Enantiomeric Aβ peptides inhibit the fluid shear stress response of PIEZO1

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Traumatic brain injury (TBI) elevates Abeta (Aβ) peptides in the brain and cerebral spinal fluid. Aβ peptides are amphipathic molecules that can modulate membrane mechanics. Because the mechanosensitive cation channel PIEZO1 is gated by membrane tension and curvature, it prompted us to test the effects of Aβ on PIEZO1. Using precision fluid shear stress as a stimulus, we found that Aβ monomers inhibit PIEZO1 at femtomolar to picomolar concentrations. The Aβ oligomers proved much less potent. The effect of Aβs on Piezo gating did not involve peptide-protein interactions since the D and L enantiomers had similar effects. Incubating a fluorescent derivative of Aβ and a fluorescently tagged PIEZO1, we showed that Aβ can colocalize with PIEZO1, suggesting that they both had an affinity for particular regions of the bilayer. To better understand the PIEZO1 inhibitory effects of Aβ, we examined their effect on wound healing. We observed that over-expression of PIEZO1 in HEK293 cells increased cell migration velocity ~10-fold, and both enantiomeric Aβ peptides and GsMTx4 independently inhibited migration, demonstrating involvement of PIEZO1 in cell motility. As part of the motility study we examined the correlation of PIEZO1 function with tension in the cytoskeleton using a genetically encoded fluorescent stress probe. Aβ peptides increased resting stress in F-actin, and is correlated with Aβ block of PIEZO1-mediated Ca²⁺ influx. Aβ inhibition of PIEZO1 in the absence of stereospecific peptide-protein interactions shows that Aβ peptides modulate both cell membrane and cytoskeletal mechanics to control PIEZO1-triggered Ca²⁺ influx.

There is a correlation between traumatic brain injury (TBI), Amyloid β peptide levels (Aβ) and the onset of TBI related diseases1. The relationship between Aβ and clinical manifestations of TBI is not understood1. Since, the initiating mechanical forces ultimately resulting in TBI appear not to visibly damage cells, a multi-step force signaling mechanism is likely involved, including changes to the cytoskeleton. A potential link in this pathway is the activity of mechanosensitive (MSC) PIEZO channels2,3 that respond to stress in the lipid bilayer4–6. Since amphipathic drugs including Aβ peptides alter membrane structure7,8 and thus membrane mechanics, we examined the effects of Aβs on PIEZO1.

How amphipaths may alter mechanical channels is suggested from studies on bacterial MSCs that respond to global forces applied through bilayer lipids9,10 or by lipid perturbations around the channels11. The recently published cryoEM structure of mouse Piezo1 in a bilayer dome underscores the connection between lipids and MSCs as an integrated system12.

Modulation of Piezo channels by amphipaths can potentially influence a number of cellular responses. For example, the differentiation of mouse neural stem cells into either neurons or glia, requires PIEZO1; inhibition of PIEZO1 sensing of cell crowding and cell division14,15. PIEZO1 channels are involved in neurite extension and inhibition of PIEZO1 can alter neuronal outgrowth16. All these processes are associated with sensing the local mechanical environment17.

In this work, we asked whether Aβ can affect PIEZO1. We tested the ability of Aβ peptides to modulate the cellular response to mechanical inputs generated by a precision fluid shear stress system, measuring
PIEZO1-mediated Ca\(^{2+}\) fluxes. A stable overexpressing PIEZO1 cell line made the response larger and more reliable. Titration of A\(\beta\) monomeric A\(\beta\)s in the femtomolar to picomolar range were effective in inhibitors, but A\(\beta\) oligomers were much less potent. PIEZO1 inhibition by D and L enantiomers of A\(\beta\)s proved equally effective, showing a lack of peptide-protein or other stereospecific interactions. The peptide-channel interaction is likely mediated via bilayer lipids. The inhibitory effects of A\(\beta\)s were similar to responses to GsMTx4, another amphipathic, stereo-nonspecific inhibitor of PIEZO1 but requiring much higher concentrations. Inhibition of PIEZO1 by A\(\beta\)s or GsMTx4 almost completely blocked cell migration in a wound healing assay. Because migration requires a pliable cytoskeleton, we measured the effect of A\(\beta\)s on cytoskeletal stresses using a genetically encoded optical force probes embedded in actin. The inhibition of PIEZO1 by A\(\beta\) and the resulting block of Ca\(^{2+}\) influx was accompanied by significantly increased actin tension. PIEZO1 activity is causally linked to cell motility and cytoskeletal reorganization.

