Basal oxidation of conserved cysteines modulates cardiac titin stiffness and dynamics

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Abstract

Titin, as the main protein responsible for the passive stiffness of the sarcomere, plays a key role in diastolic function and is a determinant factor in the etiology of heart disease. Titin stiffness depends on unfolding and folding transitions of immunoglobulin-like (Ig) domains of the I-band, and recent studies have shown that oxidative modifications of cryptic cysteines belonging to these Ig domains modulate their mechanical properties in vitro. However, the relevance of this mode of titin mechanical modulation in vivo remains largely unknown. Here, we describe the high evolutionary conservation of titin mechanical cysteines and show that they are remarkably oxidized in murine cardiac tissue. Mass spectrometry analyses indicate a similar landscape of basal oxidation in murine and human myocardium. Monte Carlo simulations illustrate how disulfides and S-thiolations on these cysteines increase the dynamics of the protein at physiological forces, while enabling load- and isoform-dependent regulation of titin stiffness. Our results demonstrate the role of conserved cysteines in the modulation of titin mechanical properties in vivo and point to potential redox-based pathomechanisms in heart disease.

1. Introduction

The passive mechanical properties of cardiomyocytes, such as stiffness, are fundamental for the normal physiology of the heart, and their dysregulation contributes to deficits in the mechanical activity of the myocardium in different forms of heart failure [1], for instance following myocardial infarction [2] or in diabetic patients [3]. A major source of passive stiffness in cardiomyocytes is titin, a giant protein in sarcomeres that also provides structural support and mechanosensing functions (Fig. 1A) [4–6]. The functional relevance of titin is exemplified by the fact that truncating variants in the titin gene (TTN) are the main cause of dilated cardiomyopathy (DCM), a disease that is the most frequent trigger of heart failure in the young and of heart transplantation worldwide [7–11].

The mechanical properties of titin are exquisitely modulated both transcriptionally through specific alternative mRNA splicing [12,13], and posttranslationally via phosphorylation [14]. However, these two mechanisms alone fail to capture the full range of titin mechanical adaptations, showing our incomplete understanding on how the stiffness of cardiomyocytes is affected by biochemical signals targeting titin [2]. The mechanical properties of titin stem from force-dependent conformational changes of the extensible I-band region of the protein (Fig. 1A). These conformational changes include extension and entropic recoil of the serially linked immunoglobulin-like (Ig) domains and the

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random-coil N2Bus and PEVK regions, and Ig domain unfolding and refolding transitions [15]. Interestingly, the I-band of titin is rich in cysteine residues, many of which appear at structurally conserved positions within Ig domains [16,17]. In vitro experiments have shown that oxidation of these cysteines has major mechanical consequences. For instance, disulfide bonds established by the triad of structurally conserved cysteines B, F and G in Ig domains stiffen titin via reduction of the contour length of the disulfide-containing unfolded state leads to domain stiffening. C: High-resolution structure of the traditionally named 127 domain of titin (also known as 191, PDB code 1tit), which contains unpaired, S-thiolation-competent cysteines 47 and 63 (positions 12674–12765 from Uniprot entry Q8WZ42). S-thiolations inhibit folding, leading to domain softening [19]. D: Map of cysteine positions along the alignment of Ig domains of human titin (breaks leave out Cys-free segments of the alignment). Structurally conserved positions B, 47, 63, F and G are indicated on top of the alignment. Positions are colored according to percentage of evolutionary conservation, from 0 (white) to 100% (dark green). The number and positions of Ig domains according to Uniprot Q8WZ42-1 (N2BA isoform) are indicated on the right. Pairs of dots mark Ig domains that are not annotated in Uniprot [18]. Ig domains are represented in grey if they are cysteine-free, in orange if they contain at least two of the triad, disulfide-competent cysteines B, F and G (disulfide-competent domains) [18], in yellow if they contain cysteines but no more than 1 cysteine B, F or G (domains incompatible with disulfide bond formation, only subject to potential S-thiolation), and in brown if at least two triad cysteines are present together with other cysteines (domains containing both disulfide-competent cysteines and cysteines that can only experience S-thiolation). The spaced-out region in the N2B titin isoform is indicated by the shaded grey area. The horizontal dashed line shows the boundary between the I- and A-band regions of titin (positions 14018–14019). E: Violin-plot distribution of percentages of evolutionary conservation of structurally conserved (SC, including BFG and 47/63 cysteines, n = 193), non-structurally conserved (NSC, n = 113), and N2Bus (n = 6) cysteines in human titin. p(SC vs NSC) = 0.0001, p(SC vs N2Bus) = 0.0001, p(NSC vs N2Bus) = 0.0033 (Kruskal-Wallis and Dunn’s multiple comparisons test). Evolutionary conservation values were calculated from the alignment of 36 titin sequences from different species (Supplementary Fig. S1 and Supplementary File S1). Horizontal bars in violin plots indicate median values. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

