ATP competes with PIP$_2$ for binding to gelsolin

Dávid Szatmári$^{1,2}$, Bo Xue$^{1,3,4}$, Balakrishnan Kannan$^1$, Leslie D. Burtnick$^5$, Beáta Bugyi$^{2,6}$, Miklós Nyitrai$^{2,6}$, Robert C. Robinson$^{1,3,7}$*

$^1$ Institute of Molecular and Cell Biology, A*STAR (Agency for Science, Technology and Research), Singapore, Singapore, $^2$ University of Pécs, Medical School, Department of Biophysics, Pécs, Hungary, $^3$ Department of Biochemistry, National University of Singapore, Singapore, Singapore, $^4$ NUS Synthetic Biology for Clinical and Technological Innovation (SynCTI), Life Sciences Institute, National University of Singapore, Singapore, Singapore, $^5$ Department of Chemistry and Centre for Blood Research, Life Sciences Institute, University of British Columbia, Vancouver, BC, Canada, $^6$ Szentagothai Research Center, Pécs, Hungary, $^7$ Research Institute for Interdisciplinary Science, Okayama University, Okayama, Japan

* rrobinson@imcb.a-star.edu.sg

Abstract

Gelsolin is a severing and capping protein that targets filamentous actin and regulates filament lengths near plasma membranes, contributing to cell movement and plasma membrane morphology. Gelsolin binds to the plasma membrane via phosphatidylinositol 4,5-bisphosphate (PIP$_2$) in a state that cannot cap F-actin, and gelsolin-capped actin filaments are uncapped by PIP$_2$ leading to filament elongation. The process by which gelsolin is removed from PIP$_2$ at the plasma membrane is currently unknown. Gelsolin also binds ATP with unknown function. Here we characterize the role of ATP on PIP$_2$-gelsolin complex dynamics. Fluorophore-labeled PIP$_2$ and ATP were used to study their interactions with gelsolin using steady-state fluorescence anisotropy, and Alexa488-labeled gelsolin was utilized to reconstitute the regulation of gelsolin binding to PIP$_2$-containing phospholipid vesicles by ATP. Under physiological salt conditions ATP competes with PIP$_2$ for binding to gelsolin, while calcium causes the release of ATP from gelsolin. These data suggest a cycle for gelsolin activity. Firstly, calcium activates ATP-bound gelsolin allowing it to sever and cap F-actin. Secondly, PIP$_2$-binding removes the gelsolin cap from F-actin at low calcium levels, leading to filament elongation. Finally, ATP competes with PIP$_2$ to release the calcium-free ATP-bound gelsolin, allowing it to undergo a further round of severing.

Introduction

Gelsolin plays important roles in the dynamics and regulation of actin filament lengths [1]. Calcium ions activate gelsolin to sever actin filaments, leading to one of the two resulting filaments being capped at its barbed end. Cellular gelsolin is mostly in the cytoplasm where F-actin severing, capping and uncapping by gelsolin, close to the plasma membrane, are thought to contribute to cell movement and membrane morphology [2–4]. Severed actin filaments, which are capped by gelsolin at their barbed ends, can be uncapped to produce directed polymerization at the plasma membrane. Phosphatidylinositides are involved in signaling to the actin cytoskeleton by modifying the activity of various actin-binding proteins, including the...
gelsolin superfamily proteins [5]. In particular, phosphatidylinositol 4,5-bisphosphate (PIP$_2$) is a major regulator of actin cytoskeletal organization [6, 7] that modulates many actin regulating proteins [8], including: actin filament capping proteins, such as gelsolin [9], CapG [10] and capping protein [11]; actin monomer-binding proteins, like profilin [12], coflin [13] and twinfilin [14]; actin filament nucleation effectors, including WASP [15], N-WASP [16] and dynamin2/cortactin [17]; actin filament crosslinking proteins, exemplified by α-actinin [18], filamin [19] and cortexillin [20]; and actin filament plasma membrane tethering proteins, represented by vinculin [21], talin [21] and ERM proteins [22]. When gelsolin localizes to PIP$_2$-rich areas of the plasma membrane [23], PIP$_2$ inhibits interactions between membrane-bound gelsolin and actin [9, 24–27] and removes gelsolin caps from actin filaments [9]. There is strong evidence to suggest that local accumulation of PIP$_2$ at the plasma membrane leads to the uncapping of gelsolin-capped filaments, resulting in rapid, directed filament elongation [9, 28].

