G-protein coupling and nuclear translocation of the human abscisic acid receptor LANCL2

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Abscisic acid (ABA), a long known phytohormone, has been recently demonstrated to be present also in humans, where it targets cells of the innate immune response, mesenchymal and hemopoietic stem cells and cells involved in the regulation of systemic glucose homeostasis. LANCL2, a peripheral membrane protein, is the mammalian ABA receptor. We show that N-terminal glycine myristoylation causes LANCL2 localization to the plasmamembrane and to cytoplasmic membrane vesicles, where it interacts with the α subunit of a G protein and starts the ABA signaling pathway via activation of adenylate cyclase. Demyristoylation of LANCL2 by chemical or genetic means triggers its nuclear translocation. Nuclear enrichment of native LANCL2 is also induced by ABA treatment. Therefore human LANCL2 is a non-transmembrane G protein-coupled receptor susceptible to hormone-induced nuclear translocation.

The human genome encodes three distinct LANCL proteins, LANCL1, LANCL2 and LANCL31, which share a high structural homology with the lanthionine synthetase component C, a cyclase involved in the synthesis of lantibiotics in Prokaryotes2. LANCL3 has been suggested to be a pseudogene3. LANCL1 has been hypothesized to be implicated in the metabolism of lanthionine metabolites in the central nervous system4. LANCL2 proved to be the human receptor of abscisic acid (ABA)5–10. ABA, a long-known plant hormone11,12, has been shown to be present also in mammals, where it affects several key functions in different cell types9,10,13,14. ABA behaves as a pro-inflammatory modulator of cells of the innate immune response7,15–17, stimulates the proliferation of human mesenchymal and hemopoietic stem cells18,19, and is involved in the control of systemic glucose homeostasis5,20–24.

LANCL2-mediated ABA signaling in mammals requires a pertussis toxin (PTX)-sensitive G protein5, eventually leading to an increase of intracellular Ca2+ levels ([Ca2+]i). The signaling pathway downstream of ABA binding to LANCL2 involves the activation of adenylate cyclase (AC), followed by overproduction of cAMP, PKA-catalyzed phosphorylation and stimulation of the plasmamembrane-bound ADP-ribosyl cyclase CD38, which converts NAD+ to cADPR and ADPR, leading to an increase of both extracellular Ca2+ entry and Calcium-induced calcium release (CICR)-mediated intracellular Ca2+ mobilization5,9,15,21.

Several indirect lines of evidence point to a G i as the G protein coupled to LANCL2: i) the sensitivity of the ABA signaling pathway to PTX in human granulocytes and in insulin-releasing cells15,21; ii) the accumulation of inositol 1,4,5-P3 (IP3) in human cells co-transfected with LANCL2 and a chimeric G protein, Goqγi5, upon stimulation with ABA7, and, iii) inhibition of the ABA-induced cAMP increase in ABA-sensitive human cells by overexpression of transducin, a βγ-subunit scavenger7. However, conclusive identification of the nature of the G protein coupled to LANCL2 has yet to be provided. For instance, the role of G αi in AC signaling is exceedingly complex, as witnessed by both AC-activating and inhibiting effects related to wide heterogeneity of the coupling receptors and of the various membrane-associated AC isoforms5,20. Moreover, the mechanism of the LANCL2-G protein coupling, specifically whether it is direct or mediated by other proteins, remains to be defined.

Interestingly, LANCL2 is not a transmembrane protein, as predicted in silico from its sequence27–29 and confirmed in vitro by the observation that it can be removed from the plasmamembrane without the use of detergents, either by mild chemical treatments30 or by inhibition of its post-translational N-terminal myristoylation28.

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In addition, the non-myristoylated LANCL2-GFP fusion protein has been found to be confined to the nucleus\(^n\)28. This observation raises the possibility that its hormone ligand ABA may affect the trafficking of LANCL2 between membranes and nucleus.

Indeed, recent findings allow to reconcile the non-transmembrane localization of LANCL2 with its hormone receptor function, as the human anion exchanger AE1 has been shown to mediate ABA transport across the plasmamembrane\(^n\)30.