2. Results

2.1. Evolutionary conservation of titin mechanical cysteines

In the light of the potential physiological relevance of titin’s mechanical cysteines, we first explored their evolutionary conservation by aligning the available titin sequences from 36 species of vertebrates [22] (Supplementary File S1). The vast majority of cysteine positions in human titin show more than 95% occupancy in the sequence alignment (i.e. sequence gaps do not typically appear at alignment positions occupied by cysteines, Supplementary Figs. S1A and B), which shows the robustness of our evolutionary conservation analyses. We present the percentage of evolutionary conservation for all cysteines belonging to Ig domains of human titin in Fig. 1D. This figure is built from a sequence alignment of both I- and A-band Ig domains of human titin (y-axis) [18], in which we highlight every cysteine position colored according to their evolutionary conservation. We found that cysteines B, F and G in Ig domains show 98% mean evolutionary conservation (Fig. 1E) and cysteines 47 and 63 are also highly conserved (94% mean evolutionary conservation, Fig. 1E). These five structurally conserved cysteines are more evolutionarily conserved than cysteines appearing in other positions in Ig domains (97% vs 89% mean evolutionary conservation, Fig. 1F). The N2Bus region also contains cysteines that can stiffen titin through disulfide bond formation [23], albeit they are less evolutionarily conserved than Ig domain cysteines (53% mean conservation, Fig. 1F, Supplementary Fig. S1C). In summary, the significant conservation of titin’s mechanical cysteines throughout evolution supports their importance for the biological function of the protein.

2.2. Titin cysteines are oxidized in basal conditions

Beyond the remarkable conservation of I-band titin’s cysteine residues and their role in the evolution of the protein in vertebrates [22], the in vivo relevance of redox mechanical modulation of titin is supported by limited data on the global oxidation of the protein [20,21,24–27], the effects of redox-active molecules on striated muscle mechanics [19,23,28,29], and the disulfide-compatible location of the majority of structurally conserved cysteines of titin [18,30]. However, the extent and location of native titin oxidations, particularly in human titin, remain largely unexplored. Recently, Loescher et al. have shown using mass spectrometry that under oxidative insults ex vivo and in vivo, induced oxidations preferentially target cysteines in the distal I-band region of murine cardiac titin through a mechanism involving protein unfolding [20]. However, our grasp of the range of redox modulation of titin remains limited since it is unknown whether and to what extent titin cysteines are oxidized also in basal, non-oxidative conditions. Whether oxidations target human titin is also unknown.

To examine basal oxidation of titin, we first developed a SDS-PAGE-based assay to define experimental conditions that minimize potential artifactual oxidations that can result from sample manipulation or inefficient thiol derivatization [31]. The assay exploits thiocysteine reactions to block reduced cysteines with an alkylation agent and label reversibly oxidized cysteines with the fluorophore monobromobimane (mBBr) (Fig. 2A) [32]. Titin oxidation was directly quantified after running 3.5% acrylamide SDS-PAGE gels, in which the slowest migrating band is contributed by several titin isoforms in the tissue [33,34]. We verified that the mBBr fluorescence signal of titin cysteines is linearly dependent on the amount of lysate analyzed (Fig. 2B, Supplementary Fig. S2A) and used Coomassie staining (Fig. 2C, Supplementary Fig. S2A) to get normalized oxidation measurements that are independent of the amount of protein loaded on the gel (Fig. 2D). The initial alkylation is done in denaturing conditions to block all reduced cysteines. This step needs to be fast and complete to limit artifactual oxidation signal. We chose N-ethylmaleimide (NEM) as the alkylation agent because of its superior reaction kinetics and its high solubility in aqueous buffers [26,31]. Using readily available cardiac samples from adult mice, we found that titin oxidation signal plateaus at NEM concentrations above 5 mM (Fig. 2E), suggesting that these conditions result in minimal artifactual oxidation. To further ensure efficient thiol blockage, in all our subsequent experiments we perfused myocardial tissue with 50 mM NEM immediately after sacrifice. The same high NEM concentration was kept during lysis. As an additional preventive measure, we subtracted the mBBr signal of samples not incubated with DTT to account for potential reduced thiols refractory to NEM blockage (see Methods). Using our in-gel mBBr fluorescence assay, we observed that...
the extent of reversible cysteine oxidation of titin is ~4 times higher than that of myosin, a partner protein of titin in the sarcomere (Fig. 2F, Supplementary Fig. S2B, to allow comparison, oxidation signals are normalized by the density of cysteines of both proteins, see Methods), confirming that a fraction of titin cysteines are constitutively oxidized in cardiac tissue.

