Binding of the protein ICln to $\alpha$-integrin contributes to the activation of ICl$_{swell}$ current

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ICl$_{swell}$ is the chloride current induced by cell swelling, and plays a fundamental role in several biological processes, including the regulatory volume decrease (RVD). ICln is a highly conserved, ubiquitously expressed and multifunctional protein involved in the activation of ICl$_{swell}$. In platelets, ICln binds to the intracellular domain of the integrin $\alpha$IIb chain, however, whether the ICln/integrin interaction plays a role in RVD is not known. Here we show that a direct molecular interaction between ICln and the integrin $\alpha$-chain is not restricted to platelets and involves highly conserved amino acid motifs. Integrin $\alpha$ recruits ICln to the plasma membrane, thereby facilitating the activation of ICl$_{swell}$ during hypotonicity. Perturbation of the ICln/integrin interaction prevents the transposition of ICln towards the cell surface and, in parallel, impedes the activation of ICl$_{swell}$. We suggest that the ICln/integrin interaction interface may represent a new molecular target enabling specific ICl$_{swell}$ suppression in pathological conditions when this current is deregulated or plays a detrimental role.

Cellular volume changes occur in a variety of physiological processes, including transepithelial transport, cell migration, proliferation, and death. As limiting excessive or prolonged volume changes is essential for normal cell function and survival, most cell types are able to counteract volume perturbations by initiating the homeostatic processes of regulatory volume increase and decrease (RVI and RVD, respectively)\cite{1,2,3,4}. Activation of a chloride conductance upon cell swelling (ICl$_{swell}$) is a key step in RVD and occurs via volume regulated anion channels (VRACs)\cite{5,6}.

The mechanisms of osmosensing are complex and far from being completely understood. Several lines of evidence implicate integrins as upstream sensors of cell volume perturbations in mammalian cells after cell swelling or shrinkage\cite{7,8,9,10,11}. Integrins are a highly conserved family of heterodimeric adhesion molecules that consist of an $\alpha$- and a $\beta$-subunit and connect the extracellular matrix to intracellular signalling proteins and the cytoskeleton, and play a critical role in many vital cellular functions, such as cell adhesion, migration, invasion, differentiation, proliferation, apoptosis, growth-factor signalling and response to hypotonicity\cite{12,13,14,15}. By linking the exterior and interior space of cells, integrins permit a bidirectional transmission (outside-in and inside-out signalling) of mechanical and biochemical stimuli across the plasma membrane\cite{16,17}. Specifically, activation of integrins by hypotonicity\cite{18} seems to trigger Cl$^-$ and organic osmolyte fluxes during RVD, possibly by recruitment of a member of the SRC family of kinases\cite{19}. Integrins also establish an intricate network of molecular interactions via their intracellular domains. The $\beta$-subunits are major players in this context, and bind signalling hubs such as talin and kindlins\cite{20,21}. The intracellular network of the $\alpha$-subunits was less extensively studied. Evidence was gathered that the intracellular domain of the platelet-specific integrin $\alpha$IIb physically interacts with the protein ICln. This interaction is part of the molecular events underlying platelet activation and capacity to form a thrombus\cite{22,23}.

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ICln is a highly conserved, ubiquitously expressed and multifunctional protein that plays a critical role in cell homeostasis and survival. ICln was suggested to be the channel or part of the macromolecular complex underlying the activation of IClswell. Despite being mainly located in the cytosol in resting cells, ICln is transposed towards the plasma membrane upon hypotonic challenge or other stimuli. In addition, ICln establishes complex interactions with diverse proteins belonging to different subcellular compartments and functional modules. For instance, ICln regulates the process of splicing and small nuclear ribonucleoproteins (snRNP) biogenesis by associating with several Smith antigen (Sm) proteins of the spliceosomal core and controlling their methylation.

The cytoplasmic tail of human integrin αIIb contains the highly conserved motif (1020)KVGGFFKR(1026) (single letter amino acid code; the numbers indicate the amino acid positions within the primary sequence of human αIIb integrin; NCBI Reference Sequence: NP_000410.2), which regulates integrin activation and acts as a recognition site for various intracellular proteins, including ICln. As the KxGFFKR motif (x denotes any amino acid) is common to several α integrin isoforms, it is important to understand whether the ICln/integrin interaction can also occur in nucleated cells and identify the corresponding function. Here we show that the interaction between ICln and the intracellular domain of integrin α-chain is essential for the activation of IClswell and is independent from the occupancy of integrin α by extracellular ligands.

Results
ICln establishes a direct molecular interaction with the intracellular domain of the integrin αIIb chain. To verify whether the ICln-integrin interaction is restricted to the process of plaquelet activation, we performed fluorescence resonance energy transfer (FRET) experiments in living NIH-3T3 fibroblasts kept in isotonic solution and transiently transfected with Canis familiaris ICln and the transmembrane and intracellular portion (amino acids 987–1039) of the human integrin αIIb chain (αIIb-short, αIIb) (Fig. 1A,B). Deletion of the extracellular portion of αIIb avoided perturbation of a possible ICln/αIIb interaction by an extracellular signal. The cDNA sequences coding for ICln and αIIb were cloned into vectors to produce fusion proteins with fluorophores suitable for FRET experiments, i.e. CFP-ICln and αIIb-YFP respectively (see Methods). After expression, αIIb-YFP clearly targeted to the plasma membrane, with a partial retention in intracellular compartments (Fig. 1A and Supplementary Fig. S1). The FRET efficiency of CFP-ICln and αIIb-YFP determined in the plasma membrane region of transfected cells following photobleaching of the FRET acceptor YFP was significantly higher (p < 0.01) with respect to that of the corresponding negative control (expression of CFP alone and αIIb-YFP) (Fig. 1B, Supplementary Table S1 and Fig. S1). Similar experiments performed on NIH-3T3 fibroblasts fixed in 3% paraformaldehyde and kept in a phosphate buffered saline led to similar results (data not shown). Also in HEK293 Phoenix cells, the FRET efficiency of CFP-ICln and αIIb-YFP determined in the plasma membrane region was significantly higher than that determined in control cells transfected with CFP and αIIb-YFP (Supplementary Table S1). The ratio of the FRET efficiency between CFP-ICln and αIIb-YFP and the stochastic interaction given by the FRET efficiency between CFP and αIIb-YFP in the two cells types was similar, being 2.9 and 2.7 in NIH-3T3 and HEK293 Phoenix cells, respectively. Accordingly, in sensitized emission N-FRET experiments, ICln and αIIb produced an N-FRET in isotonic solution that was significantly higher (p < 0.01) compared to the control (Fig. 1C,D, isotonic, and Supplementary File S1). Following exposure of cells to a hypotonic bath solution, the N-FRET of ICln and αIIb significantly increased over time, while FRET of CFP and αIIb-YFP did not significantly change (Fig. 1C,D and Supplementary File S1). These results show that in isotonic, as well as hypotonic conditions, ICln is located in close proximity to αIIb and therefore strongly suggest a direct molecular interaction between the two molecules.