Three PIP$_2$-binding sites have been identified on gelsolin: between residues 135–142, which overlaps with one of the G-actin binding sites; between residues 161–169, which overlaps with the F-actin binding site; and between residues 621–634, which overlaps with the ATP binding site [27, 29–31]. The second site is well conserved within the gelsolin superfamily. The mechanism of uncapping is not fully understood, however PIP$_2$ may either directly compete with actin for binding to gelsolin, and/or it may change the conformation of the actin-binding sites to become incompatible with binding to actin [26, 31–33]. There are several reports of a correlation between PIP$_2$ and calcium binding to gelsolin, however it is controversial whether this correlation is positive or negative [27, 29, 34].

Gelsolin can bind to the ATP-mimetic resin Cibracon-Blue and can be liberated from the resin by a range of nucleotides including ATP, ADP, GTP and GDP [35, 36]. Equilibrium dialysis experiments were used to determine dissociation constants of 0.28 μM and 1.8 μM for gelsolin with ATP and GTP, respectively, at high NaCl concentrations, while ADP and GDP showed no significant association under these conditions [37]. The affinity of ATP for gelsolin decreases (Kd = 2.4 μM) in the presence of 0.2–2.0 mM Mg$^{2+}$ (pMg 4–3, where pMg = —log [Mg$^{2+}$]). No ATP binding to gelsolin is detectable in solutions that contain more than 10 μM Ca$^{2+}$ [38]. Gremm and Wegner reported a lower affinity of calcium-free gelsolin for ATP (Kd = 32 μM) than discussed above [38]. This may reflect differences in analytical techniques, fluorescence analysis versus equilibrium dialysis, or differences in experimental buffer conditions, including pH, MgCl$_2$ and CaCl$_2$. Gelsolin does not show any detectable ATPase activity [37, 39]. The discovery of the gelsolin:ATP interaction led to the suggestion that ATP may be important in some of the multiple functions of gelsolin. The structure of the gelsolin:ATP complex revealed the basis for its sensitivity to calcium ion concentrations in that ATP interacts with both of the two halves of calcium-free gelsolin, which change conformation on binding to calcium [32, 40–45]. Thus, the loss of its ATP-binding ability is due to disruption of the interaction site within gelsolin caused by a conformational change in response to calcium [37]. The phosphate groups of ATP interact with basic residues on gelsolin domain 5 (G5, residues 514–619). These residues also comprise part of a region that previously had been determined to bind to PIP$_2$ [27]. Gelsolin is also sensitive to the type of nucleotide bound to actin; it severs filaments that contain ADP-actin but not ADP-Pi-actin units [46]. Accordingly, G4-G6 (residues 418–741) shows a preference for ADP-containing actin monomers while G1-G3 (residues 25–370) binds to ATP- and ADP-actin with comparable affinities [47]. In contrast, ATP (but not ADP) concentrations in the mM range inhibit the binding of G1-G3 to actin monomers [47].

Generally, the free calcium concentrations in resting cells oscillate on the nanomolar scale [48, 49], however in stimulated cells these concentrations can increase to micromolar levels.
Cytoplasmic free magnesium levels are regulated in the pMg 4–3 (0.5–1 mM) range [50, 51]. Intracellular ATP concentrations (2 μM—8 mM, mean value 0.5–1 mM) can change over a wide range depending on the type, stage and state of the cell, and these levels play a regulatory role in plasma membrane channel function [52, 53]. Gelsolin is activated by calcium, not by magnesium, to sever and cap filaments. However the gelsolin/actin complex is not dissociated by the lowering of calcium levels, with one bound calcium ion becoming trapped in the complex [54–61]. In this report we probe the interplay between ATP and calcium in modifying PIP$_2$ binding to gelsolin. We postulate, based on our in vitro data, which ATP plays a critical role in the recycling of gelsolin by removing gelsolin from PIP$_2$ in the plasma membrane, leading to the release of gelsolin into the cytoplasm to undergo further rounds of calcium-induced actin filament severing.

**Experimental procedures**

**Proteins**

His-tagged human wild-type gelsolin was expressed in E. coli strain Rosetta2 (DE3) pLysS cells from a pSY5 plasmid [45]. The protein was subjected to Ni-NTA affinity chromatography, HRV 3C protease cleavage, followed by gel filtration (Superdex 200, GE Healthcare) in 10 mM Tris–HCl, 150 mM NaCl, pH 8. Traces of calcium were removed by dialysis (2 mM Tris–HCl, 1 mM EGTA, pH 7.4, overnight). Rabbit skeletal muscle actin was prepared from acetone powder (Pel-Freez Biologicals) in a protocol modified from Spudich and Watt [63, 64]. Actin was stored in buffer A (2 mM Tris–HCl, 0.2 mM ATP (ATP disodium trihydrate, Sigma–Aldrich), 0.1 mM CaCl$_2$ (pCa 4), 0.1 mM DTT and 0.005% NaN$_3$, pH 7.4). Alexa Fluor$^\text{TM}$ 488 C5 Maleimide (Life Technologies) was used to label cysteine residues of gelsolin based on the manufacturer's recommendations (Thermo Fisher Scientific).

