Molecular basis of a novel renal amyloidosis due to N184K gelsolin variant

Francesco Bonì1,2, Mario Milani1,2, Riccardo Porcari3, Alberto Barbiroli4, Stefano Ricagno2 & Matteo de Rosa1,2

Mutations in gelsolin are responsible for a systemic amyloidosis first described in 1969. Until recently, the disease was associated with two substitutions of the same residue, leading to the loss of the calcium binding site. Novel interest arose in 2014 when the N184K variant of the protein was identified as the etiological agent of a novel kidney-localized amyloidosis. Here we provide a first rationale for N184K pathogenicity. We show that the mutation induces a destabilization of gelsolin second domain, without compromising its calcium binding capacity. X-ray data combined with molecular dynamics simulations demonstrates that the primary source of the destabilization is a loss of connectivity in proximity of the metal. Such rearrangement of the H-bond network does not have a major impact on the overall fold of the domain, nevertheless, it increases the flexibility of a stretch of the protein, which is consequently processed by furin protease. Overall our data suggest that the N184K variant is subjected to the same aberrant proteolytic events responsible for the formation of amyloidogenic fragments in the previously characterized mutants. At the same time our data suggest that a broader number of mutations, unrelated to the metal binding site, can lead to a pathogenic phenotype.

Gelsolin is a 80kDa protein displaying six homologous, gelsolin-like, domains (named G1 to G6) existing as two alternative splicing variants: a 24-residue shorter intracellular form and a variant secreted in the bloodstream1,2. Both isoforms bind actin fibrils, severing the filaments and preventing repolymerization. Gelsolin is the prototype of the gelsolin superfamily, whose members are all multidomain proteins, highly regulated by calcium. Gelsolin, in particular, hosts eight calcium binding sites: one for each domain and two incomplete sites, which require actin contribution. In the absence of the ion, gelsolin assumes a compact conformation, in which domains G1–G5 are wrapped around G62, masking the actin binding sites (inactive conformation). Upon increase of the ion concentration, gelsolin undergoes major conformational changes, with large interdomain rearrangements. Such molecular gymnastic is believed to happen in a sequential and cooperative fashion, ultimately leading to the open and active configuration. Active gelsolin eluded structural investigations, most probably due to its highly dynamic nature. In fact, recent studies on adseverin showed that the active conformation of gelsolin-like proteins is better described by an ensemble of structures, characterized by a larger gyration radius with respect to the inactive form3. Upon binding to actin, gelsolin adopts a novel conformation2,4, although the structures of the individual domains closely match those observed in the inactive form: new interdomain contacts are formed while others are lost2,4.

Due to its polyvalent function, gelsolin is somehow involved in a plethora of pathological processes, in particular various cancer types5. Moreover, mutations in the gelsolin gene are directly responsible for gelsolin amyloidosis (GA) with 100% penetrance. GA was first described in 19696 and for long referred as familial amyloidosis Finnish-type (FAF), characterized by deposition of amyloids in different organs and tissues. The parts of the body mostly affected are the eyes, the cranial and peripheral nerves, the skin and, at later stages, the kidneys. Such heterogeneous spectrum of symptoms makes GA very difficult to diagnose.

1CNR Istituto di Biofisica, c/o Dipartimento di Bioscienze, Università degli Studi di Milano, 20133 Milano, Italy. 2Dipartimento di Bioscienze, Università degli Studi di Milano, 20133 Milano, Italy. 3Wolfson Drug Discovery Unit, Centre for Amyloidosis and Acute Phase Proteins, University College London, London NW3 2PF, UK. 4Dipartimento di Scienze per gli Alimenti, la Nutrizione e l’Ambiente, Università degli Studi di Milano, 20133 Milano, Italy. Correspondence and requests for materials should be addressed to M.M. (email: mario.milani@unimi.it) or S.R. (email: stefano.ricagno@unimi.it) or M.D.R. (email: teo.derosa@gmail.com)
Measurements were performed in the presence of either 1 mM calcium chloride (+Ca) or 1 mM EDTA (−Ca).

|          | G2l wt Tm (°C) | G2l D187N Tm (°C) | G2l N184K Tm (°C) | G2s N184K Tm (°C) |
|----------|----------------|-------------------|-------------------|-------------------|
| +Ca      | 41.5           | 46.4              | 33.4              | 33.8              |
| −Ca      | 46.5           | 45.6              | 33.8              |                   |
| ΔTm      | 19             | 0                 | 12                | 13                |

Table 1. Summary of the thermal stability experiments. Tm values (±0.3°C) are reported for the G2l construct of the wt protein and mutants D187N and N184K and the short construct of mutant N184K. Measurements were performed in the presence of either 1 mM calcium chloride (+Ca) or 1 mM EDTA (−Ca).