Over-expression of αIIb activates IClswell. To test whether the interaction between ICln and the integrin α chain might have a functional implication on IClswell, the whole-cell patch-clamp technique was used. HEK293 Phoenix cells were transfected with the pIRE2-cGFP vector with or without αIIb, cDNA (see Methods). The bicistronic αIIb-pIRE2-cGFP vector was used to obtain the simultaneous expression of αIIb, and the transfection marker EGFP as two independent proteins in the same cell, thus allowing for the identification of cells expressing αIIb, by fluorescence microscopy. After the seal was realized and the whole-cell configuration was obtained, IClswell was elicited by reducing the extracellular osmolarity. Hypotonic shock induced the activation of a large chloride current with the biophysical fingerprints of IClswell, characterized by outward rectification and slow voltage- and time-dependent inactivation at positive potentials in both control and αIIb, over-expressing cells (Fig. 1E). The current density-to-voltage relation determined after a 10 min exposure to extracellular hypotonic solution (Fig. 1F), as well as the current density-to-time relation (Fig. 1G), showed a strong up-regulation of IClswell in cells over-expressing αIIb compared to control cells (p < 0.0001). These results suggest an involvement of the αIIb chain in the activation of IClswell.

The cell-permeable (1020)KVGGFFKR(1026) integrin peptide impairs the translocation of ICln to the plasma membrane and the activation of IClswell. The translocation of ICln to the plasma membrane after hypotonic challenge was verified by sensitized emission N-FRET in living NIH-3T3 fibroblasts exposed to the synthetic palmitoylated peptide (1020)KVGGFFKR(1026), corresponding to the amino acid sequence of the ICln recognition domain within the integrin αIIb chain. Control cells were treated with the synthetic palmitoylated peptide KAAAAAR. Palmitoylation allows the peptides applied to the extracellular solution to access the intracellular environment, with preferential, although not exclusive, targeting to the plasma membrane. Cells expressed the fusion protein CFP-ICln and the membrane label YFP-mem (i.e. the YFP protein tagged with an N-terminal palmitoylation sequence, see Methods). As previously reported, hypotonicity induced a significant increase of FRET between CFP-ICln and YFP-mem, thereby indicating a transposition of ICln towards the cell membrane (Fig. 2A). Interestingly, pre-incubation of cells with the palmitoylated (1020)KVGGFFKR(1026) peptide inhibited significantly the plasma membrane translocation of ICln in both control and αIIb over-expressing HEK293 Phoenix cells (Fig. 2B, C, Supplementary Table S1 and Supplementary File S1). These results suggest that the presence of αIIb is required for the ICln plasma membrane translocation upon hypotonic challenge, probably by targeting ICln towards the plasma membrane by a mechanism dependent on a direct molecular interaction between ICln and αIIb.

ICln and ICln/αIIb ratios were also studied by transient expression in HEK293 Phoenix cells, where the αIIb chain is essential for the activation of IClswell, thereby strongly suggesting a direct molecular interaction between the two molecules.
Figure 1. The integrin α-chain interacts with ICln and facilitates the activation of IClswell. Living NIH-3T3 fibroblasts were transiently transfected with *Canis familiaris* ICln and αIIb, and bathed in isotonic solution. (A) The FRET donor (CFP) and the FRET acceptor (YFP) were imaged before and after photobleaching of the acceptor. The green squares correspond to the bleaching ROI and the right panel represents the FRET efficiency image. (B) FRET efficiency of CFP-ICln and αIIb-YFP or CFP and αIIb-YFP determined in three different plasma membrane regions of interest (ROIs) for each cell. n represents the number of cells. **p < 0.01, unpaired Student’s t test. (C) FRET signal of CFP-ICln and αIIb-YFP or CFP and αIIb-YFP determined by sensitized emission in living NIH-3T3 fibroblasts kept in isotonic or hypotonic solution for 10 and 20 min. Bar: 10 μm. (D) N-FRET of CFP-ICln and αIIb-YFP (n = 8) or CFP and αIIb-YFP (n = 6) in isotonic solution or after 10, 15 and 20 min of hypotonic stress. Data are normalized for the value of N-FRET of CFP-ICln and αIIb-YFP measured in isotonic solution and expressed in %. Hypotonic versus isotonic: *p < 0.05, **p < 0.01, paired Student t test; CFP-ICln versus CFP: °°p < 0.01, unpaired Student t test. n indicates the number of cells.
peptide not only prevented the expected FRET increase following hypotonic stress, but indeed led to a FRET decrease. This indicates that the peptide not only abolished the translocation of ICln normally observed in hypotonicity, but additionally decreased the amount of the membrane-bound ICln compared to isotonic conditions (Fig. 2B). In contrast, preincubation of cells with the palmitoylated KAAAAAR control peptide did not affect the hypotonicity-induced translocation of ICln towards the plasma membrane (Fig. 2C). Importantly, patch-clamp experiments in whole-cell configuration showed that the impaired translocation of ICln toward the cell surface was paralleled by an impaired activation of IClnswell. Specifically, the palmitoylated (1020)KVGFKF(1026) peptide applied to the bath solution, and not the palmitoylated KAAAAAR peptide, dramatically impaired (Fig. 2D) or prevented (Fig. 2E) the activation of IClnswell in hypotonicity-stimulated HEK293 Phoenix cells. Larkin et al. previously speculated that the amino acids (84)AKFEEE(89) of ICln might be involved in binding the (1020)KVGFKF(1026) motif22. However, the cell-permeable peptide (84)AKFEEE(89) did not significantly affect IClnswell activation compared to the unrelated cell-permeable peptide ELFNDG (Fig. 2F). Therefore, we concluded that the ICln amino acids (84)AKFEEE(89) unlikely represent the main integrin interacting site.