**Phospholipid vesicle preparation**

Phospholipid vesicles were prepared by a modified protocol from James H. Morrissey, Dept. of Biochemistry, University of Illinois at Urbana-Champaign, Urbana, IL 61801, USA (https://tf7.org). A mixture of 1% PIP$_2$ (PtdIns-(4,5)-P$_2$ (1,2-dipalmitoyl)) (Cayman Chemicals), 79% PC (L-α-phosphatidylcholine) (Sigma–Aldrich) and 20% PS (3-sn-phosphatidyl-L-serine) (Sigma–Aldrich) was dissolved in 20 μM rhodamine 590 N-succinimidyl ester (Sigma–Aldrich) in chloroform. The dried lipid mixture was sonicated in an Elmasonic S100H water bath sonicator (medium strength of pulses) in 2 mM Tris–HCl, 200 mM NaCl, pH 7.4, with the temperature maintained at 40 °C, until the solution became visually homogeneous (approx. 30 mins) and small multimellar vesicles were formed. Vesicles were collected by centrifugation at 22 °C for 10 min at 5,000 x g, and stored at 4 °C.

**Gelsolin intrinsic tryptophan fluorescence**

The intrinsic tryptophan (residue numbers: 67, 111, 203, 223, 341, 392, 446, 489, 601, 637, 699, 706, 755, 759, 764) fluorescence emission assays were carried out with a Perkin Elmer LS-50 spectrofluorimeter. The excitation and emission monochromators were set to 288 nm and 332 nm, respectively, and the excitation and emission bandwidths to 5 nm. 5 μM gelsolin was incubated under physiological salt conditions (100 μM CaCl$_2$ (pCa 4), 100 mM KCl, 1 mM MgCl$_2$ (pMg 3), 0.2 mM ATP, 2 mM Tris–HCl, pH 7.4) supplemented with EGTA or CaCl$_2$ to vary the free calcium levels (calculated with Maxchelator Stanford http://maxchelator.stanford.edu/CaMgATPEGTA-NIST.htm): pCa 9: 6 mM EGTA; pCa 6: 100 μM EGTA; pCa 3: 1 mM CaCl$_2$. 

[50, 51].
Steady-state fluorescence intensities were measured after sequential addition of appropriate stock solutions to attain: 1) 2 μM PIP₂, 2) 0.5 mM ATP, and 3) pCa 6.

**Steady-state fluorescence anisotropy**

We used a fluorescent derivative of PIP₂ (PtdIns-(4,5)-P₂-fluorescein, Cayman Chemicals, NU-10010388, λ_ex = 493 nm, λ_em = 520 nm, abbreviated as PIP₂-F) and a fluorescent derivative of ATP (N⁶-(6-amino)hexyl-ATP-ATTO-532, Jena Bioscience, NU-805-532, λ_ex = 532 nm, λ_em = 553 nm, abbreviated as ATP-N) as probes for the binding of PIP₂ and ATP to gelsolin. Fluorescence intensities were measured on a Safire² monochromator microplate reader (TECAN) at 22°C. Steady-state fluorescence anisotropy values were determined according to Eq 1 [65], where \( I_{VV} \) and \( I_{VH} \), respectively, are the intensities of vertically and horizontally polarized emission on excitation with vertically polarized light, with the correction factor, \( G \), being the ratio of \( I_{VH}/I_{VV} \).

\[
r = \frac{I_{VV} - GI_{VH}}{I_{VV} + 2GI_{VH}}
\]  

The dissociation constants for ATP and PIP₂ were calculated using a hyperbolic model [66] from anisotropy data that had reached saturation during the gelsolin titrations Eq 2:

\[
r = r_f + (r_b - r_f) \left( \frac{K_d + [G] + [L] - \sqrt{(K_d + [G] + [L])^2 - 4[G][L]}}{2[L]} \right)
\]

where \( r_f \) is the anisotropy of free ligand, \( r_b \) is the anisotropy of gelsolin-bound ligand, \([L]\) is the total concentration of ligand, \( K_d \) is the dissociation constant and \([G]\) is the total concentration of gelsolin. The anisotropy of 0.5 μM ATP-N was measured in the presence of gelsolin concentrations: 0, 0.2, 1.5, 2, 3, 5 and 8 μM. Saturation was evident at 5 μM characterized by an anisotropy value of 0.15 ± 0.006. The calcium and magnesium sensitivities of ATP binding to gelsolin were characterized by the change in anisotropy of solutions containing 0.5 μM ATP-N in the presence of 5 μM of gelsolin over a range of divalent cation concentrations of pCa 2–12 or pMg 1–12, where pCa and pMg refer to, respectively. The affinity of the PIP₂-gelsolin interaction was measured by the anisotropy change of 0.5 μM or 0.25 μM PIP₂-F in the presence of 0, 0.2, 1.5, 3, 5 and 8 μM gelsolin, which showed saturation at 4 μM. Anisotropy of 0.5 μM or 0.25 μM solutions of PIP₂-F in the presence of 5 μM gelsolin was used to probe the changes in binding of the complexes under different calcium, magnesium, salt and ATP conditions.