Until recently, only two mutations were described leading to GA (D187N and D187Y). The mechanism of amyloid formation by these variants is relatively well understood, although it awaits structural confirmation. D187 is part of a cluster of residues binding the calcium ion in the G2 domain. Substitutions in such position with either N or Y impair calcium binding, causing destabilization of the G2 domain. Whether this destabilization is due to a conformational change of the native state or to an increased flexibility of the overall fold is not known, as the structure of the mutated proteins have not been reported. In either case, in the trans-Golgi network (TGN), exported gelsolin encounters calcium at millimolar concentration, sufficient for the activation and opening of the protein structure. In the presence of a destabilizing mutation, protein activation leads to the exposure of the region comprising residues 168 to 173, and the subsequent proteolysis at R172 by the endogenous protease furin. The C-terminal fragment, known as C68, is exported in the extracellular environment, where it is subjected to further aberrant proteolytic events, eventually leading to the formation of 5 and 8 kDa peptides, which are the major constituent of aggregates in GA. Such peptides harbours the mutation in position 187 and are more prone to amyloid aggregation than homologous wt fragments: that is, D187N/Y substitutions play a dual role in GA pathological mechanism, both promoting aberrant proteolysis and increasing gelsolin inherent amyloidogenicity.

Recently, other two gelsolin mutants (G167R and N184K) have been identified as associated with pathologic amyloid deposition. To date the deposition of such mutants seems to cause solely kidney symptoms, however, only a partial clinical picture is available and degree of penetrance of these mutations unknown. These mutations fall in the G2 domain of gelsolin and are in close proximity to the calcium binding site.

Here we report a structural and functional characterization of the isolated G2 domain of the N184K variant. We show that the mutation induces a destabilization of the protein fold without compromising the calcium binding site. Such destabilization confers high susceptibility to furin proteolysis, suggesting that the N184K variant is subjected to the same aberrant proteolytic cascade described for the mutants in position 187.

**Results**

Although the isolated G2 domain is commonly accepted as a valuable system to study the effects of the pathologic mutations on the structure and function of the protein, the definition of its exact boundaries has been subjected to extensive discussion. Two groups, respectively coordinated by A.R. Fersht and J.W. Kelly, mainly contributed to the dissection of the molecular bases of GA. The former group worked on a shorter construct, whose design was based on limited proteolysis studies, hereafter named G2s. The latter used a longer construct suggested by the crystal structure of the Ca-free full-length protein (G2l) (see Supplementary Fig. S1). In our work, we used the G2l construct for all the experiments in solution, nevertheless crystals grew only for the G2s, consequently, when necessary, experiments were repeated on the shorter construct to validate the structural models.

**N184K mutation destabilizes gelsolin G2 fold without compromising calcium binding.**

Previously characterized D187N/Y mutations strongly destabilize gelsolin second domain. Such destabilization is primarily due to the calcium binding impairment, which has an important role in gelsolin activation and structural stability of the individual domains.

A preliminary evaluation of N184K G2l thermal stability was assessed by denaturation experiments in the presence of SYPRO orange fluorescent probe (see Supplementary Fig. S2) to find suitable conditions for its biochemical and structural characterization. These tests suggested that a higher ionic strength was required and that the optimal conditions for the G2l mutants were 20 mM HEPES, pH 7.0, 300 mM NaCl.

Protein unfolding was monitored by CD in the far UV region in the presence of a molar excess of either CaCl₂ or EDTA. Our results on wt and D187N G2l well reproduce those reported by Huff et al. Indeed, the wt (Tm = 60.6°C) is strongly stabilized by the presence of calcium (ΔTm = +19°C), whereas the mutant shows Tm values very similar to the wt apoprotein (Table 1 and Fig. 1).

Surprisingly, N184K G2l variant exhibits a shift of the Tm value upon addition of a molar excess of EDTA, indicating that N184K is still able to bind Ca²⁺ and that the ion provides a significant stabilization (ΔTm = +12°C) of the domain. Nevertheless, the mutation destabilizes the G2 fold: The Tm of N184K is 45.6°C, 15°C lower than the Tm measured for the wt domain.

CD spectra and temperature ramps were recorded for both G2l and G2s N184K constructs. G2l and G2s CD spectra are almost identical and Tms are calculated in the presence of either calcium or EDTA (47.0 and 33.8°C, respectively) are within the experimental error, suggesting that the N-terminal extension does not influence the stability of the domain nor induces detectable changes at the secondary structure level.

**N184K mutation increases gelsolin G2 susceptibility to furin proteolysis but it does not confer intrinsic amyloidogenicity.**

D187N/Y mutations increase isolated G2 domain sensitivity to proteolysis by...
furin, which is believed to be the first event leading to gelsolin amyloidosis. Here, we wanted to test if the mutation in position 184 was conferring similar susceptibility. Such tests were performed at pH 6.5 and in the presence of 1 mM CaCl₂ to mimic conditions found in the TGN and because furin is a calcium dependent protease. The reaction was followed for roughly 16 hours and samples periodically collected to define a time course of the process. N184K G2l mutant displays high susceptibility to furin aberrant proteolysis: bands corresponding to 5 and 10 kDa fragments appear after only 1 h incubation and the G2l domain is totally processed after 16 hours. Contrary, wt G2l domain is extremely resistant to proteolysis under the tested experimental conditions (Fig. 2, top panels).

The experiment was also repeated on the wt and N184K G2s' (Fig. 2, bottom panels). These constructs, which lack a short edge strand interacting with the β1 strand, show a pattern of proteolysis very similar to that observed for the longer forms. During the proteolysis of wt domain some faint bands, possibly product of furin activity, can be observed but their intensity is not comparable with the ones visible in the experiments with the mutant. These results suggest that the N-terminal stretch does not represent a crucial protective element from aberrant proteolysis nor, in keeping with thermal stability data, has a major impact on the fold of the domain.