2.3. Preferential oxidation of I-band cysteines

To characterize the landscape of reversible titin cysteine oxidations, we resorted to mass spectrometry (MS). In this set of experiments, we focused on myocardial samples from P0 newborn mice because they contain a high proportion of the cysteine-rich N2BA titin isoform [18, 35] and therefore provide better cysteine coverage (Supplementary Fig. S3). Fluorescent titin bands were sliced from SDS-PAGE gels and treated with trypsin. The resulting peptides were subjected to LC-MS analysis and MS/MS spectra were searched against a mouse proteome database. As expected from the full titin band separation in 3.5% SDS-PAGE gels, 79 ± 1% (3 samples from different whole mouse hearts) of the identified species corresponded to titin-derived peptides, even when using a relaxed identification criterion (5% False Discovery Rate, FDR) (Fig. 3A, Supplementary Fig. S2A). The mBBr and Coomassie signals originating from titin are linearly dependent on the amount of lysate loaded in SDS-PAGE gels (all data points are the average of duplicates) (Supplementary Fig. S2A). D: Final mBBr/Coomassie ratio is independent of the amount of lysate loaded in the gel. E: Scan to determine the concentration of NEM needed to completely block initially reduced titin thiols. Duplicates were performed for each NEM concentration. F: Quantification of the oxidation of titin and myosin in four different mouse hearts using 3.5% and 12% SDS-PAGE gels, respectively (see also Supplementary Fig. S2B). The volume of lysate analyzed was adjusted so that both titin (15 μl) and myosin (3 μl) were in the linear range of detection for Coomassie. Oxidation signal is normalized by the density of cysteines of each protein (13.3 Cys/1000 amino acids for titin, 7.2 Cys/1000 amino acids for myosin, see Methods). n = 4 animals, p = 0.0286 (Mann-Whitney).

Fig. 2. Titin is oxidized in basal conditions. A: Left: Reaction scheme to label reversibly oxidized thiols with mBBr. Protein extracts in which reduced thiols have been initially blocked by NEM are run in an SDS-PAGE gel. Following electrophoresis, oxidized thiols are reduced by DTT and then labeled with the fluorescent probe mBBr. Right: example SDS-PAGE gel used to quantify mBBr fluorescence from titin bands. Coomassie staining (Coom) of the same gel is used for normalization. B, C: The mBBr and Coomassie signals originating from titin are linearly dependent on the amount of lysate loaded in SDS-PAGE gels (all data points are the average of duplicates) (Supplementary Fig. S2A). D: Final mBBr/Coomassie ratio is independent of the amount of lysate loaded in the gel. E: Scan to determine the concentration of NEM needed to completely block initially reduced titin thiols. Duplicates were performed for each NEM concentration. F: Quantification of the oxidation of titin and myosin in four different mouse hearts using 3.5% and 12% SDS-PAGE gels, respectively (see also Supplementary Fig. S2B). The volume of lysate analyzed was adjusted so that both titin (15 μl) and myosin (3 μl) were in the linear range of detection for Coomassie. Oxidation signal is normalized by the density of cysteines of each protein (13.3 Cys/1000 amino acids for titin, 7.2 Cys/1000 amino acids for myosin, see Methods). n = 4 animals, p = 0.0286 (Mann-Whitney).
Fig. 3. Landscape of cysteine oxidation in murine and human cardiac titin. A: Fraction of identified peptides belonging to murine titin (reference sequence Uniprot A2ASS6-1, which codes for the long N2BA titin isoform) when the MS/MS spectra are searched against a mouse proteome database containing common contaminants (cRAP). B: Aggregated titin coverage obtained after searching against the mouse proteome (n = 3 mouse hearts; yellow and green, peptides detected with FDR<5% and <1%, respectively). C: Redox Identification Index (RII) map for every cysteine position in the alignment of Ig domains of mouse titin, according to the color code indicated on the left (red, oxidized; blue, reduced). Average RII (AvRII) for cysteines 47, 63 and the BFG triad across all Ig domains are indicated. Cysteines shown in white were not identified and do not contribute to RII calculation. The horizontal dashed line shows the boundary between the I- and A-bands of titin (positions 14880-14881 in Uniprot A2ASS6-1). The number and positions of Ig domains are indicated on the right. Ig domains are classified and colored as in Fig. 1. D: Violin-plot distributions of RII for cysteines belonging to the I- (n = 97) and A- (n = 123) bands of murine titin. p(RII >0, I-band) = 0.0054, p (RII >0, A-band) = 0.9868 (hypergeometric test). E: Violin-plot distributions of RII for mouse titin cysteines BFG (n = 61), 47/63 (n = 14) and non-structurally conserved (NSC, n = 33). p(RII >0, BFG) = 0.0054, p (RII >0, 47/63) = 0.8137, p(RII >0, NSC) = 0.9268 (hypergeometric test). F: Violin-plot distributions of RII for mouse cardiac titin cysteines B (n = 21), F (n = 23), G (n = 17), 47 (n = 8), 63 (n = 6) and NSC (n = 33). p (RII >0, B) = 0.9959, p(RII >0, F) = 0.0001, p(RII >0, G) = 0.0250, p(RII >0, 47) = 0.53365, p(RII >0, 63) = 0.7894, p(RII >0, NSC) = 0.9268 (hypergeometric test). Horizontal bars in violin plots indicate the median values. G: Fraction of identified peptides belonging to human titin (reference sequence Uniprot Q8WZ42-1, which corresponds to the long N2BA titin isoform) when the MS/MS spectra are searched against a database containing the human proteome and common contaminants (cRAP). H: Aggregated human titin coverage obtained after searching against the human proteome (left ventricular samples from 2 different non-transplanted non-failing donor hearts; yellow and green, peptides detected with FDR<5% and <1%, respectively). I: Redox Identification Index (RII) map for every cysteine position in the alignment of Ig domains of human titin, according to the color code indicated on the left (red, oxidized; blue, reduced). Average RII (AvRII) for cysteines 47, 63 and the BFG triad across all Ig domains are indicated. Cysteines shown in white were not identified and do not contribute to calculation of RII. The dashed line shows the boundary between the I- and A-band regions of titin (positions 14018-14019 in Uniprot Q8WZ42-1). The number and positions of Ig domains are indicated on the right. Pairs of dots mark Ig domains that...
cysteine oxidations building on the conserved location of titin mechanical cysteines (Fig. 1B–E). Fig. 3C shows RII values for each cysteine position in the alignment of the Ig domains of murine titin. RII data show that titin cysteines of the mechanically active I-band are more frequently detected as oxidized than those belonging to the A-band (Fig. 3D). In addition, cysteines of the disulfide-competent triad BFG (in particular cysteines F and G) are detected as oxidized more frequently than other cysteines in titin (Fig. 3C,E,F, Supplementary Table S1). Unfortunately, we did not detect cysteine-containing peptides from the N2Bus region, so its native oxidation state could not be studied.