**Confocal microscopy imaging**

10% (v/v) rhodamine590-filled phospholipid vesicles and 5 μM Alexa488-labeled gelsolin were incubated together and a drop of the mixture was placed on a clean glass slide, which was then covered by and separated from a second slide by a parafilm gasket. In this setup, buffer conditions surrounding the phospholipid vesicles can be changed by laminar flow between the two slides. A 2 mm x 2 mm piece of tissue (KimWipes) was used to secure the vesicles. The sample was washed (2 mM Tris-HCl, pH 7.4) for 1 min and then placed on the glass slide before being covered. In the flow cell, phospholipid vesicles were kept in the field of view by the tissue fibers. Fluorescence emission-based imaging was carried out using a Zeiss LSM 510 META Confocal Microscope. The two different fluorophores were detected in two separate channels; Alexa488 (gelsolin) was excited by 477 nm laser light and emission was detected in the 505–530 nm channel, while rhodamine590 (phospholipid vesicles) was excited by 545 nm laser light and emission was detected in the 585–630 nm channel.
Statistics

The data presented were derived from at least 3 independent experiments. Values are displayed as the mean ± standard deviation.

Results

In order to probe the interplay between PIP$_2$, ATP and Ca$^{2+}$ in binding to gelsolin, we first characterized the binding of fluorescent derivatives of PIP$_2$ and ATP to gelsolin using steady-state anisotropy measurements. The fluorescent derivative of ATP (0.5 μM), N$^6$-(6-amino)hexyl-ATP-ATTO-532, named ATP-N hereafter, binds to gelsolin with a $K_d$ of 0.71 ± 0.52 μM in the absence of divalent cations, as determined by analysis of steady-state anisotropy measurements (Fig 1A). This value is within experimental error of that previously reported for the gelsolin:ATP interaction (0.28 μM), as measured by equilibrium dialysis [39]. Micromolar levels of calcium (pCa 6) ions or millimolar levels of magnesium (pMg 3) ions were able to effectively dissociate ATP-N from gelsolin (5 μM), characterized by 50% of the ATP-N remaining bound at 6.3 ± 0.2 μM calcium or 5.0 ± 1.1 mM magnesium (Fig 1B). This calcium range that induces ATP dissociation is in line with physiological calcium signaling levels and with calcium concentrations that are able to initiate conformational changes in gelsolin, which disrupt the ATP-binding site leading to ATP release [42]. The magnesium levels that release ATP-N from gelsolin are close to the reported affinity of ATP for Mg$^{2+}$ [67], which is in line with gelsolin/ATP structure [32], which does not contain cations. Thus, the Mg$^{2+}$-induced release of ATP from gelsolin by Mg$^{2+}$ alone is likely due the incompatibility of Mg-ATP with gelsolin. However, the effective concentration of magnesium is higher than normal free physiological levels pMg 4–3 (0.5–1.0 mM) [52]. We next tested the effect of ionic strength by the addition of potassium ions, which weakened the gelsolin:ATP-N interaction (S1 Fig). Under physiological potassium ion concentrations, the ATP-N interaction with gelsolin was characterized by $K_d$s of 1.27 ± 0.60 μM and 1.42 ± 0.27 μM in 100 mM and 120 mM KCl, respectively, values which were increased slightly by the addition of 1 mM MgCl$_2$ (pMg 3) ($K_d$s of 1.58 ± 0.85 μM and 2.34 ± 0.73 μM, respectively) (S1 Fig). Similarly, in 50 mM KCl and 1 mM MgCl$_2$ (pMg 3), a condition commonly used for in vitro actin polymerization experiments [63], gelsolin binding of ATP-N was determined by titration with gelsolin and via competition of ATP-N binding by unlabeled ATP, and the $K_d$ values were 1.8 μM (ATP-N) and 2.2 μM (ATP) (S2 Fig). The effect of calcium under these actin polymerization conditions recapitulated the observation made under the low salt conditions, with micromolar (pCa 6) and higher concentrations of calcium negatively impacting the gelsolin:ATP-N interaction (Panel a in S2 Fig vs. S1 Fig). Collectively, these data indicate that under physiological buffer conditions gelsolin binds to both ATP-N and unlabeled ATP in similar fashion and that calcium acts as a regulator for the binding.