In order to assess if the observed proteolytic event is due to the increased mobility of the mutated domain, we used wt and N184K G2l variants in limited proteolysis experiments with an unspecific protease. We chose trypsin for this analysis because G2 domain is particularly rich in positively charged residues, (Arg + Lys make up for more than 10% of the sequence) and a new potential cleavage site is created by the mutation itself. Nevertheless, the two variants show both an identical proteolytic pattern and a comparable kinetics of the intact domain (up to 20'), with a rapid appearance of two bands relatively stable over time (see Supplementary Fig. S3). After these early proteolytic events, further degradation appears to progress faster for the N184K fragments. This experiment further strengthens the results shown in Fig. 2, suggesting that furin cleavage is a very specific event due to an inherent susceptibility conferred by the mutation.

Fibrillogenesis of wt and N184K G2l variant was carried out at both pH 5.5 and 2.5, monitoring thioflavin T fluorescence. In the former condition, no aggregation occurs for either the wt or the mutant domain. On the
contrary, at pH 2.5, wt G2l shows a higher propensity for amyloidogenesis than the variant (Fig. 3a). Secondary structure content was monitored during the experimental time frame by CD spectroscopy (Fig. 3b,c). At lower pH the two variants show a spectrum consistent with a completely unfolded protein. Conversely at pH 5.5, the spectrum of wt is comparable to those registered at physiological conditions, whereas, the N184K mutant is partially denatured. We can therefore infer that the mutation does not increase the intrinsic propensity of gelsolin to aggregate under both denaturing and milder conditions.

Structural bases of G2 N184K instability and susceptibility to aberrant proteolysis.

To decipher the structural basis of N184K sensitivity to furin proteolysis, its structure was determined by X-ray crystallography. Although extensive crystal screenings were performed on N184K G2l construct, we failed to obtain crystals of suitable quality. As literature data and our limited proteolysis experiments suggest, the flexible N-terminal loop could prevent the growth of well-ordered crystals, thus we resorted on the use of the shorter G2s construct, lacking this N-terminal stretch.

Crystals of N184K G2s displaying two different morphologies readily appeared in different conditions. Best diffracting crystals were used to determine N184K structure at 1.05 and 1.7 Å resolution, which will be hereafter referred as the orthorhombic and trigonal forms, respectively (Table 2). Electron density was of excellent quality for both structures and allowed the tracing of all residues belonging to the core of the G2 domain. Only few amino acids were not modeled: residues 260–266 at the N-terminus of the orthorhombic crystal form, and residues 256–266, 158–159 and the side chain of 169 in the trigonal crystal. In both crystal forms the prototypical gelsolin-like fold, made of a 5-strand β-sheet sandwiched between two α-helices is maintained.

In agreement with denaturation studies, electron density for Ca2+ was clearly visible in its binding site and the ion was refined to 100% occupancy in both structures (Figs 4 and S4). In addition, other ions were found coordinated between symmetric molecules, likely contributing to the formation of the tight crystal packing, which lead to the high resolution diffraction achieved but irrelevant for the structure of G2.

Beside the C-terminal tail, the two N184K crystal forms match each other very closely, as highlighted by a Cα r.m.s.d. value of 0.25 Å (residues 160–250). A similar Cα r.m.s.d. value (0.56 Å for 102 atoms) can be calculated between the mutant in the close conformation and the wt structure.

The open conformation of the C-terminus, observed in the trigonal crystal, is likely due to crystal packing: the terminal tail interacts with residues of the N-terminus and of the β3-β4 loop belonging to a symmetric molecule
Therefore, we focused on the orthorhombic form. In this structure the mutation seems to have only a minor impact on the overall structure of the G2 domain. Interestingly, the experimental electron density of the β1-β2 turn is of lower quality with respect to the rest of the protein (Fig. S4), displaying B factors significantly above the average. Such loop (residues 167–170) is part of the aberrant proteolytic site (168-RRVVR↓A-173) and its enhanced motility could explain the higher susceptibility to furin proteolysis displayed by the mutant.

Another area of the protein characterized by relatively high B factors and poor electron density is the hinge region following the α2 helix. N184 does not directly coordinate the calcium ion, nevertheless it is hydrogen bonded to D187 (Fig. 5), the residue mutated in FAF, whose substitution to either N or Y leads to the loss of calcium binding capability. In the variant object of this study, not only the calcium is firmly bound but also the geometry of the coordination site matches perfectly the one observed in the wt structure. In the mutant, N184 is substituted with a bulkier lysine, which does not fit in the pocket occupied by the asparagine. As a result, the long flexible side-chain of lysine protrudes toward the solvent, without causing a significant rearrangement of the neighboring residues. Nevertheless, the mutation induces a weakening of a H-bond network around the Ca\(^{2+}\) coordination site involving mainly four residues: Q164, N184, G186 and the Ca\(^{2+}\) bound D187. In details, N184, in the wt protein, is H-bonded to the N atoms of both G186 and D187 and to one of the Oδ atoms of the latter (Fig. 5). All the above mentioned interactions are lost in the mutant. In addition, two longer-range polar contacts can be observed between Q164 and both D187 and N184 in the wt protein. In the mutant, Q164 is present in two alternative conformations correlated with the two conformations of T174 (refined to the same occupancy; Fig. 5): when Q164 loses the interaction with D187 it is H bonded to T174.