2.4. Titin oxidation in human hearts

We next examined the landscape of cysteine oxidation in human cardiac titin using left ventricular snap-frozen samples from 2 non-transplanted, non-failing donor hearts. The proportion of N2BA titin in adult human left ventricle is 30% [1], which ensures a good coverage of mechanically active, conserved cysteines [18]. The human samples were processed and analyzed following the same approach used for mouse samples. 74 ± 3% of identified peptides were derived from titin when the MS/MS search was done against the human proteome database at a 5% FDR (Fig. 3G). In these searches, titin coverage was 37 ± 6% (Fig. 3H). As in experiments using mouse tissues, we ran targeted searches against a database containing only human titin and the resulting cysteine peptides were validated with Vseq. mBBR-derivatized peptides also detected in –DTT samples were excluded from analysis, leading to a final 39% aggregated cysteine coverage at a FDR<2%. As in murine titin, cysteines of the I-band are more frequently detected as oxidized than those of the A-band (Fig. 3I and J) and the RII of structurally conserved cysteines in Ig domains tends to be higher than that of non-structurally conserved cysteines (Fig. 3K, Supplementary Table S1). Similar to results with murine samples, cysteine F shows the highest RII among disulfide-competent positions, although in human samples the I-band-specific cysteine 47 has the highest RII among the structurally conserved cysteines (Fig. 3KL, Supplementary Table S1). In one of the human samples, we detected two cysteines belonging to the N2Bus region (Cys4083 was detected as reduced, and Cys4124 was found both in oxidized and reduced states). In summary, MS results indicate that reversible oxidations are also present in native titin from human myocardium.

2.5. Modulation of titin mechanics by redox modifications

Our results show that evolutionary and structurally conserved cysteines of the I-band of titin are oxidized in vivo. We find reversible oxidations both in disulfide-competent cysteines and in unpaired cysteines that can establish S-thiolation adducts. These modifications have been proposed to induce opposite mechanical effects [17–19], an observation that may contribute to explain the different modulation of titin-based striated muscle stiffness under specific redox challenges [19,20,23,28,29]. To illustrate the range of regulation of titin mechanics by cysteine oxidation, we built on previous Monte Carlo simulations [19] to integrate all known mechanical effects of redox posttranslational modifications in titin Ig domains. These include reduction of contour length by disulfide bonds [18], higher unfolding and folding rates of disulfide-containing domains [18,38], and higher unfolding rates and hampered folding of S-thiolated domains [19]. In our simulations, we tuned unfolding and refolding rate constants to qualitatively reproduce the recently described unfolding/folding dynamics of native titin [33] (Supplementary Note S1, Supplementary Figs. S4A). In the simulations, a virtual human I-band titin is subject to 1 Hz triangular force pulses between 0 and a predefined peak force, and the resulting length of titin is measured (Fig. 4A and B). During the extension/relaxation cycles, titin domains unfold and refold stochastically according to their folding and unfolding rates, which are dependent on their redox state. At t = 0 s, all domains are folded and, as the simulations proceed, a fraction of domains transition to the unfolded state resulting in longer titin lengths (Fig. 4C and D). The simulation time was long enough to reach steady-state lengths at all peak forces (Fig. 4C and D, Supplementary Figs. S4B–E).