Gelsolin has been shown to have three PIP$_2$-binding sites, which have previously been characterized to display $K_d$ values in the 1–20 μM range [27, 29, 31]. In a similar strategy to that adopted for characterizing the ATP-gelsolin interaction, a soluble fluorescent derivative of PIP$_2$, PtdIns-(4,5)-P$_2$-fluorescein, named PIP$_2$-F hereafter, was used to probe the gelsolin-PIP$_2$ interaction using steady-state anisotropy measurements. In this assay the binding of PIP$_2$-F (0.5 μM) to gelsolin in the absence of divalent cations was characterized by a $K_d$ of 0.84 ± 0.39 μM (Fig 1C). This value increased to 1.35 ± 0.7 μM and 3.12 ± 1.7 μM in the presence of increased ionic strength, 100 mM and 120 mM KCl, respectively, and these values remained constant within experimental error on the further addition of 1 mM MgCl$_2$ (pMg 3) (S3 Fig). Thus, the affinities of gelsolin for PIP$_2$-F and ATP-N lie in the same range under physiological salt conditions.
Subsequently, we studied the sensitivity of PIP$_2$-F (0.5 μM) binding to gelsolin (5 μM) in the absence/presence of ATP and divalent cations (magnesium and calcium). It is known that divalent cations, calcium in particular, cause PIP$_2$ to aggregate [68, 69]. Therefore, we evaluated the effects of cations on the critical micelle concentrations (CMCs) of PIP$_2$ and PIP$_2$-F by dynamic light scattering [70]. Under the buffer conditions tested, PIP$_2$-F failed to form micelles at or below concentrations of 0.5 μM, whereas PIP$_2$ was more sensitive to cations and had a greater tendency to form micelles (S4 Fig). Thus, we used 0.5 μM PIP$_2$-F to probe its interactions with gelsolin.
The first evidence of the competition between ATP and PIP$_2$ in binding to gelsolin was observed from the elevated anisotropy (0.12, Fig 1C) of PIP$_2$-F (0.5 μM), characteristic of its association with gelsolin (5 μM), being lost on the addition of 0.2 or 0.5 mM ATP (Fig 1D). The ability of ATP to dissociate PIP$_2$-F remained constant over a wide range of Mg$^{2+}$ concentrations, from picomolar (pMg 12) to the physiologically relevant millimolar (pMg 3) range, becoming less effective above 7 mM (pMg 2.15). Hence, ATP can effectively compete with PIP$_2$ for binding to gelsolin under physiological Mg$^{2+}$ conditions. In the absence of ATP, PIP$_2$-F was able to bind gelsolin, characterized by a steady-state anisotropy value of approximately 0.12, over the ≈0–10 μM range of Ca$^{2+}$ (pCa 15–5)(Fig 1D). At higher calcium levels the anisotropy began to reduce, reaching 0.04 at 1 mM Ca$^{2+}$ (pCa 3)(Fig 1D). The PIP$_2$-F/gelsolin interaction was characterized by $K_d$s below 20 μM across the entire calcium range, in line with previously reported PIP$_2$/gelsolin affinities [27, 29, 31], and below 1 μM in the 1–10 μM range (pCa 6–5) (Panel a in S5 Fig), in the absence of KCl and magnesium. In the presence of 0.2 mM ATP in the ≈0–10 μM calcium (pCa 15–5) range, the anisotropy was close to the baseline characteristic of free PIP$_2$-F, indicating the effective dissociation of the PIP$_2$-F/gelsolin interaction (Fig 1D) (Panel a in S5 Fig) with estimated $K_d$s in excess of 300 μM (Panel a in S5 Fig). Higher calcium levels were not obtainable in the presence of ATP due to precipitation of these reagents in the absence of salt (Panel b in S5 Fig). In a control experiment, PIP$_2$ had no observable effect on the binding of calcium to FURA-2FF, implying that any calcium binding by PIP$_2$ is substantially weaker than that of FURA-2FF ($K_d$ = 25 μM, Panel c in S5 Fig), and it is unlikely to have had a significant effect in these experiments. Thus, these data suggest that ATP can release PIP$_2$-F from gelsolin effectively in the ≈0–10 μM calcium (pCa 15–5) concentration range. We confirmed that the PIP$_2$-F-gelsolin complex remained intact under actin polymerization conditions, and that PIP$_2$-F was released from gelsolin when these conditions were supplemented with 0.5 mM ATP (Fig 2A), the calculated $K_d$s ranged from 2.7 ± 0.2 μM in absence of ATP followed by a substantial rise to 65.4 ± 6.2 μM after the addition of ATP.