In the orthorhombic structure an interesting gain of motility in a region close to the furin cleavage site was observed. In order to better characterize it, we decided to perform two molecular dynamics simulations on the wt and the N184K mutant. To minimize all other possible effects except those induced by the mutation we chose to start both simulations from the same coordinates of the orthorhombic crystal structure, after the addition of the missing terminal amino acids (modeled in random conformation) with either asparagine or lysine in position 184. Overall the mutated protein displays an enhanced motility during the simulation as shown by the C\(_{\alpha}\) root mean square fluctuation (RMSF, i.e. standard deviation) of atomic positions along the trajectory (see Figure 3. N184K mutation does not increase G2 inherent propensity to form fibrils. (a) Aggregation kinetics monitored by the increase of ThT fluorescence of wt (black traces) and N184K (blue traces) G2l variants at both pH 2.5 (dashed lines) and 5.5 (solid lines). (b,c) CD spectra of wt (left panel) and N184K mutant (right panel) G2l in the fibrillogenesis conditions compared to the spectrum at physiological pH.
Supplementary Fig. S4). Consistently with the crystallographic analysis, the side chain of K184 points toward the solvent during all the 40 ns simulation, allowing a greater conformational freedom for Q164 with respect to the wt protein (Fig. 6A). Accordingly, in the mutant protein Q164 is involved in H-bond with D187 only in 10% of the simulation time (the mean distance between the two residues being 4.36 Å) whereas in the wt the two amino acids remain closer (mean distance = 3.67 Å) establishing H-bond for about 54% of the simulation time (Fig. 6B).

Table 2. Data collection and refinement statistics. + and § Values in parentheses refer to the highest resolution shells. *Rwork = Σhkli||Fo||−|Fc||/Σhkli||Fo|| for all data, except 5–10%, which were used for Rfree calculation. #Average temperature factors over the whole structure.

Figure 4. Impact of N184K mutation on the overall fold of gelsolin G2 domain. (a) Superimposition of the crystal structures of N184K (orthorhombic and trigonal crystal forms, coloured in cyan and yellow, respectively) and wt protein (pdb id 1KCQ, in light gray). Residue 184 is shown as sticks and labeled; N- and C-termini, the calcium ion and the aberrant proteolytic site are also marked. (b,c) Details of the calcium binding site in the two N184K structures; the 2Fo-Fc electron density map of the residues and the water molecules coordinating the ion are shown as a grey grid countered at 1.5 σ.
The different conformation of Q164 directly affects the dihedral of T174 side chain (Fig. 6C). In conclusion, the overall effect of the N184K mutation is transmitted by Q164 toward the furin cleavage site, causing a greater conformational freedom of $\beta_1$-$\beta_2$ loop respect to the wt (Fig. S4). Such distinct dynamical features might be the key factors in promoting the efficient furin binding and processing.

**Discussion**

Gelsolin amyloidosis is a relatively rare genetic disease caused by accumulation of aggregated proteolytic fragments of the protein gelsolin. The Finnish form of this malady (FAF) has been known for long time and it is due to mutation of aspartate 187 in either asparagine or tyrosine in the G2 domain of the protein. These variants are unable to bind calcium, which plays a stabilizing role, resulting in a misfolded protein prone to aberrant proteolysis.

Recent development of diagnostic tools for the identification and typing of amyloidosis\textsuperscript{17} promptly led to the identification of two novel variants of gelsolin carrying amyloidogenic mutations\textsuperscript{14,15}: N184K and G167R, suggesting that GA might be more common than it is believed to date. Interestingly, both mutations fall in the G2 domain and are sited in close proximity to the calcium binding site. Amyloid fibers arising from these variants, seem to accumulate solely in the kidney, whereas mutants in position 187 lead to a more systemic amyloidosis characterized by the symptomatic triad: corneal lattice dystrophy, peripheral polyneuropathy and cutis laxa. The determinants of kidney localized deposition are not known nor whether is the full-length protein or fragments forming the fibrillar tangles, as only a partial clinical description of these variants is available so far. For these reasons we focused on the molecular characterization of N184K G2 domain, to gain insights into the pathogenicity of this variant.

Our results show that N184K severely affect gelsolin stability, as it is the case of FAF and many other monogenic diseases\textsuperscript{16,18}. At the same time the mechanism behind such destabilization is subtler. In fact, the N184K variant is still able to bind calcium and the ion affects the domain stability in a similar way as observed for the wt. This result is of a pivotal importance, suggesting that a broader number of potential mutations may lead to a pathogenic phenotype. Interestingly, the primary source of the N184K destabilization is a rearrangement of the H-bond network around the calcium binding site. K184 side chain is found extruded toward the solvent in the crystal structures and throughout the MD simulation, consistent with a permanent loss of the interactions that in the wt domain involve the lost asparagine. At the same time, the mutation, although without a major impact on the fold, affects the overall motility of the domain and in particular that of the $\beta_1$-$\beta_2$ loop (some 20 Å afar). Such increased dynamicity is observed both in the crystallographic structure, in terms of higher average B factors, and in the simulations monitoring the r.m.s.f. values (Fig. S4). The effects of the mutation are transmitted toward the $\beta_1$-$\beta_2$ loop by Q164 side chain (in $\beta_1$), that loosing the permanent H-bonds with D187 and N184, can oscillate between two conformations (Figs 5 and 6A). In one of the two conformations Q164 interacts with T174 (in $\beta_2$) downward to the furin cleavage site (168-RRVVR↓A-173). Interestingly, even if in both wt and N184K mutant the region around $\beta_1$-$\beta_2$ loop displays higher B factor values respect to the average, such effect is larger in the mutant: 38.9% growth vs. 28.2%.