Here, we investigated the unusual features of LANCL2 among G protein-coupled animal hormone receptors (GPCR), by means of site-directed mutagenesis and of confocal fluorescence microscopy, fluorescence recovery after photobleaching (FRAP) and photoactivation techniques. The localization, the intracellular mobility of LANCL2 in the presence of ABA and its interaction with G\(_i\) were explored.

**Results**

**Role of N-terminal myristoylation in the subcellular localization of untagged LANCL2.** Comparison between the three LANCL genes shows that Met 19 of LANCL2 is aligned with the start methionine of the highly homologous LANCL1 and LANCL3 (Supplementary Table 1), which are cytosolic proteins27–29. Thus, transcription of LANCL2 starting from Met 19 would result in a shorter protein, lacking the myristoylated glycine and the 18-aminoacids N-terminal stretch (LANCL2sh).

Supplementary Fig. 1 shows a representative Western blot of full length LANCL2 from HeLa, Hek and the human adenocarcinoma cell line MDA-MB-468 (MDA-468), revealed with a specific mAb (see Materials and Methods). In this cell line, a 0.9 Mb region, containing the LANCL2 gene, is amplified\(^n\)31. Overexpression of LANCL2 in MDA-468 cells was documented by microarray profile analysis, by RT-PCR and by measuring \(^{3}H\) ABA binding to intact cells, in comparison with the MDA-MB-231 (a breast adenocarcinoma cell line not over-expressing LANCL2)32. The \(B_{\text{max}}\) values calculated from the relevant saturation curves by Scatchard plot analysis were 0.70 pmol ABA/mg protein for MDA-468 and 0.023 pmol ABA/mg protein for MDA-231, respectively.

Landlinger et al.\(^n\)23 demonstrated that GFP-fused and nonacids N-terminal stretch (LANCL2sh).

Results were performed 48 h after transfection. ROIs were selected in a cytoplasmic area of the cells close to the plasmamembrane avoiding the Golgi and nuclear areas (Fig. 2A).

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- **Here, we investigated the unusual features of LANCL2 among G protein-coupled animal hormone receptors (GPCR), by means of site-directed mutagenesis and of confocal fluorescence microscopy, fluorescence recovery after photobleaching (FRAP) and photoactivation techniques.**
- **The localization, the intracellular mobility of LANCL2 in the presence of ABA and its interaction with G\(_i\) were explored.**

**Effects of myristoylation on the intracellular mobility of LANCL2.** In order to investigate LANCL2 mobility in live cells, we exploited two different techniques that require the use of fluorescently tagged proteins: fluorescence recovery after photobleaching (FRAP) and fluorescence decay after photobleaching (FDAP)

FRAP explores the mobility of the tagged protein by analyzing the recovery of the fluorescent signal in a specific region of interest (ROI) after permanent photobleaching\(^n\)36. To this purpose, HeLa cells were transiently transfected with the full length LANCL2-GFP or with the truncated LANCL2sh-GFP form, and FRAP experiments were performed 48 h after transfection. ROIs were selected in a cytoplasmic area of the cells close to the plasmamembrane avoiding the Golgi and nuclear areas (Fig. 2A).
The average $t_{1/2}$ calculated for LANCL2-GFP was significantly higher than the $t_{1/2}$ of LANCL2sh-GFP (3.9 ± 0.9 s and 1.6 ± 0.2 s for LANCL2-GFP and LANCL2sh-GFP, respectively, graph in Fig. 2A; $n = 20$ cells from three different transfection experiments, $p < 0.001$).

Then, we also performed photoactivation experiments on LANCL2 fused to photoactivatable GFP (PAGFP). To compare the mobility of wild-type LANCL2 with the non-myristoylatable G2A-mutagenized LANCL2, we transiently expressed LANCL2-PAGFP or LANCL2-G2A-PAGFP in HeLa cells, and photoactivated a circular cytosolic ROI chosen next to the plasmamembrane, in cells of similar shape. Images acquired at the same time points after photoactivation clearly indicated that the fluorescence decay of LANCL2-PAGFP was much slower than that of LANCL2-G2A-PAGFP (Fig. 2B), as confirmed by calculation of the half time of recovery ($t_{1/2}$), after fitting values to one phase decay curves: $t_{1/2}$ values of LANCL2-PAGFP and of LANCL2-G2A-PAGFP were 50 ± 30 s and 3.4 ± 0.75 s, respectively (graph in Fig. 2B; $n = 34$ cells from four different transfection experiments, $p < 0.001$).