Subsequently, we used the intrinsic fluorescence emission at 332 nm from the 15 tryptophan residues evenly distributed across the 6 domains of gelsolin to probe the effects of higher concentrations of PIP$_2$ and ATP on gelsolin than could be probed by the PIP$_2$-F and ATP-N anisotropy assays. Since gelsolin contains 6 calcium-binding sites, some of which have $K_d$ values below the protein concentration used in the experiment, we turned to an EGTA-buffered calcium system to control the free calcium levels. Gelsolin was incubated under actin polymerizing conditions (100 μM CaCl$_2$ (pCa 4), 100 mM KCl, 1 mM MgCl$_2$ (pMg 3), 0.2 mM ATP, 2 mM Tris-HCl, pH 7.4), which was supplemented with EGTA or CaCl$_2$ to vary the free calcium levels. Gelsolin (5 μM) showed lower tryptophan emission levels when bound to ATP (0.5 mM) relative to PIP$_2$ (2 μM) at 1 μM (pCa 6) and 1 nM (pCa 9) CaCl$_2$ (Fig 2B). This difference was used to demonstrate the cycling between gelsolin binding to PIP$_2$ and ATP. Gelsolin showed a decrease in tryptophan fluorescence in moving from 1 nM (pCa 9) free calcium to activating levels of calcium (pCa 6 and pCa 3, Fig 2C). Subsequent addition of PIP$_2$ (2 μM) to these conditions raised the intrinsic tryptophan fluorescence at 1 nM (pCa 9) and 1 μM (pCa 6), but not at 1 mM (pCa 3) free calcium, indicating that tryptophan fluorescence is a sensitive reporter on the gelsolin/PIP$_2$ interaction in the pCa 9 to pCa 6 range (Fig 2C/1). Subsequent addition of ATP (0.5 mM) led to a reduction in the intrinsic fluorescence at both pCa 9 and pCa 6 free calcium (Fig 2C/2). Finally, the tryptophan fluorescence converged on adjusting each condition to pCa 6 free calcium, indicating that the interplay between calcium and gelsolin in PIP$_2$/ATP was in equilibrium and reversible (Fig 2C/3). Thus, the tryptophan assay shows that ATP can exert competitive effects against PIP$_2$ in the pCa 9 and pCa 6 calcium range.

Next we probed the effect of actin on the competition between ATP and PIP$_2$ in binding to gelsolin. The association of PIP$_2$-F (0.25 μM) with gelsolin (5 μM) was investigated as a...
function of free \( \text{Ca}^{2+} \) concentration under actin polymerizing salt conditions (100 mM K\(^+\), 1 mM Mg\(^{2+}\) (pMg 3)). Addition of 0.5 mM ATP caused an almost complete loss of PIP\(_2\)-F anisotropy across the entire calcium range, both in the absence and presence of 50 μM actin.
Together these data indicate that ATP can release PIP$_2$ from gelsolin in the presence of salt, calcium ions and actin.

Finally, we used confocal microscopy to determine whether ATP can be observed to release gelsolin from phospholipid vesicles. PIP$_2$-containing rhodamine590-filled phospholipid vesicles were observed to bind to Alexa488-labeled gelsolin in the absence of ATP, calcium and magnesium (S1 Movie; Fig 3A, upper panel). This gelsolin was subsequently released by the addition of 0.5 mM ATP (S1 Movie; Fig 3A, middle panel). A second round of labeled gelsolin could be bound to the phospholipid vesicles after the solution had been exchanged to remove the ATP (S1 Movie; Fig 3A, lower panel). During the release of gelsolin by ATP, some phospholipid vesicles were observed to display shape changes (S2 Movie; Fig 3B, S6 Fig). This suggests that gelsolin binds to PIP$_2$ in the phospholipid vesicles and may induce deformation or local structural changes in the vesicles and that ATP can effectively dissociate gelsolin from the phospholipid vesicles.
Discussion

ATP and PIP$_2$ have been shown to compete in binding to K-ATP channels [71]. Here, we have demonstrated that ATP can displace PIP$_2$ from gelsolin in solution under actin polymerizing buffer conditions in vitro. Furthermore, ATP is able to release PIP$_2$-bound gelsolin from the surface of phospholipid vesicles. These observations suggest that ATP is likely to dissociate gelsolin from PIP$_2$ at plasma membranes, and this ATP-driven dissociation is the missing step in recycling of gelsolin during its actin filament remodeling cycle. In a background of high cellular ATP, PIP$_2$ will not generally bind to gelsolin. However, in the situation where gelsolin-capped filaments point at the plasma membrane, the filament barbed-end bound gelsolin becomes greatly reduced in its mobility, and it is in close proximity to membrane-bound PIP$_2$ that can move within the plasma membrane and increase its local concentration by forming clusters [72]. All these factors favor the binding between gelsolin and PIP$_2$, and hence, filament uncapping. We propose that effective competition by ATP will dominate following filament uncapping and that diffusion of PIP$_2$-bound gelsolin will increase.