Similar results were obtained by NMR and MD studies\textsuperscript{7-9} on the D187N mutant where, in particular in the latter, Kazmiński and co workers not only observed a very mobile $\beta_1$-$\beta_2$ loop but even the loss of structure of the flanking strands by the end of the MD simulation\textsuperscript{7}. Such destabilization was mainly caused by the loss of the salt

---

**Figure 5.** N184K mutation perturbs the H-bond network around Ca$^{2+}$ binding site. N184K structure (orthorhombic crystal; cyan) overlaid onto the wt protein (white). N/K184, the residues forming the calcium binding site, N164 and T174 are shown as sticks and labeled. Polar interactions between N184 and N187, G186 and Q164, which are lost upon mutation, are highlighted by dashed lines as well as the new interaction between Q164 and T174.
bridge between D187 and K166, with the latter residue playing a similar role in bringing the mutation site to the β1-β2 loop, as observed for Q164 in the N184K variant.

In addition, Kazmirski et al. noticed a significant increase in the C-terminal tail dynamics, which represents a protective element from furin proteolysis due to its steric hindrance. In one of our structures, we found such a tail in an open conformation (not observed during our MD simulation), suggesting that even in the N184K mutant this stretch of the protein is more flexible and prone to unfolding. Overall, the comparison of D187N and N184K indicates that both mutations have an impact on the dynamical properties of the protein although to different extents.

Indeed, the aforementioned β1-β2 loop corresponds to the stretch of protein recognized by furin, comprising residues 168–173, and we showed that N184K mutant is promptly processed by the protease and that this event is due to specific recognition. In fact, experiments using trypsin did not show a generic susceptibility to proteolysis of the mutant, if the sensitivity of the intact domain is compared, as the reduced stability measured by thermal unfolding could have suggested. This result is in keeping with the structural analysis where we observe subtle differences only in that specific region of the G2 domain. Although furin assays suggest that N184K variant is subjected to the same proteolytic cascade of the FAF mutants, we tested if the mutation could tune the aggregation propensity of the intact G2 domain. Indeed, the gelsolin mutations in position 187 not only are responsible for the aberrant proteolysis but also the fragments formed, which contain the mutation, are more prone to aggregate compared to an equivalent wt peptide13. Moreover, aggregation efficiency is pH dependent and increases at lower values12. Our fibrillogenesis experiments in both nearly physiological and denaturant conditions showed that, while at pH 5.5 neither the wt protein nor the mutant aggregates, at low pH, when the G2 domain is totally unfolded, aggregation occurs. Unfolding leads to the exposure of gelsolin amyloidogenic core, which has been identified as the peptide spanning residues 179–19419. Interestingly, N184K mutant shows both a later onset and a longer lag phase of fibril formation. Although confirmation would come from repetition of the experiment on the proteolytic fragments, which is beyond the focus of this work, these results show that the N184K mutation does not increase the intrinsic amyloid propensity of the G2 domain. Moreover, our results suggest that the amyloidogenic peptides carrying the N184K mutation would be less prone to aggregation respect to the wt, which well correlates with the milder phenotype of the disease, not systemic but localized in kidneys.

An additional problem we had to face during the characterization of N184K G2 domain was the role of the N-terminal stretch comprising the short edge strand (β0) and the loop connecting the same to β1, or, in other words, which is the best definition of gelsolin domain 2. The issue has been already extensively discussed7,16 but...
what we believe to be the key experiment, i.e. furin proteolysis assay, was never performed in parallel on the two constructs. Our results clearly show that the β strand does not protect G2 domain from aberrant proteolysis. Therefore, the G2s construct is an appropriate and convenient model for the biochemical characterization of gelsolin G2 domain, with the added value to be suitable for crystallography studies.

In conclusion, our results show that the N184K mutation does not have a major impact on the fold of the G2 domain, while it affects the mobility of the β strand and of the loop downstream. Such clear increase of dynamicity promotes the docking of furin protease, which cleaves the protein with comparable efficiency respect to the mutants in position 187\(^{16}\). Although we do not have data in regards, we can speculate that the cleaved fragment would be further proteolysed by enzymes from the extracellular matrix with the formation of the 5 and 8 kDa peptides carrying the N184K mutation. These peptides seem to be less prone to aggregation than those originated from the 187 mutations, leading to a milder phenotype that causes a localized deposition and not a systemic disease. In this context the N184K mutation could be nearly considered a risk factor rather than a pathogenic mutation, but more clinical and molecular data are required to have a complete picture of the disease.

Methods

**Protein production.**  *Cloning, mutagenesis and expression.*  Synthetic gene coding for the second domain of human gelsolin (G2l, residues 133–266) was purchased from Eurofins Genomics and subcloned into a pET-28 plasmid at the Nde I/Xho I sites (Novagen) carrying a 6xHis-tag at the N-terminus. G2l mutants (D187N and N184K) were produced by site-directed mutagenesis using G2l wt as template and the phusion mutagenesis kit (Thermofisher). The shorter construct of N184K (G2s, res 151–266) used for crystallographic studies was also obtained by site-directed mutagenesis according to the two-step protocol described in ref. 20.