**The (61)ISLHA(65) ICln motif is the prime integrin binding site.** In order to identify the binding site of the integrin (1020)KVGFKF(1026) motif in ICln, we applied 1H−15N HSQC spectroscopy. Binding of the (1018)MWKVGFFKRNR(1028) integrin peptide to ICln was probed by using both full-length *Canis familiaris* ICln (cICln, Fig. 3A) and a truncated form (TeICln, Fig. 3A) lacking the C-terminal part. TeICln folds into a pleckstrin homology domain-like structure consisting of two nearly orthogonal antiparallel β-sheets forming a β-barrel capped by an α-helix near the C-terminus (60) and Fig. 3C). Figure 3B shows an overlay of NMR spectra of unligated (green) and ligated (red) TeICln. It can be seen that most of the residues are unchanged and only a subset of residues labeled by their position in the primary sequence) display small but significant chemical shift changes. The most pronounced chemical shift changes were observed for S62 (shown in red in Fig. 3A,C), I61 and A65 (shown in orange in Fig. 3A,C). Most interestingly, a continuous stretch of residues displayed significant chemical shift changes (I61-A65). Additionally, the following residues (shown in yellow in Fig. 3A) exhibited chemical shift changes: S55, Y58, L63, Y79, V80, A84, S90 and residues in the C-terminal region (A91-L98). The most pronounced chemical shift changes were observed for A65 (shown in red in Fig. 3A,C). Figure 3A also refers to D. (E) Original recordings obtained by whole-cell patch-clamp in hypotonic (top) or isotonic (bottom) conditions in control (left panels) or αIIb, transfected (right panels) HEK293 Phoenix cells stimulated with voltage increments as depicted in the lower inset. Cells over-expressed both αIIb and the transfection marker EGFP as separate proteins, or EGFP only. (F) The current density-to-voltage relationships measured 10 min following hypotonic shock, and G, the current density-to-time relationships were fitted with second order polynomials, following application of the extra-sum of squares F test (αIIb, hypotonic, versus control, hypotonic: p < 0.0001).

We set out to verify whether a peptide comprising the ICln motif (61)ISLHA(65) could affect the transposition of ICln to the plasma membrane and the activation of IClnswell. The translocation of ICln to the plasma membrane after hypotonic challenge was investigated by sensitized emission N-FRET in living NIH-3T3 fibroblasts transfected with the fusion protein CFP-ICln and the membrane label YFP-mem. Pre-incubation of cells with 50 μM of the cell-permeable (palmitoylated) synthetic peptide (57)EYPTISLHALSRDR(70) (pal-ICln peptide; numbers indicate the amino acid positions within the primary sequence of hICln) encompassing the (61)ISLHA(65) motif not only abrogated the FRET increase between CFP-ICln and YFP-mem normally observed after hypotonic stress, but led to a FRET decrease compared to the isotonic condition (Fig. 4A,B), again indicating an interference with ICln transposition and localization to the plasma membrane. Accordingly, patch-clamp experiments in whole-cell configuration showed that the
palmitoylated (57)EYPTISLHALSRDR(70) peptide applied to the bath (extracellular) solution (Fig. 4C,E), and not a palmitoylated peptide with a scrambled amino acid sequence (RIYSASDLEPRLTH, pal-control, Fig. 4D,E), strongly inhibited ICl swell in HEK293 Phoenix cells. The effect of the peptide (57)EYPTISLHALSRDR(70) on IClswell was also investigated by adding it to the pipette filling (intracellular) solution used in patch-clamp experiments. This manoeuvre allows the peptide to gain access to the intracellular space after establishing the whole-cell configuration. The non-palmitoylated (57)EYPTISLHALSRDR(70) peptide did not interfere with the activation of the current (Fig. 5A). In contrast, adding the palmitoylated peptide to the intracellular space significantly reduced IClswell (Fig. 5B).

Integrity of the (61)ISLHA(65) motif is essential for the association of ICln to the αiib-chain and the activation of IClswell. To further confirm the identity of the ICln amino acid(s) involved in the interaction with the integrin α chain, amino acid substitutions (I61G, S62G, A65N) were made within the integrin-recognition domain of Canis familiaris ICln (cICln, Fig. 6A) and FRET was determined by sensitized emission in NIH-3T3 fibroblasts transiently transfected with wild type (wt) or mutated (mut) CFP-ICln and αIIb-YFP. The FRET between CFP-ICln mut and αIIb-YFP measured in the plasma membrane region in isotonic and hypotonic solutions was significantly lower with respect to that measured between CFP-ICln wt and αIIb-YFP (Fig. 6B). These results confirm that the ICln amino acids I61, S62 and A65 are critically involved in the interaction with αIIb. To investigate the functional consequences of the impaired ICln/integrin interaction, IClswell was measured by patch-clamp experiments in whole-cell configuration in HEK293 Phoenix cells.
transfected with ICln wt, ICln mut or a control vector. Similar to previous results\textsuperscript{23}, overexpression of ICln significantly up-regulated ICl\textsc{swell}. In contrast, in cells expressing ICln mut, ICl\textsc{swell} did not differ from the endogenous current measured in control cells (Fig. 6D,E). ICln mut resulted functionally impaired compared to the wild type.
also following expression in NIH-3T3 fibroblasts (Supplementary Fig. S2). These results, together with those shown in Figs 2, 4 and 5, strongly indicate that interfering with the ICln/integrin interaction impairs the activation of IClswell.

Discussion

Although integrins are in general recognized as osmosensors in RVD\textsuperscript{7,42}, the molecular mechanisms by which they influence the swelling-activated chloride current IClswell are incompletely understood. This lack of information was due in part to the molecular identity of VRACs remaining elusive for more than 30 years\textsuperscript{43,44}. In 2014, two independent laboratories identified the leucine-rich repeat containing protein 8 A (LRRC8A) as an essential component of the channel that mediates ICl swell\textsuperscript{45,46}. Recently, the structure of the human and Mus musculus homo-hexameric LRRC8A channel was described by cryo-electron microscopy\textsuperscript{47–50}. Overall, these studies provide the basis for understanding the precise mechanism of activation of VRACs and identifying their molecular partners. LRRC8A and other members of the LRRC8 family form heteromers, of which the composition controls the VRAC single-channel conductance, rectification, anion selectivity and inactivation kinetics\textsuperscript{51}. When reconstituted into lipid bilayers in a two-compartment system, purified LRRC8 complexes formed anion channels activated either by osmolality gradients between two compartments both containing hyperosmolar solutions, or low ionic strength in the absence of osmotic gradients between the two compartments\textsuperscript{51}. These findings suggest that LRRC8 complexes may include a sensor for osmolarity and low ionic strength. However, other authors recently showed that low ionic strength is neither sufficient to activate VRAC on intracellular compartments, nor indispensable to keep plasma membrane-localized VRACs active, and rather suggested that the diacylglycerol/protein kinase D pathway specifically and potently activates VRAC\textsuperscript{52}. Therefore, in a cellular context, other factors,
including binding partners, post-translational modifications\(^5\), second messengers or even other channels\(^6\), may contribute to VRAC activation besides a mere reduction in the intracellular ionic strength.