Simulations of the canonical N2BA titin at a low peak force of 10 pN show that disulfides and S-thiolations result in longer titin lengths (i.e. lower stiffness) (Fig. 4C), while at a peak force of 100 pN, the effect of disulfides reverses leading to overall titin stiffening (Fig. 4D). At this high peak force, S-thiolation maintains its softening effect. Additive mechanical modulation occurs if both oxidative modifications are present in titin simultaneously (Fig. 4C and D). To have a broader view of the extent of titin softening/stiffening induced by redox modifications, we did simulations at a range of peak forces and calculated the ratio of titin steady-state peak lengths between the oxidized and the reduced conditions. Results show that the softening effect of S-thiolation remains fairly constant, whereas at 50–80 pN peak force, the contribution of disulfides transitions from stiffening to softening (Fig. 4E, Supplementary Fig. S4F). This dual behavior stems from the fact that disulfides favor mechanical unfolding of Ig domains (softening effect), while also reducing the contour length of unfolded domains and increasing folding rates (stiffening effects). At low peak forces in which Ig domain unfolding rates are low, the softening effect is more prominent; while at high peak forces in which Ig unfolding is more frequent, the stiffening effects prevail. Beyond modulation of steady-state titin stiffness, our simulations also illustrate that both disulfides and S-thiolations induce a more dynamic state of titin by favoring Ig domain unfolding reactions, particularly at low forces (Fig. 4F, Supplementary Fig. S4G).

In the human heart, the short N2B isoform is expressed to higher levels than the longer and softer N2BA [1,39]. Since alternative splicing occurs at the region of titin with the highest density of cysteines (Fig. 1D), we also ran Monte Carlo simulations for the N2B isoform of titin. In contrast to the results obtained with N2BA, we find that disulfides do not induce softening of N2B titin at any peak force (Fig. 4G, Supplementary Fig. S4H-N). Interestingly, S-thiolation softens N2B titin to a greater extent (20% vs. 10% for N2BA titin at 50 pN peak force, Fig. 4E,G), reflecting the higher density of S-thiolation-competent Ig domains in N2B (Supplementary Table S2). For the same reason, the extent of modulation of titin dynamics by redox modifications is also different in N2B and N2BA titins (Fig. 4H, Supplementary Fig. S4O), although in both isoforms oxidations increase protein dynamics by
favoring more Ig domain unfolding.

Our MS results were scarce with regards to the oxidation state of the cysteines in the N2Bus region of titin. Monte Carlo simulations show that potential disulfides in the N2Bus [23] would boost the overall stiffening effect of this redox modification, especially in the short N2B isoform (Supplementary Fig. S4P,Q). Indeed, under conditions in which N2Bus cysteines form disulfides, disulfides always stiffen titin. Taking together all the results from the Monte Carlo simulations, we conclude that disulfides in the N2Bus [23] would boost the overall stiffening potential disulfides in the N2Bus [23]. Monte Carlo simulations show that cysteines in the N2Bus region of titin. Although several in vitro experiments over the last years have shown that cysteine oxidation is a strong modulator of titin mechanics, evidence that such modulation is functionally relevant in vivo has been lacking [17]. Our results showing that titin is oxidized in basal conditions both in mice and in humans, together with the recent observation that murine titin cysteines are particularly sensitive to oxidative stress [18–21], demonstrate the in vivo relevance of these mechanisms of mechanical modulation.

In vitro, conserved cysteines B, F and G can be induced to form disulfide bonds. Establishing a nanomechanical function for such disulfide bonds, which titin is reduced, or oxidized by disulfides, S-thiolation adducts or both (color code is indicated at the bottom; peak forces are indicated in the insets). The parameters of S-thiolations used in these simulations correspond to S-glutathionylations (Supplementary Note S1). Graphs show the average of 10 independent simulations and SD is indicated by shaded areas. For subsequent analyses, we considered times longer than 1 h, in which the length of titin fluctuates around steady state values (region outside the grey shading). E: Ratio of oxidized vs reduced N2BA titin length at different peak forces. F: Ratio of oxidized vs reduced N2BA unfolding events in simulations at different peak forces. G: Ratio of oxidized vs reduced N2B titin length at different peak forces. H: Ratio of oxidized vs reduced N2B unfolding events in simulations at different peak forces. In panels E–H, n = 10 simulations; error bars: SD. In panels E and G, the effect of the Ox/Red length on titin mechanics is indicated on the right. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