![Fig 4. Model of the severing, capping, uncapping and inactivation/release cycle of gelsolin.](https://doi.org/10.1371/journal.pone.0201826.g004)
The cartoon presented in Fig 4 details the postulated stages of this remodeling cycle under standard cellular conditions. Activation: elevation of calcium levels leads to a conformational change in gelsolin that releases ATP and allows gelsolin to recognize an actin filament. Severing: competition for actin-actin interactions by gelsolin-actin interactions leads to the severing of the filament. Capping: gelsolin remains bound to the barbed-end of the severed filament, preventing its elongation. Uncapping: when a gelsolin-capped filament encounters PIP$_2$ in the plasma membrane, the cap is removed through an unknown mechanism. The uncapped filament is then free to elongate and exert force on the plasma membrane. Release: gelsolin is released from PIP$_2$ at the plasma membrane through ATP competition, leading to diffusion of the gelsolin:ATP complex away from the plasma membrane. Gelsolin will return to its inactive state in low calcium environments. Thus, gelsolin is likely removed from PIP$_2$ at the plasma membrane in an ATP-dependent manner that distinguishes it from other PIP$_2$-sensing actin-regulating proteins, allowing gelsolin to cycle in a background of elevated PIP$_2$.

Supporting information

S1 Fig. The interplay among calcium, magnesium and potassium ions on ATP-N binding to gelsolin. In the absence of divalent cations, the anisotropy of ATP-N (0.5 μM) increased with increasing gelsolin concentration and this interaction was potassium ion dependent. The K$_d$s calculated from the binding curves fitted with Eq 2 showed diminishing affinities with increasing potassium ion concentrations (blue triangles). Inclusion of 1 mM MgCl$_2$ slightly weakened the affinity of ATP-N for gelsolin (magenta circles) at 100 mM and 120 mM KCl (K, potassium ions, KM, potassium and magnesium ions).

S2 Fig. ATP binding to gelsolin under actin polymerization conditions. (A) The calcium-dependent gelsolin binding affinity for ATP under actin polymerization conditions was determined by monitoring the change in ATP-N anisotropy on gelsolin titration at different calcium ion concentrations. Each data point arises from the K$_d$ calculated from a titration similar to Fig 1A. K$_d$s were calculated to be between 1.51 μM and 2.35 μM in the 1 nM to 10 μM calcium concentration range, and the value increased to 8.05 ± 1.32 μM above 100 μM calcium. (B) The anisotropy of ATP-N (0.5 μM) bound to gelsolin (5 μM) was measured upon titrating with unlabeled ATP in the absence of calcium. 50 μM ATP was sufficient to remove the gelsolin-bound ATP-N. The affinity of unlabeled ATP for gelsolin, under actin polymerizing salt conditions, was calculated to be K$_d$ = 2.2 ± 0.17 μM measured by labeled/unlabeled ATP competition on gelsolin, which is similar to the affinity of ATP-N derived from Panel c in S1 Fig (K$_d$ = 1.58 ± 0.8 μM). Data were analyzed by the method of Kubala (Kubala et al. 2004) with the modification that the change in anisotropy of ATP-N was substituted for the change in fluorescence emission as the indication of ATP competition.

S3 Fig. The interplay between calcium ions, magnesium ions and ionic strength on PIP$_2$-F binding to gelsolin. In the absence of divalent cations, the anisotropy of PIP$_2$-F (0.5 μM) increased with increasing gelsolin concentration and this interaction was potassium ion dependent. The K$_d$s calculated from the binding curves fitted with Eq 2 showed diminishing affinities with increasing potassium ion concentrations (blue triangles). Inclusion of 1 mM MgCl$_2$ slightly weakened the affinity of PIP$_2$-F for gelsolin (magenta squares) at 100 mM and 120 mM KCl (K, potassium ions, KM, potassium and magnesium ions). (B) Anisotropy of PIP$_2$-F (0.5 μM) in the presence of gelsolin (5 μM) as a function of KCl or NaCl concentration. Data were fitted with simple sigmoidal curves. The decrease in anisotropy indicates that PIP$_2$-
F is dissociated from gelsolin by increasing potassium or sodium ion concentrations, with the half-effective concentrations of $141.2 \pm 3.0 \, \text{mM}$ and $143.3 \pm 1.6 \, \text{mM}$, respectively. This indicates the effect of KCl on gelsolin:PIP$_2$F binding is largely ionic and nonspecific.

(S4 Fig. Assessment of the solubility of PIP$_2$ and PIP$_2$-F in the experimental buffers. (A) Determination of the actual concentrations of PIP$_2$-F after incubation with different cations by light absorbance at 494 nm. T, 2 mM Tris-HCl, pH 7.4; K, 100 mM KCl; C, 1 mM CaCl$_2$; M, 1 mM MgCl$_2$. (B) Determination of critical micelle concentrations of PIP$_2$ and PIP$_2$-F (Panel inset) by dynamic light scattering.