Moreover, as shown in Supplementary Fig. 2, LANCL2 is capable of rapid lateral diffusion within the plasmamembrane. Together, these observations indicate that the full length LANCL2 is membrane-associated also in the cytoplasm. When protein myristoylation was prevented by single point mutation, association with the cell membranes was lost, as indicated by its faster free diffusion inside the cell.
LANCL2 dynamics after ABA treatment. The fact that untagged LANCL2 localized to the cell nucleus when not myristoylated (Fig. 1G–I) suggested to explore whether nuclear translocation of the protein might occur as part of its signaling pathway, following ABA binding. To investigate whether LANCL2 migrated into the nucleus following addition of ABA, we used the full-length form of LANCL2 fused to PAGFP; the finding that LANCL2-GFP and untagged LANCL2 behaved similarly in their nuclear localization upon G2A mutation (Fig. 1H,I, respectively), was taken as a demonstration that the GFP tag did not significantly modify the intrinsic ability of LANCL2 to migrate into the nucleus.

At 48 h after transfection with LANCL2-PAGFP, HeLa cells of similar shape and size were subjected to photobleaching in a 8 μm-diameter cytoplasmic ROI, with or without (control) the simultaneous addition of 5 μM ABA; the photoactivation ROI was chosen close to the plasmamembrane, 10–20 μm far from the nearest nuclear membrane, whereas the nuclear ROI covered most of the nuclear area (Fig. 3A). The fluorescent signal from LANCL2-PAGFP rapidly distributed away from the photoactivated region, slowly increasing also inside the nucleus; the dynamics were recorded for 25 min, when a steady-state level of nuclear fluorescence was reached (Fig. 3B, left panel).

At 25 min after photoactivation, the mean nuclear fluorescence, relative to the pre-photoactivation frames, was significantly higher in the ABA-treated cells compared with untreated controls (4.18 ± 1.4 vs. 2.51 ± 0.5, respectively, n = 22 cells from three different transfection experiments, p = 0.005).

Compared with the nuclear enrichment of LANCL2-PAGFP in the absence of ABA, that of PAGFP alone, compared with the nuclear enrichment of LANCL2-PAGFP in the absence of ABA, that of PAGFP alone, used as a control, showed a quantitatively similar increase (the plateau of nuclear fluorescence relative to pre-stimulation values was 2.2 ± 0.3), but significantly different kinetics (1/2 of 2.2 ± 0.4 and 6.6 ± 1.3 min for

Figure 2. Intracellular mobility of different recombinant forms of LANCL2. (A) A representative FRAP experiment. EGFP signal in HeLa cells overexpressing LANCL2-GFP (left column) or LANCL2sh-GFP (right), before (prePB) and at indicated time points after photobleaching (T0). Graph: kinetics of mean FRAP, with s.d., of LANCL2-GFP (blue) and of LANCL2sh-GFP (red trace); the mean half-life constants are indicated in the abscissa as dotted lines. (B) A representative FDAP experiment. PAGFP signal in HeLa cells overexpressing LANCL2-PAGFP (left column) or LANCL2-G2A-PAGFP (right), before (prePA) and at different time points after photoactivation (T0). Graph: kinetics of mean FDAP, with s.d., of LANCL2-PAGFP (blue) and of LANCL2-G2A-PAGFP (red trace); the mean half-life constants are indicated in the abscissa as dotted lines.
PAGFP and LANCL2-PAGFP, respectively, \( p < 0.001 \); Supplementary Fig. 3). Thus, the slight increase of nuclear fluorescence of LANCL2-PAGFP in the unstimulated cells could be due to a low extent of ABA-independent, spontaneous, LANCL2 nuclear migration, rather than to non-specific nuclear import of a PAGFP-tagged protein.  