The composition of the extracellular matrix modulates the characteristics of \(I_{\text{Cl}\text{swell}}\) through integrin receptors, and knock-down of \(\beta_1\) integrin impaired \(I_{\text{Cl}\text{swell}}\) activation in skeletal muscle-derived C2C12 cells\(^5\). The integrin \(\beta\) chains seem to play a fundamental role in this context. The research group of Browe and Baumgarten showed that mechanical stretch of \(\beta_1\) integrin leads to the activation of a chloride current with the biophysical and pharmacological fingerprints of \(I_{\text{Cl}\text{swell}}\)\(^5\)–\(^6\), thus suggesting that the role of integrins in RVD would be an outside-in transduction of a mechanical stimulus with \(I_{\text{Cl}\text{swell}}\) as the final effector. Accordingly, knockdown of integrin \(\beta_1\) also reduced RVD of adherent Ehrlich ascites cells\(^4\), and knockdown of integrin \(\beta_3\) almost completely blocked \(I_{\text{Cl}\text{swell}}\) activation in basilar artery smooth muscle cells\(^6\). As integrin \(\beta 3\) co-precipitated with Src and the ClC-3 protein, these authors suggested that the integrin \(\beta 3\)/Src/ClC-3 signalling pathway might be involved in the activation of \(I_{\text{Cl}\text{swell}}\)\(^6\).

In order to elucidate the molecular events underlying \(I_{\text{Cl}\text{swell}}\) activation, we set out to explore the role of the interaction between integrins and ICln. ICln interacts with the (1020)KVGFFKR(1026) motif of the platelet-specific integrin \(\alpha I I b\), thereby influencing platelet activation\(^2\),\(^2\)–\(^4\). We found that ICln can interact with the \(\alpha\) chain also in cells unrelated to platelets, in isotonic as well as hypotonic conditions (Fig. 1A–D). Moreover, overexpression of the transmembrane and intracellular portion of the \(\alpha I I b\) chain (\(\alpha I I b\)) upregulated \(I_{\text{Cl}\text{swell}}\) (Fig. 1E–G). Omission of the extracellular portion of the integrin \(\alpha I I b\) chain was adopted to avoid modulation of \(I_{\text{Cl}\text{swell}}\) by a possible alteration of cell adhesion to the extracellular substrate, thereby allowing for an unbiased exploration of the intracellular signalling events towards \(I_{\text{Cl}\text{swell}}\) activation. Importantly, this approach showed

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**Figure 5.** The palmitoylated (57)EYPTISLHALSRDR(70) ICln peptide applied to the intracellular solution impairs the activation of \(I_{\text{Cl}\text{swell}}\). Current density-to-voltage (left) and current density-to-time (right) relationships determined by patch-clamp in whole cell configuration in HEK293 Phoenix cells in hypertonic solution and after 20 min of hypotonic shock with 50 \(\mu\)M of the (A) unmodified or (B) palmitoylated ICln peptides (EYPTISLHALSRDR) or control peptides (RIYSASDLEPRLTH) in the pipette filling (intracellular) solution. Data were fitted with second order polynomials, following application of the extra-sum of squares F test (hypotonic, ICln peptide versus control peptide: n.s.; pal-ICln peptide versus pal-control peptide: \(p < 0.0001\)).
that the ICln/integrin interaction occurs regardless of the activation state or occupancy of the integrin α-chain by extracellular ligands. These findings are in agreement with those of Larkin et al. which showed that ICln bound equally to purified αIIb and to integrin from resting or activated platelets, and that binding was not affected by direct integrin activation with Mn²⁺ or by inhibitors of integrin occupancy.

The primary sequence of the site of the integrin αIIb-chain responsible for binding to ICln was previously determined to be represented by the highly conserved (1020)KVGFFKR(1026) amino acid motif, and was confirmed in the present study (Fig. 2). However, the identity of the ICln site involved in binding to the