3. Discussion

The high nucleophilicity of the thiol side chain makes cysteine the most reactive protein amino acid. In addition, cysteine residues are sensitive to irreversible oxidation, which can result in protein aggregation and degradation [40]. As a consequence, proteins tend to include cysteine only if the associated functional benefits compensate the risks stemming from its peculiar physicochemical properties. Indeed, cysteine is among the least frequent but most conserved residues in proteins [41], reflecting key functional roles including configuration of enzyme active sites or metal chelation sites [42–44], establishment of disulfide bonds [38], deployment of electron transport systems [45], or as redox sensors [46]. Although several in vitro experiments over the last years have shown that cysteine oxidation is a strong modulator of titin mechanics, evidence that such modulation is functionally relevant in vivo has been lacking [17]. Our results showing that titin is oxidized in basal conditions both in mice and in humans, together with the recent observation that murine titin cysteines are particularly sensitive to oxidative stress challenges [20], demonstrate the in vivo relevance of these mechanisms of mechanical modulation.
Carlo simulations have allowed us to explore the range of oxidative mechanical modulation of titin (Fig. 4). The simulations suggest that the extent and direction of mechanical modulation depends on the specific titin isoform, the biochemical nature of the oxidative modification (disulfide vs S-thiolation, as nicely demonstrated experimentally in Ref. [20]) and the range of mechanical load experienced by titin. Two observations stemming from the Monte Carlo simulations are relevant for the redox modulation of titin mechanics at physiologically relevant forces, which are generally assumed to be no higher than 10 pN/titin molecule [15,54–56]. First, due mainly to increased Ig unfolding rates, oxidations render the titin filament much more dynamic, up to one order of magnitude at low forces (Fig. 4F,H). More frequent Ig unfolding can change the landscape of interactors and prime residues for phosphorylation [20], resulting in modulation of titin-based mechanosignaling [1], whereas high folding rates enabled by disulfide bonds can sustain titin’s contribution to active muscle contraction [33,54,57]. Monte Carlo simulations also illustrate that disulfides can result both in stiffening or softening of titin (Fig. 4E,G), a consequence of the opposing mechanical effects of disulfides on contour length and the (un)folding rates of Ig domains. This observation, together with the softening effects of S-thiolation, can reconcile seemingly contradictory results of experiments using muscle preparations. For instance, treatment of human cardiomyocytes with the reducing enzyme thioredoxin results in drops in passive tension during oscillatory changes in length [23] while incubation with DTT results in increased passive tension during stepwise length increases [19]. Similar DTT incubations also increase passive tension of rat and human skeletal fibers [29]. Regarding oxidative modifications, treatment of mouse cardiac muscle with H$_2$O$_2$ induces increased passive tension in oscillatory protocols [28], whereas specific S-thiolation reactions lead to softening during stepwise extension of human cardiomyocytes [19,20], an effect also observed in human and rat skeletal muscle [29]. Interestingly, specific induction of disulfides by treatment with protein disulfide isomerase leads to myocyte stiffening, suggesting that the stiffening effect of disulfides prevails under the experimental conditions tested [20].

Overall, our results show that the redox state of conserved cysteines of the I-band of titin is an important modulator of titin mechanics in basal conditions (Fig. 5). Indeed, our simulations illustrate that the range of titin mechanical modulation achieved by redox posttranslational modifications can be ample, particularly for the induction of more dynamic states of the protein that can influence mechanosignaling cascades. In this context, we speculate that altered redox signaling may cause titin mechanical dysfunction during disease [21,52], for instance contributing to myocardial stiffening following myocardial infarction [2,20].

4. Methods

4.1. Human subject research

Human subject research was carried out in accordance with principles outlined in the Declaration of Helsinki. Procedures for procurement of human left ventricular tissue from 2 non-failing donor hearts that were not used for transplantation were approved by the Ethical Committee of the Medical University of Graz (28-508 ex 15/16) and the Instituto de Salud Carlos III (CEI PI 65.2017-v1). Upon ice-cold cardioplegia, cardiac biopsies were harvested from the left ventricular free wall, quickly frozen in liquid nitrogen and stored at -80°C. Both hearts had echocardiographic evidence of preserved ejection fraction (>50%) and a clinical history that was free of cardiac abnormalities.

4.2. Animal research

All post-mortem animal samples used in this work were from CD1 mice housed and maintained in the animal facility at the CNIC (Madrid, Spain) in accordance with Spanish and European Legislation (Directive 2010/63/EU amended by Regulation EU 2019/1010).

4.3. Cysteine conservation analysis

The longest currently available titin sequences for the 37 species reported in Ref. [22] were aligned using Clustal Omega, with the only exception of chicken, for which no titin sequence with enough coverage was found in Uniprot, GenomeNet or NCBI databases. Sequence identification codes are listed in Supplementary Note S2. Conservation values for all cysteine positions in the human titin sequence (Uniprot Q8WZ42) were obtained as percentages of conservation not considering gaps in the alignment. The alignment and homology models of titin Ig
domains of human titin have been published before [18]; alignment of mouse titin Ig domains appearing in Uniprot was obtained similarly using Clustal Omega (Supplementary Note S3). These Ig domain alignments were used to classify titin cysteines for structural conservation and mass spectrometry analyses.