Results

Fluid shear stress assay. We first created a cell line that expressed a fluorescently tagged version of human PIEZO1 called hPIEZO1-1591-EGFP. The fusion protein encoded internal GFP and N-terminal His tag did not significantly affect the channel properties. The cDNA was integrated into the genome of HEK293T cells using a lentivirus vector. The single channel conductance of this stably overexpressing PIEZO1 cell line (hP1-CL) was indistinguishable from that of hPIEZO1 in transiently transfected HEK293T cells (Supplementary Fig. 1).

A\(\beta\) peptide inhibition of PIEZO1. hP1-CL cells were grown on fibronectin-coated microfluidic chambers (Supplementary Fig. 2) and loaded with the calcium indicator, Fluo-4-AM and stimulated with 10 ms pulses of 15 dynes/cm\(^2\). Changes in fluorescence were monitored by a CCD camera at 1 Hz. Shown are intensity changes at four indicated time points. The maximum response is observed at \(\Delta t = 22\) s, long after termination of the stimulus. (Panel B). The L forms of A\(\beta\)(1-40) or (1-42) prepared by Peptide Method I at 10 \(\mu\)M inhibited the shear-induced elevation of [Ca\(^{2+}\)] response (4 independent experiments with SEM). (Panel C). PIEZO1 inhibitory potencies of the enantiomeric A\(\beta\) peptides (prepared by Peptide Method I) and of the commercially available monomeric peptide AgS, called Aggresure L-A\(\beta\)(1-40). The K\(_i\) (half-maximal concentration for inhibition) was determined based on the peak response plotted as a function of concentration. Each concentration is an average of 4 independent experiments with SEM. We observed K\(_i\) differences between the D and L forms of each peptide (diastereomeric pairs are identified with horizontal double-headed arrows). (Panel D) Monomeric A\(\beta\) peptides prepared by Peptide Method II were tested. The monomeric L-A\(\beta\)(1-40) was nearly identical to AgS in its ability to inhibit PIEZO1 function with a K\(_i\) of \(\sim 50\) nM (4 independent experiments with SEM).
PIEZO1 response (Supplementary Fig. 3A, black squares) at the same magnitude of shear stress (Supplementary Fig. 3A represents the response of 4 independent experiments). Cytoskeletal disruption of hP1-CL cells with either cytochalasin D (Supplementary Fig. 3A, green triangles) or colchicine (Supplementary Fig. 3A, dark blue triangles) produced cells unresponsive to fluid shear, suggesting a requirement of the PIEZO1 shear response for cytoskeletal integrity coupled to the bilayer. This is similar to our observation that mouse PIEZO1 whole cell currents required an intact cytoskeleton to respond to cell indentation.

We tested two Aβ peptides and found that both inhibited the shear response (Fig. 1B). Incubation of hP1-CL with 10 μM L-Aβ(1-40) or L-Aβ(1-42) peptides (see Peptide Method I in Methods section below) inhibited the fluid shear response (Fig. 1B, red circles and black squares, represents 4 independent experiments). Neither the addition of scrambled L-Aβ(1-40) peptide to hP1-CL cells nor expression of Amyloid Precursor Protein (APP)-YFP in hP1-CL had an inhibitory effect on the shear response (Supplementary Fig. 4 left and middle panel). The right panel (Supplementary Fig. 4) shows the average response of 4 independent experiments with SEM.

As a control, we examined the responses to the known inhibitor, D-GsMTx418. At 5 μM, we observed complete inhibition of the response in the shear assay (Supplementary Fig. 3A, light blue diamonds, 3 independent experiments with SEM). Supplementary Fig. 3B shows that responses to shear stress were similar in the stably transfected cell line and in transiently transfected cells, whereas cells expressing only mCherry produced no response (Supplementary Fig. 3B). Suppressing PIEZO1 expression with miRNAs also inhibited the response, but scrambled miRNA sequences had no effect (Supplementary Fig. 3B). These data are summarized in Panel B (Supplementary Fig. 3) and is an average of 3 independent experiments with SEM.