Recombinant plasmids (G2l_wt/pET28, G2l_D187N/pET28, G2l_N184K/pET28 and G2s_N184K/pET28) were transformed in the *E. coli* strain BL21 pLys (DE3) (InvitrogenTM). Cell cultures were grown in LB Broth at 37 °C. Expression of G2l and G2s was induced by addition of 1 mM IPTG.

**Purification.**  Typically, 10–20 mg of cells were resuspended in lysis buffer (20 mM sodium phosphate, pH 7.4, 0.5 M NaCl, protease inhibitors (Complete, EDTA-free, Roche), 10 μg/ml of Deoxyribonuclease I) and lysed by sonication. The clarified crude extract was incubated with 2 ml of Ni\(^{2+}\) resin (Ni SepharoseTM 6 Fast Flow, GE Healthcare) for 2 hours at 4 °C, and after extensive washing with the lysis buffer, G2l protein was eluted with 20 mM NaPO\(_4\), 0.5 M NaCl, 0.3 M Imidazole.

After cleavage of the 6xHis-tag (5 U of thrombin/mg of G2 for 1 h at room temperature) the sample was passed through a HiLoad 16/600 SuperdexTM75 column equilibrated with 20 mM HEPES, 300 mM NaCl, pH 7.4.

The N184K G2s variant was further purified on a MonoQ 5/50 GL column equilibrated with 10 mM TrisHCl, 20 mM NaCl, pH 8.5 and eluted with a linear NaCl gradient (from 20 mM to 1 M).

**Thermal stability.**  *Thermofluor assay.*  The experiments were carried out using a MJMini™ Personal Thermal Cycler (BIO-RAD), heating the samples from 10 to 99 °C. Fluorescent Sypro Orange (Sigma) was used to monitor G2l thermal denaturation. 2.5 μl of G2l (final protein concentration 5 mg/ml) were mixed with 3.5 μl of a 1:500 dilution of Sypro Orange commercial solution and 19 μl of either 50 mM Sodium Acetate, pH 5.5 or 50 mM Sodium citrate, pH 2.5. 10 μl of Thioflavin T (ThT) was added to the sample and aliquots of 100 μl each were collected at 0, 60, 180 min and 20 h.

To block the reaction SDS loading buffer 1X (Bio-Rad) +0.7 M β-mercaptoethanol was added to each aliquot which was promptly heated at 90 °C for 3 minutes. Proteolysis reaction was monitored by SDS-PAGE.

**Circular dichroism.**  CD measurements were performed with a J-810 spectropolarimeter (JASCO Corp., Tokyo, Japan) equipped with a Peltier system for temperature control. All measurements were performed in 20 mM HEPES, 100 mM NaCl, pH 7.4 at 0.2 μl/ml protein concentration. All temperature ramps were recorded from 10 to 95 °C (temperature slope 50 °C/hour) in a 0.2 cm path length cuvette and monitored at 218 nm wavelength. Tm was calculated as the minimum of the first-derivative of the traces. Spectra before and after unfolding ramp were recorded (260–190 nm).

**Furin assays.**  Furin cleavage assays were performed using 1 U of commercial furin enzyme (BioLabs) and 1 mg/ml of G2l/G2s variants in 20 mM MES, pH 6.5, 100 mM NaCl, 1 mM CaCl\(_2\). The final volume of each reaction mix was 50 μl and the reaction was incubated at 37 °C. To monitor the reaction, 12 μl aliquots of the reaction mix were collected after 0, 60, 180 min and 20 h.

To block the reaction SDS loading buffer 1X (Bio-Rad) +0.7 M β-mercaptoethanol was added to each aliquot which was promptly heated at 90 °C for 3 minutes. Proteolysis reaction was monitored by SDS-PAGE.

**Limited proteolysis.**  WT and N184K G2l were dissolved at 1 mg/ml in PBS, pH 7.4. Trypsin (PROMEGA) was added to the sample at a concentration of 2 ng/μl at room temperature. Proteolysis samples (10, 20, 30, 60, 90, 120 minutes) were analysed by SDS-homogeneous 15% PAGE (GE Healthcare).

**Fibrillogenesis.**  WT and N184K G2l were diluted 1 mg/ml in either 50 mM Sodium Acetate, pH 5.5 or 50 mM Sodium citrate, pH 2.5. 10 μl of Thioflavin T (ThT) was added to the sample and aliquots of 100 μl each were loaded in Costar 96-well black-wall plates then covered with sealing films. The plates were incubated at 37 °C and subjected to 900 rpm double-orbital shaking using a BMG LABTECH FLUOstar Omega plate reader. Thioflavin T fluorescence emission at 480 nm was monitored by bottom reading using 445 nm as excitation wavelength and fluorescence was monitored in three or more replicates for each well until it reached a plateau.
Crystallization, structure solution and analysis. Crystallization trials were performed at 20 °C using an Oryx-8 crystallization robot (Douglas Instruments) and several commercial screens in a sitting-drop set up: 0.2 or 0.3 μl of 8 mg/ml N184K G2s in 20 mM HEPES, 300 mM NaCl, pH 7.4, supplemented with 5 mM CaCl2 were mixed with 0.2 or 0.1 μl of the reservoir solution, respectively. Best diffracting crystals were obtained in either 200 mM calcium acetate, 100 mM MES pH 6.0, 20% PEG 8000 (orthorhombic form) or 0.2 M lithium sulfate, 0.1 M TrisHCl pH 8.5, 40% PEG 400 (trigonal form). The former was cryoprotected in a reservoir solution supplemented with 20% glycerol before being flash-cooled in liquid nitrogen. X-ray diffraction data were collected at beamline ID29 (ESRF, Grenoble) at 100 K. Data were processed with MOSFLM21 and scaled with SCALA22.