4.4. In-gel determination of reversibly oxidized thiols

Our protocol for in-gel determination of reversibly oxidized thiols was adapted and optimized from previous reports [22, 32]. Protein extracts were obtained by cryopulverization of myocardial tissue followed by homogenization in sample buffer (50 mM Tris-HCl, 10 mM EDTA, 3% SDS, pH 6.8; 40 μl per mg of tissue) containing 50 mM NEM, unless indicated otherwise. Samples were run on SDS-PAGE gels (3.5% for analysis of titin, 12% for other proteins) in the absence of reducing agents. Following electrophoresis, oxidized thiols were reduced by incubation of the gel with 10 mM DL-dithiothreitol (DTT) (Sigma-Aldrich) in 50 mM ammonium bicarbonate, pH 8.8 at 60 °C. After 3 washes of 20 min with sample buffer, the newly reduced thiols were labeled by incubation with 5 mM mBBr (Merck Millipore) in sample buffer during 2 h at room temperature in the dark. The excess of mBBr was removed by three washes with destaining solution (40% ethanol, 10% acetic acid) lasting 1 h, overnight and again 1 h. Fluorescent bands were visualized using a Gel-Doc (BioRad) with UV excitation (standard filters for ethidium bromide). Coomassie staining of the same gel was used to normalize fluorescence signals. Quantification of the bands was done by densitometry using Quantity One. Fluorescence signals coming from a replicate gel not treated with DTT were subtracted (Supplementary Fig. S2B). When comparing oxidation of titin and myosin, we normalized the oxidation signals by their density of cysteines (Uniprot entries A2ASS6 and Q02566 for mouse cardiac titin and myosin, respectively. All gels in a comparative quantification were analyzed from the same unprocessed GelDoc image.

4.5. Mass spectrometry

Titin bands were sliced from SDS-PAGE gels used for fluorescence quantifications, dried and washed by incubation with 200 μl high purity water (Fluka CHROMASOLV™ LC-MS) for 10 min at 1200 rpm (4 times). The resulting gel pieces were dehydrated by 2 incubations with 100% acetonitrile and 1 incubation with 50 mM ammonium bicarbonate (pH 8.8) in acetonitrile (both incubations, 15 min shaking at 1200 rpm). After drying using a SpeedVac, the gel pieces were incubated with modified trypsin from porcine pancreas (Sigma Aldrigh) in digestion buffer (10% acetonitrile in 50 mM ammonium bicarbonate, pH 8.8) for 2 h in ice to allow the diffusion of the inactive enzyme, and then at 37 °C overnight. Digested tryptic peptides were extracted with 1% trifluoroacetic acid in acetonitrile (15 min incubation, 1200 rpm shaking) and dried. Finally, they were resuspended in 1% trifluoroacetic acid by pulse-vortexing and sonication, and desalted using OMIX commercial columns (Biomaster group). Peptides were then injected into a reversed phase C-18 nano-column (Acclaim PepMap RSLC, 75 μm internal diameter and 50 cm length), and eluted to be analyzed in a hybrid quadrupole-Orbitrap Q Exactive mass spectrometer (Thermo Scientific) for protein identification. A continuous acetonitrile gradient consisting of 0–30% A for 120 min, 50–90% B for 5 min (A = 0.1% formic acid; B = 98% acetonitrile, 0.1% formic acid) at a flow rate of 200 nl/min was used to elute tryptic peptides from the nano-column to a nanospray emitter for real time ionization and induced fragmentation. High resolution mass spectra were acquired in a data-dependent manner with dynamic exclusion by combining a MS spectrum (from 400 to 1500 m/z, 120,000 resolution) followed by the MS/MS spectra (60,000 resolution) from the 15 most intense species. For protein identification, tandem mass spectra were extracted and charge state was deconvoluted by Proteome Discoverer 1.4.0.288 (Thermo Fisher Scientific). All MS/MS spectra were analyzed using SEQUEST (Thermo Fisher Scientific). Full proteome search databases (mouse: UniProtKB/Swiss-Prot April27_2016, 48736 sequences; human: UniProtKB/Swiss-Prot, November 2019, 74333 sequences) were supplemented with 116 cRAP proteins (common Repository of Adventitious Proteins, Global Proteome Machine). SEQUEST searches allowed two missed cleavages and used 20 ppm and 20 mA precursor and fragment mass tolerances, respectively. NEM (+125.047679 Da)- or mBBr (+190.074228 Da)-modified cysteines, and oxidation of methionine were specified as variable modifications. In these initial searches, a target-decoy fixed value PSM validation strategy was used to filter peptides according to XCorr values (more than 1.2 in doubly-charged peptides, or more than 1.4 in triply-charged peptides for <5% FDR, and more than 1.42 in doubly-charged peptides, or more than 1.79 for triply-charged peptides for <1% FDR). Since the vast majority of the identified peptides belonged to titin in searches against full proteome databases, to optimize coverage of cysteine peptides we repeated the search using titin-only reduced search databases [36]. Peptide identifications were validated using Vseq, an in-house developed tool that evaluates the mass tolerance, intensity, fragmentation goodness and quantitative value of MS/MS scans [37]. Maximum matched ions value for a sequence was calculated considering the number of residues multiplied by 2, to cover the complete theoretical length of the main fragmentation series B and Y. Since experiments were acquired using a label-free strategy and using HCD for ion-induced dissociation, we set a threshold of 25% minimum matched ions, which corresponds to at least 50% of the Y-HCD-enhanced fragmentation series. We also filtered peptide identifications according to their E-scores, which is defined as the dot product of the intensities for all matched ions regardless of their charge state. We applied an E-score threshold of >0.01. We verified that none of the unidentified scans showing a very low XCorr and matched ions value was above this limit. To estimate final FDR values, we repeated the searches considering the same parameters against the corresponding decoy databases. FDR was below 2% in all cases. To minimize false positive detection of oxidized cysteines, we analyzed in parallel the same samples without including DTT in the derivatization reactions. For each sample, cysteine positions belonging to mBBr-peptides identified both in the −DTT and +DTT treated specimens were excluded from the analysis (27 ± 5% and 30 ± 4% of the oxidized cysteine-containing peptides identified in mouse and in human).