We examined whether the Aβ could inhibit PIEZO1 currents in the patch. We formed outside/in patches from the hP1 cell line and tested whether 10 μM of Aβ (1-40) or Aβ (1-42) (Peptide Method I) inhibited channel activity, a concentration that blocked the shear response. Surprisingly, neither peptide inhibited outside/in patch currents (Supplementary Fig. 5, represents 3 independent experiments).

Peptide binding specificity to the channel was assessed using the L and D enantiomers of Aβ(1-40) and Aβ(1-42), (c.f.22). For all these experiments, each data point is an average of 4 independent experiments with SEM. The ratio of inhibitory constants, Kᵢ, between the L and D Aβ peptides was ~300-fold for Aβ(1-40) (Fig. 1C, blue double arrow) and ~70-fold for Aβ(1-42) (Fig. 1C, red double arrow). Using the scrambled peptide as an estimate of non-specific binding, at least a > 10,000-fold difference between L and D forms was expected because the D-form will be incapable of specific interactions. Since we observed only a 300-fold difference, it indicated a mechanism that does not involve peptide-channel interactions. The monomeric peptide AgS (AggreSure L-Aβ(1-40); Fig. 1C, black squares) inhibited PIEZO1 with a Kᵢ about two orders of magnitude lower (~30 fM), and indicated that the sensitivity of enantiomeric peptides is related to state of the peptide (i.e. aggregate versus monomer).

### Table 1. Calculated Kᵢ for inhibition of hPIEZO1-mediated, shear stress-induced [Ca²⁺]ᵢ increase by enantiomeric Aβ peptides prepared by Method 1 (monomer, rows 2–5) or by Method 2 (oligomer, rows 6–9), and for the commercially available monomeric peptide, AggreSure 40 (row 1).

| Peptide     | Kᵢ [M]   | Kₑ₄₀ | n  | nₑ₄₀ | Adjusted R² |
|-------------|----------|------|----|------|-------------|
| AggreSure 40| 31.2 e-15| 6.8 e-15| 0.50 | 0.06 | 0.987       |
| L40 monomer | 51.5 e-15| 9.9 e-15| 0.49 | 0.05 | 0.997       |
| D40 monomer | 112.6 e-15| 14.3 e-15| 0.34 | 0.02 | 0.998       |
| L42 monomer | 261.5 e-15| 19.7 e-15| 0.40 | 0.04 | 0.986       |
| D42 monomer | 725.5 e-15| 98.8 e-15| 0.37 | 0.09 | 0.990       |
| L40          | 15.8 e-12| 2.58 e-12| 0.48 | 0.05 | 0.990       |
| D40          | 4.9 e-9  | 683 e-12| 0.34 | 0.03 | 0.998       |
| L42          | 345 e-12 | 49.11 e-12| 0.52 | 0.04 | 0.995       |
| D42          | 26.6 e-9 | 6.7 e-9  | 0.49 | 0.05 | 0.989       |

PIEZO1 inhibition by monomeric peptides. We prepared monomers of four Aβ peptides (see Peptide-Method II) and titrated their inhibitory effects on the shear response of hP1-CL cells (Fig. 1D, 4 independent experiments with SEM indicated for each concentration). The difference between the Kᵢ values of AgS peptide (~30 fM) and monomerized L-Aβ(1-40) (~52 fM) was negligible (Table 1). The Kᵢ for all the monomeric peptides were in the fM to sub-pM range, with Kᵢ values for enantiomeric pairs of Aβ(1-40) and Aβ(1-42) differing by only 2- to 3-fold (Table 1). These results are consistent with the assumption that peptides prepared by method I differed in oligomeric state (or conformation) from AgS in a way that reduced PIEZO1 inhibitory potency by 3–6 orders of magnitude. The similar efficacy of the D and L peptides suggests that they only interact with PIEZO1 through non-contact (lipidic) mechanisms.

We were unable to washout the channel inhibition within 20 min or by increasing the shear stress to 25 dynes/cm². In conventional solution kinetics, the upper limit for the association rate constant (kₐ) is usually considered to be ~10⁶ M⁻¹ s⁻¹, a diffusion-controlled limit. The dissociation rate constant (kᵢ) therefore should be less than 10⁻⁴ s⁻¹, and the time constant for achieving binding equilibrium at a peptide concentration ~Kᵢ will be on the order of 10⁴ s (a few hours). If the dissociation constant is 100 fM, the time constant would be about 10⁵ s (a day, or so). The fact that we achieved equilibrium in much shorter times suggests that the local membrane concentration was much higher than in solution.
Titration of the Aβ response showed a negatively cooperative Hill coefficient <1 suggesting that higher concentrations buffer the monomers into less effective oligomers. In contrast to Aβ inhibition, GsMTx4 had a much higher Kᵢ of ~250 nM²³,²⁴ with a Hill coefficient of ~ 1.9 (Supplementary Fig. 3C), and channel activity was readily restored by GsMTx4 washout (data not shown).

