (or intensity bins) that are to be considered. From the co-occurrence matrix, several texture features can be measured. Haralick correlation is one of these features that measures the likelihood of finding two pixels of similar intensity separated by a given distance. Calculation of the co-occurrence matrix



and corresponding Haralick correlation values was repeated at many orientation angles and spatial distances, resulting in an “ $m \times n$ ” matrix of Haralick correlation values, where  $m$  is the number of angles (values are symmetric about  $180^\circ$ ) and  $n$  is the number of pixel offset. Other Haralick features that were considered in the comparison analysis include Contrast, Uniformity (also known as homogeneity or angular second moment), local Homogeneity (or Inverse Difference Moment (IDM)), Entropy and Correlation. We also measured a variance value, which is the sum of the Haralick contrast values at an offset distance of 1 pixel. We expected the variance value to be close to zero for homogenous images and close to one for patterned images.

### 5.9. Statistics

For all experiments, data were collected from at least three independent experiments. ImageJ was used for cell counting in 10 fields per each condition. For calcium transients, signals from myocytes were collected and analyzed from 10 fields per sample and averaged. Statistical significance between experimental groups was determined using ANOVA. ANOVA Power analysis was completed using GraphPad Prism7 software. A  $p$  value of  $<0.05$  was considered statistically significant.

### Declarations

#### Author contribution statement

O. Sbaizero: conceived and designed the experiments; analyzed and interpreted the data; wrote the paper.

D. Borin: performed the experiments.

B. Pena and S. Chen: performed the experiments; analyzed and interpreted the data.

C. Long: conceived and designed the experiments; wrote the paper.

M. Taylor and L. Mestroni: Conceived and designed the experiments; contributed reagents, materials, analysis tools or data; wrote the paper.

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#### Competing interest statement

The authors declare no conflict of interest.

#### Additional information

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