Analysis of the decaying fluorescence in the cytoplasmic photoactivated ROI revealed a greater LANCL2 decrease in the ABA-treated than in the untreated cells: the plateau of the fluorescence values relative to the first photoactivated frame and fitted to a single exponential curve, was 0.32 ± 0.09 for control, and 0.22 ± 0.076 for ABA-treated cells (\( n = 17 \) cells, \( p < 0.001 \); Fig. 3B, right panel). Moreover, the fluorescence decay in the photo-activated cytoplasmic ROI was faster in ABA-treated compared with control cells (the half-life was 50 ± 30 s for control, and 23 ± 13 s for ABA-treated cells; \( n = 17 \) cells, \( p = 0.02 \)).

The higher nuclear content of LANCL2 in the ABA-treated cells, in agreement with the greater and faster LANCL2 decrease in the cytoplasmic photoactivation region, indicates that ABA stimulates LANCL2 migration into the nucleus.

**Role of localization on LANCL2 function.** Next, we investigated the role of the myristoylation-dependent membrane localization of LANCL2 in ABA perception/signaling.

ABA binding to LANCL2 eventually results in the stimulation of AC mediated by the βγ subunit complex released from an activated Gi-protein.  

To determine whether LANCL2 directly interacted with the α subunit of a Gi-protein, we performed FRET experiments, by labelling LANCL2 with a donor fluorophore (EGFP) at the C-terminus (LANCL2-GFP), and the α subunit of a chimeric Gi protein, known to be activated by LANCL2, with an acceptor fluorophore (TagRFP, Gi-RFP). As a negative, non membrane-associated control, we used the EGFP-tagged soluble short form of LANCL2, LANCL2sh-GFP (Supplementary Table 1).

Co-transfection of LANCL2-GFP and Gi-RFP in HeLa cells generated a measurable FRET signal, indicating that the donor (EGFP) and the acceptor (TagRFP) were sufficiently close to allow resonance energy transfer (Fig. 4A–C). Interestingly, FRET between LANCL2 and the Gi α subunit was observed not only on plasmamembranes (identified by their intense signal at the border of the cells, and analyzed by manually drawn ROIs), but also in intracellular regions: the average apparent FRET efficiency calculated for the LANCL2-Gi couple in plasmamembrane ROIs (20 ± 7%) was similar to the FRET calculated in whole-cell regions (20 ± 5%; \( n = 40 \) cells in five different transfection experiments, Fig. 4G). Conversely, the interaction was lost (FRET efficiency 1 ± 2%) when the Gi-RFP was co-transfected with soluble LANCL2sh-GFP (Fig. 4G).

According to these results, we can infer that the interaction with the α subunit of the chimeric Gi-protein requires a membrane-associated, myristoylated, LANCL2, the close proximity between LANCL2 and Gi occurring both on the plasmamembrane and on intracellular membranes.
We then compared the ABA-binding ability of purified recombinant LANCL2 (LANCL2-gst) and of its truncated short form (LANCL2sh-gst). Both forms were cleaved from the GST tag (Supplementary Table 1), produced as described in6, but cleaved in the absence of DTT, a more physiological condition leading to a higher binding affinity than previously observed. Indeed, specific and saturable [3H]ABA binding occurred to both proteins with similar affinities (Kd values were 8.9 ± 1 μM and 10.6 ± 1.4 μM for LANCL2 and LANCL2sh, respectively, from three different experiments).

Thus, the 18 aminoacids N-terminal stretch of LANCL2, which includes the myristoylation site, is not necessary for ABA binding.

Finally, we explored the ability of different LANCL2 forms to trigger the ABA-signaling pathway, by measuring the increase of the intracellular concentration of cAMP ([cAMP]i) in transfected HeLa cells following addition of ABA (as described in15).

HeLa cells transfected either with untagged LANCL2, or with LANCL2-GFP, or with the untagged mutagenized G2A LANCL2 (LANCL2-G2A) were incubated with 5 μM ABA and the [cAMP]i was measured after 30 s. The [cAMP]i increased in HeLa cells overexpressing untagged LANCL2 (165 ± 19% over basal, unstimulated values, p < 0.02) or LANCL2-GFP (170 ± 21%, p < 0.02). Conversely, the [cAMP]i was not significantly modified compared to basal values in cells transfected with the non-myristoylatable LANCL2-G2A (94 ± 17%, p = 0.6), Fig. 4H.