Co-localization of peptide and channel. Monomerized red fluorescent Aβ(1-42) peptide at 100 pM completely inhibited the PIEZO1-mediated Ca²⁺ response to shear stress (Fig. 2-Left Panel, 4 independent experiments with SEM). We incubated 100 pM of fluorescent Aβ with fluorescent hP1-CL cells for 10 min and investigated possible PIEZO1 co-localization with the peptide (Fig. 2-Right Panels marked A–F) using Structured Illumination Microscopy (SIM). Panels A–C are images near the bottom of the cell. Panel A is the peptide (red) and Panel B is the PIEZO1 channel (green). Panel C is the overlay of the red and green fluorescent channels showing yellow puncta where there is overlap indicating co-localization (indicated by arrows). Note that not all channels are associated with Aβ(1-42) peptide. Panels D–F are images near the top of the cell which also show co-localization of the peptide and PIEZO1 (indicated by arrows). Piezo proteins are known to segregate into spatial domains²⁵,²⁶ in the absence of Aβs, and the altered environment of the channels may favor Aβ association.

Aβ peptides affect PIEZO1-mediated cell migration. PIEZO1 channels are involved in neurite extension¹⁶ and other types of cell motility. This led us to test the effects of Aβ peptides on hP1-CL motility in a wound healing assay. Cells were grown to confluence on glass coverslips half-coated with PDMS (Polydimethylsiloxane). The PDMS layer was then peeled off allowing us to accurately measure the collective cell migration rates across the newly exposed glass surface.

Surprisingly, hP1-CL cells migrated ~10-fold faster than native HEK293T cells (Fig. 3 Top panel) showing the involvement of PIEZO1 channels in cell sheet movement. Addition of either D-GsMTx4 (5 µM) or enantiomeric forms of monomeric Aβ(1-40) (10 pM) slowed cell migration by over 10-fold, consistent with PIEZO1 involvement in cell movement (Fig. 3 Bottom left panel). Figure 3 Bottom right panel shows the average velocity for 3 independent experiments with SEM.

Aβ peptides are involved in cytoskeletal remodeling. Cell movement in the presence of PIEZO1 channels appears to rely on cytoskeletal integrity and plasticity, consistent with the ability of cytoskeletal disrupting agents to prevent PIEZO1 activation (Supplementary Fig. 4A). We examined forces within the cytoskeleton during channel inhibition. By transient transfection, we introduced a genetically encoded optical probe of actin stress (cpst-FRET) into stable hP1-mCherry-CL cells¹⁹ (Fig. 4A), resulting in probe expression into F-actin. The fluorescence spectrum of hP1-mCherry-1591 does not significantly overlap with the spectrum of the actin force
For each experiment described below, we analyzed and averaged (with SD) 15 cells from 3 different experiments. After establishing baseline actin stress for 30 min, addition of the peptides resulted in an increase in actin stress (Fig. 4B). 10 pM of monomeric L-Aβ(1-40) (red circles) appeared slightly more effective than D-Aβ(1-40) (blue triangles) or 5 µM D-GsMTx4 (black squares) in increasing actin-associated tension. Figure 4C shows that the fluid shear stress associated with a simple bath exchange had no effect on actin stress (red circles). 10 pM of monomerized scrambled Aβ peptide (black squares) also had no effect on actin stress. However, removal of Ca2+ from the extracellular bath greatly increased the resting actin stress (blue triangles). The steady state changes in actin stress (at t = 60 min) are summarized in Fig. 4D.

Interestingly, FRET images of hP1-mCherry-CL cells show lamellipodia containing the actin probe (Fig. 4E). Addition of D-GsMTx4 or Aβ(1-40) enantiomers led to a drop of F-actin tension and its retraction from the lamellipodia (Fig. 4E,F). Our results support a model where Ca2+ influx through PIEZO1 channels maintains the cytoskeletal plasticity required for cell migration. Control experiments with native HEK293T cells revealed no effects on actin tension by bath Ca2+ removal, D-GsMTx4, or Aβ(1-40). Three independent experiments were averaged (SEM).