Figure 6. Modification of the (61)ISLHA(65) motif impairs the activation of IClswell. (A) Sequence of the amino acids 57–70 of human (h), Canis familiaris (c) wild type (wt) and mutated (mut) ICln. In the latter, the amino acids establishing the strongest interaction with the integrin peptide were mutated as indicated. (B) Living NIH-3T3 fibroblasts were transfected with Canis familiaris wild type (wt) or mutated (mut) CFP-ICln and αIIb-YFP. N-FRET of CFP-ICln wt and αIIb-YFP (n = 13) or CFP-ICln mut and αIIb-YFP (n = 9) was determined by sensitized emission in isotonic solution or after 10, 15 and 20 min of hypotonic stress. Data are normalized for the values of N-FRET of CFP-ICln wt and αIIb-YFP in isotonic solution and expressed in %. ICln wt versus ICln mut: *p < 0.05, **p < 0.01, unpaired Student’s t-test. n refers to the number of cells. (C) Original recordings obtained in hypertonic (top) or hypotonic (bottom) conditions in HEK293 Phoenix cells expressing Canis familiaris ICln mut or wt or in control cells. Current was elicited with voltage increments of 20 mV from −120 to +100 mV applied from a holding potential of 0 mV (lower right inset). Cells overexpressed both ICln and the transfection marker EGFP as separate proteins, or EGFP only as the negative control. Experiments were done in whole-cell configuration. (D) Current density-to-voltage relationship measured 10 min following hypotonic shock, and (E) current density-to-time relationship showing upregulation of IClswell with respect to the control in ICln wt-transfected (p < 0.0001) and not in ICln mut -transfected cells. Data were fitted with second order polynomials, following application of the extra-sum of squares F test.
integrin α-chain remained to be established, and was determined here by NMR spectroscopy to be the (61) ISLHA(65) motif (Fig. 3). FRET experiments in NIH-3T3 cells show that cell-permeable synthetic peptides mimicking the ICln binding site to integrin or the integrin binding site to ICln not only impeded the expected hypotonicity-induced FRET increase between ICln and the plasma membrane probe YFP-mem, but also reduced FRET below the basal levels measured in isotonic conditions (Figs 2B and 4A,B). These data indicate that the cell-permeable synthetic peptides prevented the hypotonicity-induced translocation of ICln towards the cell membrane and also reduced the membrane-resident pool of ICln. To explain our findings, we propose a model (Supplementary Fig. S3) where the cytosolic, integrin-bound and membrane bound forms of ICln exist in a dynamic equilibrium. In isotonic conditions (Supplementary Fig. S3A), the cytosolic form of ICln largely prevails and a significant fraction of ICln is bound to integrin α, as denoted by FRET experiments (Fig. 1B,D). Hypotonicity (Supplementary Fig. S3B) induces the transposition of ICln towards the cell membrane (Fig. 2A), thereby altering the dynamic equilibrium in favor of the membrane-bound ICln, without a substantial increase of the integrin-bound ICln pool (Fig. 1D). The palmitoylated (1020)KVGFFKR(1026) integrin peptide may compete with the integrin α-chain endogenously expressed in these cells for binding to ICln (Supplementary Fig. S3C). Such a competition would have interfered with and weakened or prevented the ICln/α-chain interaction. The palmitoylated (1020)KVGFFKR(1026) peptide may have realized a non-functional interaction with ICln (not sufficiently stable, or inappropriately oriented), which failed in leading to an association of ICln to the plasma membrane. This may have produced an alteration of the dynamic equilibrium between the membrane-bound and unbound ICln in favor of the bound form, thus leading to the observed decrease in N-FRET between ICln and the membrane probe following exposure to the hypertonic medium (Fig. 2B). These findings indicate that binding to integrin is an essential prerequisite for stabilization of ICln at the plasma membrane in hypotonicity. It is also conceivable that ICln may have recruited the membrane-resident ICln in a complex that was not available for interaction with the membrane probe YFP-mem. A similar reduction of FRET between ICln and the membrane probe in hypotonicity could be observed with the palmytoilated ICln peptide (Fig. 4A,B). Importantly, the lack of translocation of ICln provoked by the integrin as well as the ICln peptide was accompanied by an impaired activation of IClswell (Figs 2D,E and 4C). To have an inhibitory effect, the ICln peptide needed to be targeted to the cell membrane (Fig. 5A,B), further suggesting that its molecular target is located in close proximity of the cell periphery. We may envision that only a palmitoylated ICln peptide can reach the correct localization, sufficient local concentration or take the proper orientation to bind α integrin. Therefore, an intracellular, non-palmitoylated water-soluble ICln peptide would be inactive, while a palmitoylated membrane-targeting peptide would bind to and displace ICln from α integrin (Supplementary Fig. S3D), thus preventing the association of ICln with the plasma membrane and hampering the activation of IClswell.

Mutations within the (61)ISLHA(65) ICln motif (Fig. 6A) abrogated the ICln interaction with the integrin αIIb chain (Fig. 6B) and rendered ICln inactive, i.e. unable to increase the IClswell intensity above the endogenous control (Fig. 6C–E). Overall, these experiments indicate that the interaction between ICln and the integrin α-chain promotes the activation of IClswell. It is important to highlight that the direct physical interaction between ICln and integrin αIIb per se does not require α integrin interaction with extracellular ligands, as it was characterized in a cell system overexpressing a truncated form of the α-chain void of the extracellular domain. We therefore suggest that the α-chain provides a scaffolding system for ICln, facilitating the translocation and/or stabilizing the ICln protein in close proximity of the cell membrane, an essential prerequisite for IClswell activation. However, we do not exclude modulation of the ICln/integrin interaction by an outside-in signalling pathway in the physiological setting. We propose a model of integrins orchestrating the regulation of cellular volume by acting as osmosensors as well as osmoeffectors within an integrin–ICln–IClswell pathway (Supplementary Fig. S4). The osmotic signal, sensed by the β-chain, would trigger an intracellular signalling cascade ending on ICln, thus leading to its recruitment at the sub-membrane region and binding to the α-chain. This would promote the association of ICln with the plasma membrane, its accumulation in this compartment and activation of IClswell as the final event. It is also conceivable that ICln may dissociate from integrin prior association with the plasma membrane; this hypothesis is consistent with the mild increase of FRET between ICln and integrin α following hypotonic stress (Fig. 1D), while FRET with the membrane probe is more pronounced (Fig. 2A). The mechanism by which ICln would activate IClswell once recruited at the plasma membrane deserves further investigation. It is conceivable that ICln may take part in the macromolecular complex of the IClswell channel.

The concept of an outside-in signalling mediated by integrin and triggered by hypotonicity is well supported by several findings. In rat liver, the active conformation of integrin β1 rapidly appeared after hypo-osmotic perfusion, indicating integrin activation in response to hepatocyte swelling. Hypo-osmotic hepatocyte swelling triggered the activation of Src-type kinases, mitogen-activated protein (MAP)-kinases and downstream metabolic events, which were fully abolished by integrin αIIb inhibitory proteins exhibiting an Alanine-Glycine-Aspartic Acid (RGD) motif. In rabbit ventricular myocytes, the activation of IClswell induced by mechanical stretch of β3 integrin was associated to an activation of focal adhesion kinases (FAK), Src, the angiotensin II receptor, and the EGF receptor37–40. In basilar artery smooth muscle cells, hypotonic solution increased Src phosphorylation in a time-dependent manner, a phenomenon that was significantly enhanced by integrin β3 overexpression and reduced by integrin β3 knockdown, thus suggesting that integrin β3 mediated the Src phosphorylation induced by hypotonic solution61.

Mapping of the residues displaying frequency changes upon binding of the (1020)KVGFFKR(1026) motif on the 3D structure of ICln reveals a distinct binding site on one edge of the β-sandwich (Fig. 3C). Interestingly, this binding mode is analogous to ubiquitin (Ub) binding of the pleckstrin-homology GLUE domain of the human ESCRT-II EAP45 (also called VPS36)65. Figure 7 shows a structural superposition of the EAP45 GLUE-Ub complex and ICln using the pleckstrin-homology domain as templates. The structural similarity is clearly visible and
convincingly supports a similar protein-binding mode. Interestingly, EAP45 GLUE binding to Ub was found to be coupled to phosphorysitol binding, thus suggesting a structural dynamic link between recognition of ubiquinated cargoes and endosomal phospholipids binding in the context of multivesicular bodies sorting. The similar protein-binding mode might reflect a functional resemblance as ICln displays a dynamic and regulated incorporation in membranes depending on changing osmotic conditions.