N184K G2s crystal structure was solved by molecular replacement with the program PHASER23, using G2 wt structure (pdb ID 1KCO) as searching model. Structures were refined with both refmac524 and Phenix refine25 and manual model building performed with Coot26. Pymol was used to both analyse the structures and prepare the figures27.

Molecular dynamics simulation. The molecular dynamics simulations were performed using the program NAMD28. Briefly, we started from the orthorhombic structure of N184K and added 6 missing amino acids at the C-term (261–266) in random conformation resulting in a model composed of 118 amino acids (from Gly148 to Ala266) and a Ca2+ ion. In order to produce the model for wt simulation we modified the same pdb introducing the mutation K184N.

Using the program psfgen (part of the namd2 package28) H-atoms were added and partial charges assigned to every atom in the two models. Then for each model a box was built around the protein (67.1 × 69.4 × 66.6 Å3) and filled with 8,892 and 8,896 water molecules in N184K or wt, respectively. The charge equilibration of the two systems was performed adding NaCl 0.2 M (37/36 Cl− and 34/34 Na+ ions, in N184K/wt using the vmd package; http://www.ks.uiuc.edu/Research/vmd/). Initially, harmonic constraints to fix protein atom positions during energy minimization of the solvent were applied for 0.5 ns, before equilibrating the systems at 310 K for additional 0.5 ns. The simulation was then started, and lasted 40 ns, with 2 fs time steps. During the simulation periodic boundary conditions in NPT ensemble, with Langevin temperature control (T = 300 K) and Langevin piston Nose-Hoover method28, were used to maintain constant temperature and pressure, respectively. Van der Waals interactions were handled with a cutoff of 12 Å, and switched off using a smoothing function beyond 10 Å. The electrostatic interactions were treated with the Particle Mesh Ewald method30, using a grid of 100 points along each dimension of the simulation box.

References

1. Sun, H. Q., Yamamoto, M., Mejillano, M. & Yin, H. L. Gelsolin, a multifunctional actin regulatory protein. J Biol Chem 274, 33179–33182 (1999).
2. Nag, S. et al. Ca2+-binding by domain 2 plays a critical role in the activation and stabilization of gelsolin. Proc Natl Acad Sci USA 106, 13713–13718, doi: 10.1073/pnas.0812374106 (2009).
3. Chunnarnsilpa, S., Robinson, S., Grimes, J. M. & Leyrat, C. Calcium-controlled conformational choreography in the N-terminal half of adipin. Nat Commun 6, 8254, doi: 10.1038/ncomms9254 (2015).
4. Choe, H. et al. The calcium activation of gelsolin: insights from the 3A structure of the G4-G6/actin complex. J Mol Biol 324, 691–702 (2002).
5. Li, G. H., Arora, P. D., Chen, Y., McCulloch, C. A. & Liu, P. Multifunctional roles of gelsolin in health and diseases. Med Res Rev 32, 999–1025, doi: 10.1002/med.20231 (2012).
6. Meretoja, J. Familial systemic paramyloidosis with lattice dystrophy of the cornea, progressive cranial neuropathy, skin changes and various internal symptons. A previously unrecognized hereditary disorder. Ann Clin Res 3, 314–324 (1969).
7. Kazmi, S. L. et al. Loss of a metal-binding site in gelsolin leads to familial amyloidosis-Finnish type. Nat Struct Biol 9, 112–116, doi: 10.1038/nsb745nsb745 [pii] (2002).
8. Isaacson, R. L., Weeds, A. G. & Fersht, A. R. Equilibria and kinetics of folding of gelsolin domain 2 and mutants involved in familial amyloidosis-Finnish type. Proc Natl Acad Sci USA 96, 11247–11252 (1999).
9. Kazmi, S. L., Howard, M. J., Isaacson, R. L. & Fersht, A. R. Elucidating the mechanism of familial amyloidosis-Finnish type: NMR studies of human gelsolin domain 2. Proc Natl Acad Sci USA 97, 10706–10711, doi: 10.1073/pnas.180310097180310097 [pii] (2000).
10. Rattanawamy, G., Huff, M. E., Su, A. I., Rion, S. & Kelly, I. W. Destabilization of Ca2+-free gelsolin may not be responsible for proteolysis in Familial Amyloidosis of Finnish Type. Proc Natl Acad Sci USA 98, 2334–2339, doi: 10.1073/pnas.04145259841452598 [pii] (2001).
11. Chen, C. D. et al. Urinary and serum gelsolin in patients with familial amyloidosis in the Golgi through a defect in Ca2+ stabilization. EMBO J 20, 6277–6287, doi: 10.1093/emboj/20.22.6277 (2001).
12. Solomon, J. P. et al. The 8 and 5 kDa fragments of plasma gelsolin form amyloid fibrils by a nucleated polymerization mechanism, while the 68 kDa fragment is not amyloidogenic. Biochemistry 48, 11370–11380, doi: 10.1021/bi901368e (2009).
13. Maury, C. P., Nurmiaho-Lassila, E. L. & Rossi, H. Amyloid fibril formation in gelsolin-derived amyloidosis. Definition of the amyloidogenic and effective acceleration of amyloid formation of mutant Asn-187 and Tyr-187 gelsolin peptides. Lab Invest 70, 558–564 (1994).
14. Sethi, S. et al. Renal amyloidosis associated with a novel sequence variant of gelsolin. Am J Kidney Dis 61, 161–166, doi: S0227-6686(12)01042-6 [pii] 10.1016/j.ajkd.2012.07.016 (2013).
15. Efebera, Y. et al. Novel gelsolin variant as the cause of nephrotic syndrome and renal amyloidosis in a large kindred. Amyloid 21, 110–112, doi: 10.3109/13506129.2014.891502 (2014).
16. Huff, M. E., Page, L. J., Balch, W. E. & Kelly, I. W. Gelsolin domain 2 Ca2+ affinity determines susceptibility to furin proteolysis and familial amyloidosis of Finnish type. J Mol Biol 334, 119–127, doi: S0022283603011665 [pii] (2003).
17. Sethi, S. et al. Laser microdissection and mass spectrometry-based proteomics aids the diagnosis and typing of renal amyloidosis. Kidney Int 82, 226–234, doi: 10.1038/ki.2012.108 (2012).
18. Yue, P., Li, Z. & Moult, J. Loss of protein structure stability as a major causative factor in monogenic disease. J Mol Biol 353, 459–473, doi: 10.1016/j.jmb.2005.08.020 (2005).
19. Maury, C. P., Nurmiaho-Lassila, E. L., Boysen, G. & Liljestrom, M. Fibrillogenesis in gelsolin-related familial amyloidosis. Amyloid 10 Suppl 1, 21–25 (2003).
20. Wang, W. & Malcolm, B. A. Two-stage PCR protocol allowing introduction of multiple mutations, deletions and insertions using QuickChange Site-Directed Mutagenesis. Biotechniques 26, 680–682 (1999).
21. Battye, T. G., Kontogiannis, L., Johnson, O., Powell, H. R. & Leslie, A. G. iMOSFLM: a new graphical interface for diffraction-image processing with MOSFLM. *Acta Crystallogr D Biol Crystallogr* **67**, 271–281, doi: 10.1107/S0907444910048675 (2011).