Therefore, LANCL2 myristoylation and a functional interaction between LANCL2 and the chimeric Gi subunit (Fig. 4H) are essential for the ABA-triggered activation of adenylate cyclase, both requirements being met at the plasmamembrane and at internal membranes in the cytoplasm.

Conversely, binding of ABA does not require association of LANCL2 with the plasmamembrane, as it is not significantly affected by the truncation of 18 N-terminal aminoacids. This observation suggests that an unmyristoylated, and thus freely-diffusible, LANCL2 could be able to bind ABA and activate other, cAMP-independent, signaling pathways.

Discussion
Here, we demonstrated that N-terminal glycine myristoylation causes LANCL2 localization to the plasmamembrane and to cytoplasmic membrane vesicles as well. Interestingly, evidence has recently been provided of
non-canonical subcellular sites for G protein activation: these include endosomal localization of functionally active GPCR\(^{39-41}\), and the interaction of the GIV/Girdin protein, a non-receptor guanine nucleotide exchange factor, with heterotrimeric G proteins at the Golgi\(^{42,43}\) and at still undefined intracellular sites\(^{44}\). The effectiveness of N-myristoylated LANCL2 in coupling to G\(_i\) physically and functionally (FRET efficiency and [cAMP], increase, Fig. 4) also when associated to internal membrane vesicles represents a further example of G-protein mediated signal initiation at non-canonical intracellular sites.

The finding of nuclear localization of LANCL2 after exposure of the cells to ABA (Fig. 3) suggests a possible role for LANCL2 in ABA-triggered transcriptional events. Indeed, stimulation by ABA of the transcription of several cytokines regulating the proliferation of human mesenchymal and hemopoietic stem cells has been previously observed\(^{18,19}\), and ABA-induced transcriptome variations are well established in plants\(^{45}\).

A still unanswered question is whether demyristoylation of native LANCL2 (e.g., via specific enzymes or limited proteolysis targeting the N-terminus) is required for nuclear import and whether ABA binding itself can boost demyristoylation. So far, we were unable to identify any demyristoylated form of LANCL2, which could however escape detection for several reasons (e.g., transient nature due to rapid nuclear clearance, or low specificity of anti-myristoyl anchor antibody).

The facts that LANCL2 is intracellular and that its hormone ligand ABA is an anion at physiological pH values, and thus unable to diffuse across the membrane lipid bilayer, imply that a transmembrane transport is required to allow extracellular ABA to reach its receptor. Indeed, influx of ABA has been recently demonstrated to occur through the transmembrane bicarbonate/chloride exchanger 1 (AE1, band 3 protein) in erythrocytes\(^{30}\). Moreover, the bidirectional nature of ABA transport across the plasmamembrane may also support paracrine mechanisms, already observed with this hormone\(^{10,17-19,21}\), based upon release of ABA from one cell and its perception/signaling in a neighboring cell. In any case, the fact that ABA enters the cells through a transmembrane transporter is unique among animal hydrophilic hormones, which bind to the extracellular domain of integral membrane receptors.

In conclusion, this study highlights some non-canonical features of the mammalian ABA receptor: i) LANCL2 is a membrane-bound, but not an integral membrane protein, whose myristoylation allows coupling to a G\(_i\) protein and activation of adenylate cyclase; ii) upon ABA-binding LANCL2 can translocate into the nucleus, to our knowledge an unprecedented feature for a membrane-associated, G-protein coupled, receptor.

The diversified LANCL2 intracellular behavior might account for the emerging multifunctionality of LANCL2\(^{46}\), for the heterogeneity of cell targets and for the pleiotropy of ABA-stimulated regulatory effects in these cells, impacting on fundamental systemic functions\(^{7-10,15-19,47}\).

These results broaden our knowledge on the mechanisms underlying hormonal receptor-G protein interaction in animals, by providing the first observation of a non-transmembrane G protein-coupled receptor capable of hormone-induced nuclear translocation. Thus, LANCL2 appears to be uniquely endowed with two key features (G protein-coupling and nuclear translocation), each one typical of a different family of animal hormone receptors, i.e. peptide and steroid hormone receptors, respectively.