Discussion

An important advantage to the shear stress approach, relative to techniques such as patch clamp27, is the ability to measure PIEZO1 channel function without altering cellular integrity. We know that the channel's environment clearly affects the channel's response to mechanical stress21. Assays intended to mimic the in situ situation must leave the environment minimally altered. We have shown that monomeric Aβ peptides can block PIEZO1 activity at fM concentrations, and that other forms of the peptides (i.e. aggregated) are less potent. Using enantiomers we demonstrated that the Aβ effects are dominated by long range effects that are not stereospecific similar to the observations seen with GsMTx422.

We examined the electrophysiological effects of Aβs using patch clamp. In control cells, channel activity in response to stretch was obvious18,21. We could not observe PIEZO1 inhibition by Aβ peptides at concentrations that inhibited the response to fluid shear stress. The inability to observe patch current correlations with the extreme sensitivity of the shear stress assay is a warning that extrapolation of patch results does not always apply.
In situ situations. The stresses in patch experiments may alter the binding of the $A\beta$ s or the ability of the peptide perturbations to reach the channels. How may $A\beta$ s affect channel function? Mouse PIEZO1 gate by local bilayer tension. In neurons, PIEZO1 activity appears concentrated in cholesterol-rich lipid raft domains containing Stomatin-like 3 protein (STOML3). These membrane nanostructures can redirect the mechanical forces that modulate MSCs. The insertion of $A\beta$ peptides into a confined lipid domain compresses the surrounding lipid and channels. Alternatively, $A\beta$ disrupts the boundary lipids of PIEZO1, as suggested by cryoEM. Perhaps $A\beta$ s perturb these structural lipids rendering the channel inactive. $A\beta$ peptides are known to concentrate in lipid rafts. The extremely low concentration of monomeric peptide needed to inhibit channel function (fM) may reflect a partition coefficient that concentrates monomeric peptides in constrained domains containing PIEZO1.

Another possibility is that $A\beta$ affects PIEZO1 activity indirectly. Activation of PIEZO1 channels may result from forces created through the cytoskeleton. Cells generate traction forces by interactions with the extracellular matrix (ECM) and can use integrin rich focal adhesion regions to transmit tension to the bilayer activating...
PIEZO1 channels. The connection between PIEZO1 (Fam38a) and integrins was demonstrated prior to the discovery of PIEZO1’s mechanical channel properties and activation of PIEZO1 channels through the ECM was shown by Poole et al. The disruption of PIEZO1 activity by Aβ would be attributed to inhibition of any step along a signaling pathway.

Aβ peptides vary considerably in their structure and oligomeric states. Moreover, opinions vary regarding the toxicity of the various peptide forms, ranging from soluble dimers to fibers. Recent work suggests that Aβ monomers are random coils.

Although Aβ peptides are often associated with Alzheimer’s disease, monomeric Aβ peptides at low concentrations can exert positive physiological effects on synaptic plasticity and neuronal survival. Giuffrida et al. reported the interesting observation that monomeric Aβ (1–42) is neuroprotective, preventing trophic deprivation in developing neurons and protecting mature neurons against excitotoxic death at low concentrations.

We have shown the connection between PIEZO1-mediated Ca²⁺ influx and cytoskeleton remodeling and cell mechanics. The elevation in Aβ peptide concentration following head trauma might be intended to limit the PIEZO1 response to mechanical activation. Consistent with our results on the loss of efficacy with oligomer formation, Giuffrida et al. suggested that oligomerization decreases the concentration of monomers, and may explain a negative Hill coefficient (Table 1). These results also suggest that other amphipathic molecules may also exhibit environmentally sensitive affinities as a function of bilayer tension.

The coupling of PIEZO activity to motility is clear from the effects of over-expression in the HEK293T cells, and inhibition by Aβ peptides. Collective cell migration is related to tissue remodeling events involved in wound healing and cancer, and is distinct from anchorage-independent (“amoeboid”) cell migration observed in the setting of Fam38a (PIEZO1) depletion. Linking the flux of Ca²⁺ through PIEZO1 to cytoskeletal changes allows for rapid and coordinated cell migration. The ubiquitous expression of Aβ peptide raises the question of how it may influence mechanical responses throughout the organism.