22. Evans, P. Scaling and assessment of data quality. *Acta Crystallogr D Biol Crystallogr* **62**, 72–82, doi: 10.1107/S0907444905036693 (2006).

23. McCoy, A. J. *et al.* Phaser crystallographic software. *J Appl Crystallogr* **40**, 658–674, doi: 10.1107/S0021889807021206 (2007).

24. Vagin, A. A. *et al.* REFMAC5 dictionary: organization of prior chemical knowledge and guidelines for its use. *Acta Crystallogr D Biol Crystallogr* **60**, 2184–2195, doi: 10.1107/S0907444904023510 (2004).

25. Afonine, P. V. *et al.* Towards automated crystallographic structure refinement with phenix.refine. *Acta Crystallogr D Biol Crystallogr* **68**, 352–367, doi: 10.1107/S0907444912001308 (2012).

26. Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. Features and development of Coot. *Acta Crystallogr D Biol Crystallogr* **66**, 486–501, doi: 10.1107/S0907444910007493 (2010).

27. Schrödinger, L. L. C. *The PyMOL Molecular Graphics System, Version 1.8* (2015).

28. Phillips, J. C. *et al.* Scalable molecular dynamics with NAMD. *J Comput Chem* **26**, 1781–1802, doi: 10.1002/jcc.20289 (2005).

29. Feller, S. E., Zhang, Y. H., Pastor, R. W. & Brooks, B. R. Constant-Pressure Molecular-Dynamics Simulation - the Langevin Piston Method. *J Chem Phys* **103**, 4613–4621, doi: 10.1063/1.470648 (1995).

30. Darden, T., York, D. & Pedersen, L. Particle Mesh Ewald - an N.Log(N) Method for Ewald Sums in Large Systems. *J Chem Phys* **98**, 10089–10092, doi: 10.1063/1.464397 (1993).

Acknowledgements
We thank Prof. Martino Bolognesi and Dr. Eloise Mastrangelo for helpful discussions and critical reading of the manuscript. The financial support of Telethon – Italy (Grant no. GEP15070) is gratefully acknowledged. We acknowledge the European Synchrotron Radiation Facility for provision of synchrotron radiation facilities and we would like to thank ID29 staff members for assistance in using their beamline.

Author Contributions
F.B., R.P., A.B., M.M. and M.d.R. performed the experiments; F.B., M.M. and M.d.R. designed the study; F.B., R.P., A.B., M.M., S.R. and M.d.R. wrote the paper.

Additional Information
Accession codes: Atomic coordinates and structure factors for the N184K variant of gelsolin G2 domain were deposited in the Protein Data Bank, with accession codes 5FAF and 5FAE.

Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Bonì, F. *et al.* Molecular basis of a novel renal amyloidosis due to N184K gelsolin variant. *Sci. Rep.* **6**, 33463; doi: 10.1038/srep33463 (2016).

© The Author(s) 2016