Conservation of ABA as a stress hormone regulating cell responses to environmental stimuli in plants, lower Metazoa (sponges and hydroids)\(^{48,49}\) and mammals, suggests its origin in a common precursor to animals and plants. This ancient evolutionary root may indeed place this hormone in a category of its own compared to the other animal hormones.

**Materials and Methods**

**Reagents and antibodies.** The mouse monoclonal antibody against human LANCL2 (LANCL2 mAb) was produced by Dr. C. Fresia at Molecular Biotechnology Center (MBC), Turin (Italy), as previously described\(^ {30}\). Primary antibody anti-human vinculin was a kind gift of E. Turco (MBC, Turin). HRP-conjugated secondary antibodies were from Santa Cruz Biotechnology (Santa Cruz, CA); Alexa-488-conjugated ones were from Thermo Fisher Scientific (NY, USA). 2-hydroxymyristic acid (HMA), tissue culture media and supplements were purchased from Sigma-Aldrich (Milan, Italy). HMA, dissolved in chloroform and dryed out with N\(_2\), was resuspended at a 10 mM concentration in 1% fatty acid free BSA in HBSS, through overnight shaking\(^ {33,34}\).

**Cell cultures and transfection.** HeLa, HEK-293, MDA-MB-468, MDA-MB-239 cells were cultured in Dulbecco’s modified Eagle’s complete cell culture medium (DMEM with 10% fetal bovine serum, 50 U/ml penicillin, 50 μg/ml streptomycin). Cells were seeded at 15,000/cm\(^2\), on 25 mm glass coverslip Menzel-Glaser thickness N1.5 (Tecnovetro SRL, Monza, Italy), on 8-wells chambered coverglasses (Lab-Tek, Nunc, Thermo Fisher Scientific, NY, USA), or on 35 mm Petri dishes 24 hours before transfection, and maintained in a humidified 5% CO\(_2\) atmosphere at 37°C. Cells were transfected with the calcium-phosphate method\(^ {50}\), adding to the fresh complete medium a 10% volume of transfection reagent composed of Hepes Buffered Saline, 100 mM CaCl\(_2\) and 30 μg/ml endotoxin-free DNA (prepared with EndoFree Plasmid Midi Kit, QIAGEN, following manufacturer instructions). Medium was replaced after 12–16 h.

The ready-to-use construct CellLight\(^ \text{®} \) Plasma Membrane-RFP (Molecular Probes, Thermo Fisher Scientific, Waltham, MA USA) was transfected into cells 24 h after LANCL2-GFP transfection, using BacMam 2.0 technology; it was obtained by expressing the red fluorescent protein (RFP) fused to the myristoylation/palmitoylation sequence from Lck tyrosine kinase, providing accurate and specific targeting to cellular plasmamembrane.

**Western Blot.** Western blot analyses were performed following standard methods, as previously described\(^ {38}\): 5–50 μg of different cell lysates (1% SDS, 5 mM EDTA, 10 mM DTT, 1:200 of protease inhibitors cocktail, Sigma-Aldrich) were loaded on a 10% polyacrylamide gel.
Recombinant proteins and [3H]ABA binding. Expression and purification of the recombinant proteins were performed as previously described. BL21 (DE) E. coli cells utilized were grown in Luria–Bertani medium (Difco, BD Italy, Roma, Italy).

The release of LANCL2-gst or of LANCL2sh-gst (Supplementary Table 1) was achieved with the PreScission Protease (GE Healthcare), by incubating the GSH-Sepharose-bound fusion protein for 16 h at 4°C following manufacturer instruction, but without DTT: the protease cleavage left 9 amino acids at the N-terminus of the protein, considered in the Mw calculations.

To compare specific [3H]ABA binding of recombinant LANCL2-gst and LANCL2sh-gst produced in E. coli, we proceeded as previously described.

[3H]ABA binding experiments to intact MDA-MB-468 and MDA-MB-231 cells, were performed as previously described.