In order to semi-quantitatively describe the landscape of identified cysteine redox states, we calculated a Redox Identification Index (RII) for each cysteine as follows:

$$RII = \frac{1-(#Red)}{#Replicates} + 1-(#Ox) = 0-(#Ox&Red)$$

(1)

where #Ox and #Red are the number of replicates in which the cysteine is detected as oxidized or reduced, respectively. #Ox&Red is the number of replicates in which the cysteine is detected in both oxidized and reduced forms and #Replicates is the total number of replicates in which the cysteine is detected. As NEM- and mBBr-modified peptides might be detected with different efficiencies, due to different ionization and sampling during MS fragmentation, we followed a restrictive approach that simply took into account whether each cysteine was detected modified with NEM and/or mBBr in each replicate sample. Therefore, RII can take ±1, ±2/3, ±1/3 and 0 values for cysteines identified in three replicates; ±1, ±1/2, and 0 for cysteines identified in two replicates; and ±1 and 0 for cysteines identified in one replicate. Although the coarseness of RII values may lead to dispersion of experimental data due to missing identifications, we expect this effect to be averaged out when RII from multiple cysteine positions are analyzed together (see Fig. 3). Indeed, we have calculated the global RII for each cysteine conservation category by counting and comparing all mBBr- and NEM-modified peptides of all cysteines belonging to each conservation category, a procedure that results in RII values that are very similar to the ones obtained by averaging individual RII values (Supplementary Table 1).

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raw data used for cysteine validation and RII calculations can be found in Supplementary Files S2 and S3.

4.6. Monte Carlo simulations

We constructed a simplified model of titin’s I band to predict the length of the molecule under force upon redox modifications using Monte Carlo simulations. Simulations calculate the length of the two entropic regions of the protein (PEVK and N2Bus) and the Ig domains, according to the Freely Jointed Chain model of polymer elasticity and the Bell’s model of force-dependent reactions. The length of the entropic regions and the number of Ig domains in each isoform were determined according to human titin sequence in Uniprot, further curated in Ref. [18]. Titin domains were classified according to their cysteine content and arrangement, which was used to determine the modifications they could be target of (Supplementary Table S2) and their mechanical and kinetic parameters (Supplementary Tables S3–S6). The constructed model was subject to an oscillating force protocol (triangular waves, 1 Hz, 10 ms simulation step) for 3 h. Pilot simulations were also conducted using 1 ms simulation steps, leading to equivalent results. In the initial state all domains are folded. If oxidative modifications are present, we consider they occur at all potential target sites. In conditions involving disulfide bonds, all domains capable of disulfide isomerization contain the BG disulfide in the initial state. We averaged 10 independent simulations for each condition. The code for the simulations and the subsequent analysis of the data was written in Igor Pro (Wavemetrics).

4.7. Statistics

Statistical tests were run using Graph Pad Prism 8.3.1 for Windows. To evaluate the tendency of different categories of cysteines to be oxidized (RII > 0), we estimated the corresponding p-values using the hypergeometric distribution function implemented in Excel (Microsoft). Errors are given by SEM, unless indicated otherwise.

Author contributions

EHG and JAC conceived and led the project. EHG, IMM, CSC, MRP, DG and RPJ analyzed titin sequence. EHG, IMM, CSG, NV, DVC and CBC did in-gel determination of reversible cysteine oxidations. EHG, IMM, DG and RPJ analyzed titin sequence. EHG, IMM, CSC, MRP, DG and RPJ analyzed titin sequence. EHG, IMM, CSG, NV, DVC and CBC did in-gel determination of reversible cysteine oxidations. EHG, IMM, EBK, EC and JD did and analyzed mass spectrometry experiments. MA, SS, and PPR procured and characterized human heart samples. EHG, AFT, JD and IMM did statistical analysis of experimental data. IMM did Monte Carlo simulations. EHG, IMM and JAC drafted the manuscript with input from all authors.

Data and material availability

Data and materials are available on reasonable request to the corresponding authors. Mass spectrometry raw data have been deposited in PeptideAtlas (http://www.peptideatlas.org/, Identifier: PASS01609).

List of supplementary materials

Supplementary Files S1–S3 and Supplementary Information File including Supplementary Figs. S1–S4, Supplementary Notes S1–S3, Supplementary Tables S1–S6 and Supplementary References.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have influenced the work reported in this paper.

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Appendix A. Supplementary data

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