We could detect neither endogenous ICln nor recombinant αIIb in immunoprecipitates of recombinant αIIb or endogenous ICln from HEK293 Phoenix cells. This finding is opposed from what was observed in platelets, in which the co-immunoprecipitation was indeed detected, and is indicative of a dynamic interaction, or that only a small fraction of ICln is available for interaction with integrin at a given time. This may represent a characteristic of non-platelet cells and supports the hypothesis that the physiological function of this interaction differs from that found in platelets.

Integrins and the integrin interactome are emerging as novel targets in the control of inflammatory diseases and cancer. As ICln is involved in numerous pathophysiological processes, its pharmacological inhibition is regarded as a potential strategy in the treatment of several life-threatening conditions. For example, in the heart, blockers of Cl⁻ currents such as DIDS may have antiarrhythmic properties. Furthermore, pharmacological inhibition of ICln antagonizes apoptosis by preventing apoptotic volume decrease and downstream apoptotic events. Hence, ICln inhibitors could find application in the treatment of certain forms of heart failure. Although not unanimously, ICln blockers have been shown to impair the motility of cells, delay their proliferation rate and induce cell cycle arrest, thus suggesting that ICln suppression might be beneficial in the treatment of some forms cancer. In brain, ICln contributes to the release of the excitatory neurotransmitters glutamate and aspartate, and is thought to promote the excitotoxic death of neuronal cells in stroke and traumatic brain injury.

In addition, involvement of ICln in the pathogenesis of exercise-induced asthma was suggested. ICln is inhibited by a wide variety of classical non-specific Cl⁻ channel blockers such as 4,4'-Diisothiocyanato-2,2'-stilbenedisulfonic acid (DIDS), 5-nitro-2-(3-phenylpropyl-amino)benzoic acid (NPPB) and tamoxifen. Interestingly, tamoxifen is a selective estrogen receptor modulator that is widely used in the treatment and prevention of breast cancer (https://www.cancer.gov/about-cancer/treatment/drugs/tamoxifen). ICln blockade may in part contribute to its anti-proliferative actions. Accordingly, astrocye-specific LRR8A knockout mice are protected from brain damage after ischemic stroke. Overall, these lines of evidence point to a potential beneficial effect of ICln suppression in several pathological conditions.

In summary, the protein ICln directly binds the highly conserved (1020)KVGFFKR(1026) amino acids of the intracellular domain of the integrin αIIb via the (61)ISLHA(65) amino acid motif lying in the β4 - β5 loop and β5 sheet of its pleckstrin-homology-like N-terminal portion. The stability of this interaction is required for the proper activation of the current ICln during RVD. The findings presented here show that perturbation of the binding of ICln to the integrin α chain leads to ICln attenuation, and suggest that the ICln/α integrin interaction interface could be exploited as a new pharmacological target to obtain specific ICln suppression.

**Figure 7.** 3D structural homology between ICln and the pleckstrin-homology GLUE domain of human ESCRT-II EAP45. Regions of structural similarity are indicated in red (ICln) and orange (GLUE). GLUE binding partner Ubiquitin (Ub) is shown in yellow. Most shifted residues of ICln are shown as cyan spheres. The location of GLUE residue F68 demonstrated to be crucial for the interaction with Ub is also shown in blue. Figures were prepared using the programs TopMatch and PyMOL (The PyMOL Molecular Graphics System, V. r. p., Schrödinger, LLC).
Methods

Cell culture and transient transfection. Human embryonic kidney (HEK)293 Phoenix cells were cultured in Minimum Essential Eagle Medium (Sigma-Aldrich, St. Louis, MO, USA) supplemented with 10% fetal bovine serum (Cambrex Bio Science, Milan, Italy), 2 mM L-glutamine, 100 U/ml penicillin, 100 μg/ml streptomycin and 1 mM pyruvic acid (sodium salt). Mouse NIH-3T3 fibroblasts were cultured in Dulbecco’s modified Eagle’s medium (Sigma-Aldrich) supplemented with 10% newborn bovine serum (Cambrex Bio Science), 100 U/ml penicillin and 100 μg/ml streptomycin. The cells were maintained at 37 °C, 5% CO₂, 95% air and 100% humidity. Subcultures were routinely established every second to third day by seeding the cells into 100 mm diameter Petri dishes following trypsin/EDTA treatment.

For FRET experiments, cells were seeded on 6-well plates and grown overnight to 80% confluence. Cells were then transfected with 8–12 μg of polyethyleneimine (Sigma-Aldrich) and 2–3 μg of DNA (NIH-3T3 fibroblasts) or with 3 μg of plasmid DNA by the calcium phosphate co-precipitation method (HEK293 Phoenix cells). After 8 hours, the medium was replaced. Twenty-four hours post-transfection, cells were seeded on glass coverslips. FRET experiments were done 48 hours after transfection.

For post-clamp experiments, cells were seeded into 30 mm diameter Petri dishes, grown overnight to 50% confluence and transfected with 3 μg of plasmid DNA by the calcium phosphate co-precipitation method (HEK293 Phoenix cells) or with 8–12 μg of polyethyleneimine and 2–3 μg of DNA (NIH-3T3 fibroblasts). Eight hours after transfection, cells were transferred on glass coverslips (10 mm diameter); electrophysiology measurements were performed 24–32 hours after transfection. The clamp efficiency was about 50%. For post-clamp experiments on untransfected cells, HEK293 Phoenix cells were seeded directly on glass coverslips and measurements were performed 16–24 hours after seeding.

Cloning procedures, plasmid constructs and site-directed mutagenesis. Starting from human or Canis familiaris (from MDCK cell line) cDNA, the ORF of full length ICln was amplified by PCR using standard protocols. The cDNA sequence corresponding to amino acids 987–1039 of the human integrin αIIb chain (αIIb short - αIIb) was amplified by PCR from the pcDNA3.1 vector (kindly provided by Prof. N. Moran) containing the full-length CDNA of αIIb. The PCR products encoding ICln and αIIb were cloned in frame into the mammalian expression vectors pECFP-C1 and pEYFP-N1 (Clontech, Mountain View, CA, USA) respectively, to produce the fusion proteins CFP-ICln (CFP was fused to the Nt of ICln) and αIIb-YFP (YFP was fused to the Ct of ICln) suitable for FRET experiments. To label the plasma membrane, the pEYFP-Mem vector (Clontech), encoding for YFP targeted to the plasma membrane by a palmitoylation sequence (YFP-mem) was used. CFP and YFP were used as the FRET donor and acceptor, respectively.