**Methods**

**PIEZO1 cell lines.** The vector hPI-1591-EGFP-mCherry with an N-terminal his tag was amplified using a BamH1 forward primer and EcoR1 reverse primer. The gel purified PCR product was used in an InFusion reaction with pBabe puro vector DNA that had been cut with BamH1 and EcoR1. The resulting construct, pBabePuro-NHis-hP1-1591-EGFP, was treated with restriction enzymes MluI and XhoI. A PIEZO1 cDNA fragment containing inserted EGFP at amino acid residue 1591 and with MluI and XhoI ends was inserted by ligation (pBabePuro-NHis-hP1-1591-EGFP). Integrity of changes was confirmed by DNA sequencing.

**Primers.**

InfHP1BabeBamHisF – GGCGCCGGCCGGATCCTCAGCCACCATGCACCATCATC
InfHP1BabeR1rev – CACCGGTACTGAATTCCTACTACTCTCACGAGTCC.

**Generation of stable cell line.** To generate viral particles, AmphoPak cells in 30 mm dishes were grown to 25% confluency and transfected with DNA (1 µg per dish) using Mirus transfection reagent according to manufacturer’s recommendation. After 2 days, the media was collected and filtered through a 45 micron filter.

1 ml freshly harvested virus plus 2 µg/ml polybrene were added to adherent HEK293T cells grown to 25–50% confluency in 30 mm dishes. Fresh medium was added after 3 hrs, and the cells were allowed to incubate overnight. Lentiviral infection was repeated after 24 hours. The twice infected cells were allowed to incubate for an additional 24 hours, and subjected to puromycin selection (Sigma) at four concentrations between 0.25 to 4.0 µg/ml. Single colonies were generated from cells grown at 2.8 µg/ml puromycin. The resulting clonal hPI-CL cell line was used in subsequent experiments.
**Shear stress assay.** The microfluidic chambers had glass coverslip bottoms coated with human fibronectin (Invitrogen) and fluid flow guided by PDMS channels. Cells were cultured in the chamber for 3 days, with daily medium changes. On the experimental day, chambers were rinsed with isotonic solution (75 mM NaCl, 5 mM KCl, 2 mM MgCl₂, 1 mM CaCl₂, 10 mM HEPES, adjusted to 320 mOsm with mannitol). A stock of Flu-4 AM was diluted to 5 µM in isotonic saline and incubated with cells for 10–30 min at 37°C. Chambers were then rinsed twice with isotonic solution and incubated for 8 minutes to completely Flu-4 AM de-esterification. A single fluid shear stress pulse of ~15 dynes/cm² for 8–10 ms was applied to the chamber using a high-speed pressure servo (HSPC-1 from ALA). Peptide stocks were kept at 4°C until used, dissolved to desired concentrations, and immediately added to cells. Each chamber had a minimum of 15 cells. Images of the cellular response to shear stress were captured by an EM-CCD camera (C9100 model, Hamamatsu) at a nominal rate of 1 Hz. To quantify the response, the average intensity in the field of view was background subtracted. Image intensity was adjusted for the rate of bleaching. Each condition was repeated in four separate chambers, and the results averaged. The responses were normalized to the control experiments, and the inhibition was calculated as previously described. The data points were fit by a Hill plot with the equation:

\[ y = \frac{V_{max} \cdot x^n}{(k^n + x^n)} \]  

where \( V_{max} \) is the maximum response, \( n \) is the Hill coefficient, and \( k \) is related to the concentration at half-maximal response. The extracted parameters are shown in Table 1.

**Electrophysiology.** Cell-attached and outside-out patches were used as previously described. All experiments were at room temperature. Pressure or suction steps were applied to the membranes using a high speed pressure clamp (ALA). The bath solution contained (in mM): 150 KCl, 10 HEPES, 1 MgCl₂, 1 CaCl₂ at pH 7.4 that clamped the resting membrane potential to zero. The pipette solution was 150 mM KCl at pH 7.4.