(S5 Fig. Assessment of the effects of calcium and ATP on the binding of ATP-N and PIP$_2$-F to gelsolin. (A) Kds were calculated from the fit data with Eq 2 of gelsolin:PIP$_2$-F binding under different calcium concentrations in the absence or presence of ATP. Part of the data have been shown in Fig 1D (white triangles). In the absence of ATP, PIP$_2$-F (0.5 μM) binds to gelsolin, as reflected by the high affinity, across a wide range of calcium concentrations. In the presence of ATP, the interaction of PIP$_2$-F with gelsolin was very weak below 10 μM calcium ($K_d = 563.0 \pm 3.6 \, \text{μM}$ at 10 μM calcium). (B) In the absence of gelsolin, calcium directly precipitates ATP-N (C, open triangles) with a half-maximum value of 70.4 ± 0.7 μM. Magnesium has no effect on the fluorescence emission of ATP-N either in the absence (M, orange circles) or presence of 10 μM calcium (MC, purple squares). Data were fitted with simple sigmoidal curves. (C) The reported FURA-2FF dissociation constant for calcium ions is 25 μM (A.G. Scientific Inc.). The calcium-dependent fluorescence emission profile of FURA-2FF (2 μM) over pCa range of 5–7 was similar in the presence and absence of 20 μM PIP$_2$, which were fitted with simple sigmoidal curves. The half-saturation pCa values were $6.265 \pm 0.004$ and $6.217 \pm 0.011$ in the absence and presence of PIP$_2$, respectively, suggesting that PIP$_2$ does not interact with calcium with a $K_d$ less than 25 μM. The measurement was carried out with a Perkin Elmer LS-55 spectrofluorimeter ($\lambda_{ex} = 340 \, \text{nm}$, $\lambda_{em} = 505 \, \text{nm}$).

(S6 Fig. Line scans of fluorescence intensity versus distance along the arrows shown in Fig 3B. (A) Profiles of gelsolin-Alexa488 (red) and PIP$_2$-containing vesicles filled with rhodamine590 (blue) in the absence of ATP. (B) Gelsolin-Alexa488 (red) was released and the size of phospholipid vesicle (filled by rhodamine590, blue) was changed by 0.5 mM ATP treatment in Fig 3B.

(S1 Movie. The influence of ATP on the binding of Alexa488-labeled gelsolin to PIP$_2$-containing rhodamine590-filled phospholipid vesicles. Time course for Fig 3A. Gelsolin-Alexa488 (5 μM, red, top left) was incubated with rhodamine590-filled PIP$_2$-containing membrane vesicles (blue, top right) and visualized by confocal microscopy. The merged image indicates that gelsolin and vesicles colocalized (bottom left). After ATP (0.5 mM) was added the majority of gelsolin-Alexa488 was released from the vesicles. Following removal of ATP via buffer exchange and addition of fresh gelsolin-Alexa488 (15 μM), gelsolin re-associated with the vesicles. The movie plays six times faster than the real time course.

(S2 Movie. Shape changes of phospholipid vesicles during the release of gelsolin by ATP. Time course for Fig 3B. After the addition of ATP (0.5 mM) the vesicle changed morphology concurrently with the release of gelsolin (red) from the vesicle surface (blue). The movie plays...
six times faster than the real time course. 

(MP4)

**Acknowledgments**

DS, BX, BK and RCR thank A*STAR for support. This research was supported by grants from the Hungarian Science Foundation (OTKA) Grants K109689 (to BB) and K112794 (to MN), by the UNKP-16-4 New National Excellence Program of the Ministry of Human Capacities and by the UNKP-17-4 New National Excellence Program of the Ministry of Human Capacities (to BB).

**Author Contributions**

**Conceptualization:** Dávid Szatmári, Bo Xue, Leslie D. Burtnick, Miklós Nyitrai, Robert C. Robinson.

**Data curation:** Beáta Bugyi.

**Funding acquisition:** Robert C. Robinson.

**Investigation:** Dávid Szatmári, Bo Xue, Balakrishnan Kannan.

**Methodology:** Dávid Szatmári, Bo Xue, Balakrishnan Kannan.

**Supervision:** Miklós Nyitrai, Robert C. Robinson.

**Writing – original draft:** Dávid Szatmári, Bo Xue, Balakrishnan Kannan, Leslie D. Burtnick, Beáta Bugyi, Miklós Nyitrai, Robert C. Robinson.

**Writing – review & editing:** Dávid Szatmári, Bo Xue, Balakrishnan Kannan, Leslie D. Burtnick, Beáta Bugyi, Miklós Nyitrai, Robert C. Robinson.

**References**