For electrophysiology experiments, ICln or αIIb ORFs were cloned into the Xhol and BamHI restriction sites of the bicistronic mammalian expression vector pIRE2-EGFP (Clontech). The use of vectors bearing an internal ribosome entry site (IRES) allows for the simultaneous expression of two individual proteins (αIIb, or ICln and the enhanced green fluorescent protein (EGFP)) from a single bicistronic mRNA without the production of fusion proteins. Therefore, EGFP expression occurs only if preceded by αIIb, or ICln expression. Single transfected cells could be optically individuated by detecting the fluorescence emitted by EGFP (excitation maximum = 488 nm; emission maximum = 507 nm). Control experiments were conducted in cells transfected with the pIRE2-EGFP vector with no insert and therefore overexpressing only the EGFP protein.

For protein purification, the ORFs of Canis familiaris ICln and its truncation mutant (ICln159X, referred to as TcICln) were cloned in frame with a hexahistidine tag into the bacterial expression vector pET3-His, allowing for the expression of N-terminal His₆-tagged proteins in Escherichia coli strains.

Mutagenesis of the Canis familiaris ICln sequence was carried out using the Site-Directed Mutagenesis kit (Stratagene, La Jolla, CA, USA) with the following primers: 5′GGATATCCCGCGGTTGGCTTGCATAGTTGTCACGGGG3′, forward, and 5′CCCTGGACACGTATGCAAGCCACCGGTGGGGTATTCC3′, reverse. All plasmid inserts were sequenced prior to use in experiments (Microsynth AG, Switzerland).

FRET experiments. Acceptor photobleaching experiments were performed in living or fixed (for 10 min in 3% paraformaldehyde at room temperature) NIH-3T3 or HEK293 Phoenix cells as previously described. Forty-eight hours post-transfection, cells were transferred to a FCS-2 chamber (Biotechns, Inc. Butler, PA, USA) and imaged with a TCS SP2 AOBS confocal microscope (Leica Microsystems, Wetzlar, Germany), using an argon laser and a 63X (1.4 NA) oil immersion objective. During acquisition, cells were bathed in isotonic solution (in mM: NaCl 90, KCl 5, CaCl₂ 2, MgCl₂ 2, Glucose 5, HEPES 10, Mannitol 80, pH 7.4) or in phosphate buffered solution. CFP was excited at 458 nm and the fluorescence acquired between 465 and 505 nm; YFP was excited at 514 nm and the fluorescence acquired between 525 and 600 nm. YFP photobleaching in whole cells was performed with 4 sequential illuminations (4 frames, 1024 × 1024 resolution, 200 Hz) at 514 nm (setting the AOTF at 100% and the zoom at 8×). FRET efficiency was calculated from the ratio of the CFP fluorescence evaluated before (CFPₚₑ) and after (CFPₚₒ) YFP photobleaching, using the formula:

\[
\text{FRET efficiency} = 1 - \frac{\text{CFP}_{\text{post}}}{\text{CFP}_{\text{pre}}} \]

in three (NIH-3T3 cells) or 4–5 (HEK293 Phoenix cells) different plasma membrane regions of interest (ROIs) for each cell. ROIs were drawn within the bleaching ROI, on the pre-bleach YFP image, in areas with a clear targeting of αIIb-YFP at the cell periphery. n in Fig. 1B represents the number of analyzed cells.

For sensitized emission experiments, living NIH-3T3 cells were kept in isotonic solution for 5 minutes. The donor (CFP), FRET and acceptor (YFP) images were acquired, and the isotonic solution was replaced with hypotonic solution (in mM: NaCl 90, KCl 5, CaCl₂ 2, MgCl₂ 2, Glucose 5, HEPES 10, pH 7.4). Donor, FRET and
acceptor images were then acquired after 10, 15 and 20 minutes. The sensitized emission (F_{sen}) was calculated as already described\(^{79,80}\) and according to the formula:

\[
F_{sen} = \frac{M_M - M_D \cdot \beta - M_{DA} \cdot (\gamma - \alpha \cdot \beta)}{(1 - \beta \cdot \delta)}
\] (2)

M indicates the measured fluorescence (subtracted of the background) in the ROIs. Donor emission with donor excitation is indicated as \(M_D\), acceptor emission with donor excitation (FRET channel) is indicated as \(M_M\), and acceptor emission with acceptor excitation is indicated as \(M_{DA}\). The parameters \(\alpha, \gamma, \delta\) and \(\beta\) have been calculated in samples expressing only CFP or only YFP according to\(^{80}\). The N-FRET indices were calculated in each cell from the mean intensities derived from 3 plasma membrane ROIs according to the formula:

\[
N - FRET = \frac{F_{sen}}{M_D}
\] (3)

where \(F_D\) is defined as \(M_D + F_{sen}\). In Figs 1D and 6B, \(n\) indicates the number of analyzed cells.

Visualization of CFP- and/or YFP-expressing cells and detection of FRET in the experiments shown in the Figs 2A–C, 4A,B was performed on an IX70 inverted microscope (Olympus, Tokio, Japan) equipped with a Polychrome 4 monochromator (TILL Photonics, Planegg, Deutschland) and a cooled charge-coupled device camera (TILL Imago SVGA) controlled by the TILL Vision software (versions 3.3 and 4.0). Experiments were performed by changing three separate Olympus BX cubes equipped with the appropriate filter combinations for CFP (in nm: excitation filter 436/20, beamsplitter 455 DCLP, emission filter 480/40), YFP (in nm: excitation filter 510/20, beamsplitter 530 DCLP, emission filter 560/40), and FRET measurements (in nm: excitation filter 436/20, beamsplitter 455 DCLP, emission filter 560/40) (AHF Analysentechnik, Tübingen, Germany). FRET signals were determined in areas of interest of the cell in the membrane region before, 5 and 15 minutes after hypotonic stimulation. \(n\) indicates the number of cells analyzed; for each cell, ten ROIs were taken into account\(^{28–30}\).

To obtain the N-FRET image shown in Fig. 1C, the following image processing steps were carried out. ROIs were designed corresponding with cells expressing only CFP or YFP. From these ROIs, correction factors were measured and calculated. With these factors, sensitized emission images were obtained applying the formula (2) in a pixel-by-pixel manner using the math or the Image calculator tools of ImageJ software. The sensitized emission images were ratioed to the \(F_D\) image to obtain the N-FRET pictures. To suppress excessive noise in dim parts of the images, the N-FRET images were smoothed with a spatial filter and a mask was created, by setting a threshold equal to the background from this image. Subsequently, this mask was applied to the original, unfiltered N-FRET image\(^{80}\). The N-FRET images obtained were then optionally smoothed.