**microRNAs.** Two microRNAs targeting PIEZO1 expression were cloned using the BLOCK-IT expression vector kit (Invitrogen) according to the manufacturer’s specifications. The following primers were used:

338_top  TGCTGTAGACAATCTTAGACGTGTCAGTCAGTGGCCAAAACGGTCGTCAAGTA
338_bot  CCTGTAGACAATCTTAGACGTGTCAGTCAGTGGCCAAAACCGTGGTCTAAGATT
3767_top TGCTGTAGACAATCTTAGACGTGTCAGTCAGTGGCCAAAACCGTGGTCTAAGATT
3767_bot  CCTGTAGACAATCTTAGACGTGTCAGTCAGTGGCCAAAACCGTGGTCTAAGATT

The two miRNAs were introduced into one vector (chaining) according to the manufacturer’s specifications.

**Cell Migration.** Half of a glass coverslip was coated with catalyzed PDMS (polydimethylsiloxane) (9:1) and allowed to harden. Cells were cultured on the complete coverslip kept in tissue culture dishes. After achieving confluency, the PDMS was peeled from the glass. The location of the front edge of the confluent cells was monitored for 5 hr at 37°C with 5% CO₂-supplemented DMEM media containing 10% bovine serum. Test peptides were added to the culture media at the start of the 5 hr experiment. To analyze the average migration speed cell front, images were processed using Cell Profiler (open source software, [http://cellprofiler.org/](http://cellprofiler.org/)) to measure the area covered by the cells in each frame. Dividing by the width of the frame, an average (with SEM) speed of movement in a direction perpendicular to the cell front was calculated. The datasets generated and/or analyzed during the current study are available from the corresponding author on request.

**Super resolution microscopy.** Samples were imaged on a Visitech VTI-iSIM attached to a Nikon TE2000 inverted microscope. Images of samples excited at 488 nm and 568 nm were captured at 60 z-intervals of 200 nm.

**Cytoskeleton stress probe.** To measure cytoskeletal stress, hP1-mCherry-CL cells were transiently transfected with cDNA encoding the cpst-actin FRET probe. The FRET ratio of 528 nM and 475 nM was measured by simultaneous imaging using an image splitter. Baseline actin stress was measured at 37°C in isotonic solution for 30 min before peptide addition (see above).

**Transient transfection.** Transient transfections were done at least 24 hours before experiments using Mirus TransIT®-293 reagent according to manufacturer’s specification.

**Peptide preparation.** The enantiomeric forms of human Aβ(1-40) and Aβ(1-42) peptides were chemically synthesized by and purchased from Anaspec or Bachem. The scrambled Aβ(1-40) peptide, HiLyte (555) fluorescent peptide Aβ(1-42), and AggreSure peptide were purchased from Anaspec.

**Peptide preparation methods. Method I.** To one mg of peptide, 70 µl 1.0% NH₄OH was added. The peptide was then diluted with bath buffer (see above) to a concentration of approximately 1 mg/Ml, and gently vortexed and frozen at −80°C. This method generated oligomeric peptide.
Method II. (monomer procedure43): One mg of peptide was dissolved in 0.5 ml TFA (trifluoro acetic acid) and sonicated for 10 min. TFA was removed with streaming argon after which 800 μl of 1,1,3,3,3-Hexafluoro-2-propanol (THIP) (Sigma-Aldrich) was added and incubated 1 hr at 37 °C. This solvent allows aggregated peptides to dissociate into monomers by interfering with interactions used to stabilize aggregate peptides. The solvent was removed by streaming argon, and 800 THIP was added and frozen to −80 °C. The material was lyophilized and dissolved in 5 mM DMSO (Sigma-Aldrich) and then diluted in buffer. All peptides were stored at −80 °C.

Peptide sequences were as follows:

\[
\begin{align*}
\text{A}\beta(1-40) & \quad \text{DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGV} \\
\text{A}\beta(1-42) & \quad \text{DAEFRHDSGYEVHHQKLVFFAEDVGSNKGAIIGLMVGGV} \\
\text{A}\beta(1-40) & \quad \text{Scrambled AEHDHSVKLKEGAYMIFVDQGHVF} \\
\end{align*}
\]

Data Availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author on reasonable request.

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Author Contributions
M.M.M. performed all shear stress experiments and analyzed the data. L.Z. performed all cloning experiments. F.S. and S.H. edited the manuscript. P.A.G. designed the experiments, was involved in the analysis, drafting, and editing of the manuscript as well as performing experiments. All authors discussed various aspects of the project.

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