Immunofluorescence and fluorophores. HeLa or HEK-293 cells were grown on 18- or 25-mm diameter coverslips, previously coated with poly-L-lysine (Sigma-Aldrich), for 24 hours before transfection, 2-HMA treatment or fixation. Specimens were treated for fixation at RT for 10 min with para-formaldehyde (4% in PBS), then permeabilized for 8 min with 0.1% Triton X100, blocked for 10 min with 3% BSA, incubated for 1 hour with anti-LANCL2 mAb 5 μg/ml in BSA 3%, and finally stained for 30 min with 10 μg/ml Alexa Fluor 488 secondary antibody (Thermo Scientific). Mounting medium was Mowiol + 2.5% DABCO (Sigma-Aldrich).

For live analysis cells were maintained in complete DMEM without phenol red, supplemented with 20 mM Hepes.

Excitation/emission maxima of fluorophores utilized were: Alexa 488 495/519, EGFP 488/507, PAGFP 504/517, TagRFP and CellLight®-RFP 555/584.

Localization images were obtained using a Leica TCS SP2 confocal microscope (Leica Microsystems, Heidelberg, Germany) equipped with argon/He-Ne laser sources and Leica HCX PL APO CS 63.0 X NA1.40 oil objective.

To exclude that the EGFP-fusion could affect subcellular distribution, we treated HeLa cells overexpressing untagged LANCL2 with the N-myristoyl transferase inhibitor 2-hydroxymyristic acid (HMA), then stained the protein with a specific anti-LANCL2 mAb; as a further control to ensure that no intracellular morphological changes affecting LANCL2 localization could have been induced by the HMA treatment, we used the mutated form of LANCL2 bearing the G2A mutation (therefore not susceptible to undergo myristoylation), fused to EGFP.

cDNA constructs. cDNA encoding human LANCL2 was subcloned from previously available constructs into the following expression vector: pEGFP-N1, pCDNA3.1+, pPAGFP-N1, pGEX-6P-1. When compatible, restriction enzyme sites were used for subcloning; otherwise, and for truncated LANCL2 forms (LANCL2sh-gst, LANCL2sh-GFP), specific primers containing the desired restriction enzyme sites were designed for PCR amplification from plasmid (TibMolBiol, Genova, Italy). The G2A site-specific mutagenesis was obtained with the QuickChange Lightning Site-Directed Mutagenesis kit (Agilent Technologies; Santa Clara, CA), following manufacturer’s instructions, applied on the corresponding/analogous/appropriate vector.

EGFP and PAGFP tags were fused at the C-terminus of LANCL2 protein forms.

The EGFP coding sequence contained an A207K mutation to eliminate potential dimerization of the EGFP moiety.

The α subunit of the chimeric G protein was subcloned into pTagRFP-C (Evrogen, Moscow, Russia), thus resulting N-terminally fused to TagRFP monomeric protein.

All cDNA constructs (Supplementary Table 1) were confirmed by sequencing (TibMolBiol).

FRAP and photoactivation experiments. HeLa cells were transiently transfected with LANCL2-GFP or LANCL2sh-GFP for FRAP experiments, or with LANCL2-PAGFP or LANCL2-G2A-PAGFP for FDAP experiments, and processed 48 h after transfection. Images were acquired by a Nikon A1 confocal microscope (Nikon Corporation, Tokio, Japan), equipped with a 60 X PlanApo oil immersion objective (NA 1.40). All images, corrected for fluorescence background, were analysed with ImageJ 1.48v (Wayne Rasband, Nat. Inst. of Health, USA), and the obtained data with the Excel software (Microsoft).

For FRAP experiments, the EGFP fluorescence emission was bleached in a 6 μm-diameter ROI for 0.12 s, with the 405 nm laser line, zooming to the ROI to minimize the photobleaching duration. Fluorescence recovery was followed for 120 s, exciting the sample with the 488 nm Argon Laser-line.

At each time point the mean fluorescence value (Ft) in the bleached ROI was analysed as described: each ROI F0 was normalized for the F0 of a ROI not subjected to photobleaching, to correct for photobleaching of EGFP due to the imaging procedure. Photobleaching-corrected Ft values were subtracted of first post-photobleaching values and normalized for the mean fluorescence value measured before photobleaching.