**Patch-clamp recordings.** Patch-clamp experiments were performed as previously described\(^{29,30}\). Shortly, single cells expressing EGFP were selected by fluorescence microscopy and voltage-clamped using the whole-cell patch-clamp technique. The resistance of the glass pipettes was 3–8 MΩ when filled with the pipette solution (in mM: CsCl 125, MgCl\(_2\) 5, EGTA 11, raffinose 50, ATP 2, HEPES 10, pH 7.2 (adjusted with CsOH)). The hypertonic bath solution was composed of (in mM): NaCl 125, CaCl\(_2\) 2.5, MgCl\(_2\) 2.5, HEPES 10, mannitol 100, pH 7.4 (adjusted with NaOH). Fast exchange of the hypertonic bath solution with a hypotonic bath solution (in mM: NaCl 125, CaCl\(_2\) 2.5, MgCl\(_2\) 2.5, HEPES 10, pH 7.4) was obtained using a perfusion system with a flow rate of 5 ml/min and a bath volume of ~300 μl. For data acquisition, an EPC-10 amplifier (HEKA Elektronik, Lambrecht, Germany) controlled by a Macintosh computer running Patch Master (HEKA Elektronik) software was used. Access resistance was measured and calculated. Current values were normalized to the membrane capacitance to obtain the current density. Each data set was obtained from cells from at least 3 independent subcultures with control experiments on the same subculture.

**Protein expression and purification.** cICln and TcICln were overexpressed and labelled with \(^{15}\)N as N-terminal His\(_{6}\)-tagged fusion proteins in E. coli BL21(DE3) strain (Agilent Technologies, Santa Clara, CA, USA). For protein expression and uniform labeling with \(^{15}\)N, transformed E. coli BL21 (DE3) (Agilent Technologies) were first grown in LB-medium (Sigma-Aldrich) and then diluted 1:100 into 1 liter of minimal medium (48 mM Na\(_2\)HPO\(_4\), 22 mM KH\(_2\)PO\(_4\), 8.56 mM NaCl and 18.7 mM 15NH\(_4\)Cl (Euriso-top, Germany), supplemented with 20 ml of a 20% (w/v) glucose solution (Sigma-Aldrich), 2 ml of 1 M MgSO\(_4\), 0.3 ml of 1 M CaCl\(_2\) and 100 μg/ml ampicillin) and further cultured at 37 °C in an orbital shaker. Protein expression was induced by addition of isopropyl-1-thio-β-D-galactopyranoside to a final concentration of 0.5 mM. Bacterial cells were harvested 4 h after induction. The bacterial pellet obtained from a 1 liter 15N labelled M9 minimal medium culture was resuspended in 30 ml of lysis buffer (30 mM K\(_2\)HPO\(_4\), pH 8.0) and lysed using a French-press. To purify over-expressed proteins, the bacterial lysate was cleared by centrifugation and loaded onto an IMAC column (HisTrap, GE Healthcare, Chicago, IL, USA) pre-equilibrated with lysis buffer. Following extensive washing with lysis buffer, bound protein was eluted with a buffer containing 50 mM K\(_2\)HPO\(_4\), pH 8.0 by using a NaCl gradient. Fractions containing cICln or TcICln were pooled, concentrated in Amicon Ultra 10 KDa ultrafiltration units (Millipore, Burlington, MA, USA) and loaded onto a Hilo load 16/60 Superdex 75 column (GE Healthcare) equilibrated in 50 mM K\(_2\)HPO\(_4\), 100 mM NaCl, pH 7.2. Fractions containing cICln or TcICln were collected, pooled and concentrated. For the following NMR experiments the buffer was changed to 50 mM K\(_2\)HPO\(_4\), pH 6.6.
NMR spectroscopy. NMR experiments were performed at 25 °C using Inova 800 MHz spectrometers (Varian Inc., Palo Alto, CA, USA). 1H,15N HSQC spectra were recorded with carrier frequencies for 1H and 15N at 4.73 ppm and 120.0 ppm, respectively. The sample contained 0.8 mM recombinant canine ICln, 50 mM K2HPO4, pH 6.6 in 90% H2O and 10% D2O. All spectra were processed using NMRPipe/NMRDraw and analyzed with Sparky.

Salts, chemicals and drugs. All salts and chemicals used were of “pro analysis” grade. Palmitoylated and non-palmitoylated synthetic peptides were synthesized and purified by HPLC by the Organic Synthesis Core (Royal College of Surgeons, Dublin 2, Ireland) or purchased from GenScript (Piscataway, NJ, USA).

Statistical analysis and sequence alignment. All data are expressed as arithmetic means ± S.E.M. For statistical analyses, GraphPad Prism software (version 4.03 for Windows, GraphPad Software, San Diego, CA, USA) was used. Significant differences between means were tested by paired or unpaired, two-tailed, Student’s t-test as appropriate. The current density-to-time and current density-to-voltage relationships were fitted with linear regression or second order polynomials (Y = A + BX + CX2). For detecting significant differences between those data, the extra-sum of squares F test was applied. Statistically significant differences were assumed at p < 0.05 (**p < 0.05; ***p < 0.01; ****p < 0.001). n indicates the number of cells tested, except where otherwise specified. Sequence alignment was done with Gene Runner software, version 6.5.48.

Data Availability. The datasets used and analysed during the current study are available from the corresponding author on reasonable request.

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**Acknowledgements**

The authors sincerely thank Prof. Niam Moran and Prof. Johannes Fürst for helpful comments and suggestions and acknowledge the expert secretarial assistance of Elisabeth Mooslechner. D.A.C. was supported by the Harvard/Salzburg-Stipendium 02/00/28060/2013/064 (Stadt Salzburg).

**Author Contributions**

R.K., M.P. and S.D. conceived the study. A.S., G.T., S.R., D.A.C., M.T., K.L., C.N. and S.D. performed and analyzed the experiments. W.P., R.K. and M.P. acquired funding. S.D. wrote the original draft, C.N. and W.P. reviewed and edited and all authors read and approved the final manuscript.

**Additional Information**

Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-48496-4.

**Competing Interests:** The authors declare no competing interests.

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