Fluorescence recovery traces were fitted with a single exponential equation (GraphPad Prism, version 5.01 for Windows, California, USA), as follows:

$$F_t = F_{\text{max}}(1 - e^{-t/t_\text{r}})$$

which allowed to calculate the half-time constant (t½), the time it takes for the fluorescence to recover to 50% of the asymptote (plateau) intensity (Fmax), as $$t_{\text{r}} = \frac{\ln 2}{\tau}$$.

For each experimental condition, all curves were averaged and t½ values of different populations were compared using Student’s t-test.

ROIs of different size were analyzed, yielding similar results.
Photoactivation was performed by exciting the PAGFP with the 405 nm laser line inside a 8 μm-diameter ROI for 0.063 s. The fluorescence decay in the photoactivated region was imaged for 3 min for fast-diffusing LANCL2-G2A-PAGFP and for 25 min for LANCL2-PAGFP, with 488 nm laser line.

As control, we used PFA-fixed HeLa cells expressing LANCL2-PAGFP to monitor the potential photobleaching of PAGFP during the observation time.

No bleaching was observed in fixed cells in the time interval of our experiments, nor in ROIs chosen in living transfected HeLa cells after whole-cell photoactivation (data not shown).

After photoactivation, the mean fluorescence value (Ft) of the ROI at each time point was corrected for the fluorescence measured before the photoactivation; then it was normalized to F0, where t0 is the first frame after the photoactivation.

Data were fitted with a single exponential decay equation, according to

$$F_t = (F_0 - F_{\infty}) e^{-\frac{t}{t_{1/2}}} + F_{\infty}$$

where $F_{\infty}$ is the normalized fluorescence value at infinite time. The half-life ($t_{1/2}$) for each experimental condition was averaged, and the different populations were compared using Student’s t-test.

**FRET imaging and analysis.** Fluorescence (or Förster) Resonance Energy Transfer (FRET) allows the study of protein interactions, measuring a non-radiative transfer of energy from an excited state or a fluorophore (donor) to a different fluorophore (acceptor), over distances comparable to the size of proteins. As stated by the non-invasive method of sensitized emission (SE) applied to live specimens, following donor excitation, FRET results in a decrease in donor emission with a simultaneous increase in acceptor emission, due to a dipolar interaction between the molecules.

The expression of EGFP and TagRFP tagged proteins in HeLa cells grown in 8-wells chambered coverglasses (Lab-Tek, Nunc, Thermo Scientific) was visualized 2 days after transfection by confocal microscopy using a Leica TCS SP5 AOBS (Leica Microsystem, Heidelberg, Germany) equipped with an oil immersion objective 63x/1.4 NA. Light collection configuration was optimized according to the combination of chosen fluorochromes, selecting the spectral windows by the acousto-optic beam splitter of the Leica SP5 scanning head.

The Leica “LAS AF” software package was used for image acquisition. FRET efficiency (FE) was measured using the Sensitized Emission method. Briefly, measurements were performed by detection of the fluorescent signals of the donor, FRET and acceptor in a line by line sequential scan acquisition, obtained with the Leica Microsystem “FRET SE Wizard” software. The same process was applied to donor-only and acceptor-only reference specimens, on cells chosen to visually display mean fluorescence intensity. From the images obtained, we calculated the calibration coefficients required to correct for excitation and emission cross-talk.

FE was calculated in plasmamembrane-representing or in whole-cell-including ROIs, for 8–15 cells in each experiment, according to $FE = (B - (A - \gamma C))/C$, where A, B, C are the intensities of the donor, FRET and acceptor emissions and $\gamma$ and $A$ are the calibration factors correcting for donor cross-talk and acceptor cross-excitation, respectively.

**Statistical Analysis.** Data were compared using an unpaired Student’s t test. Statistical significance was set at P-value < 0.05. Statistical analysis was performed using the GraphPad Prism Software, or Microsoft Excel software.

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Author Contributions
C.F. and E.Z. designed the study. C.F., T.V., L.G., V.B., S.B., L.S., C.U., M.D.B. and M.P. performed the experiments; C.F. analysed the data; C.F., C.U., A.D.F. and E.Z. interpreted data. C.F., A.D.F. and E.Z. wrote the manuscript.

Additional Information
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