LuTHy: a double-readout bioluminescence-based two-hybrid technology for quantitative mapping of protein–protein interactions in mammalian cells

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Abstract

Information on protein–protein interactions (PPIs) is of critical importance for studying complex biological systems and developing therapeutic strategies. Here, we present a double-readout bioluminescence-based two-hybrid technology, termed LuTHy, which provides two quantitative scores in one experimental procedure when testing binary interactions. PPIs are first monitored in cells by quantification of bioluminescence resonance energy transfer (BRET) and, following cell lysis, are again quantitatively assessed by luminescence-based co-precipitation (LuC). The double-readout procedure detects interactions with higher sensitivity than traditional single-readout methods and is broadly applicable, for example, for detecting the effects of small molecules or disease-causing mutations on PPIs. Applying LuTHy in a focused screen, we identified 42 interactions for the presynaptic chaperone CSPα, causative to adult-onset neuronal ceroid lipofuscinoses (ANCL), a progressive neurodegenerative disease. Nearly 50% of PPIs were found to be affected when studying the effect of the disease-causing missense mutations L115R and ΔL116 in CSPα with LuTHy. Our study presents a robust, sensitive research tool with high utility for investigating the molecular mechanisms by which disease-associated mutations impair protein activity in biological systems.

Keywords: CSPα; LuTHy; missense mutations; protein–protein interactions; quantitative

Subject Categories: Genome-Scale & Integrative Biology; Methods & Resources; Network Biology

Method

Introduction

Protein–protein interactions (PPIs) play an essential role in the proper functioning of living cells (Perkins et al, 2010; Cafarella et al, 2017). They transmit information between signaling proteins, regulate enzymatic activities of proteins, and control the cellular tasks of molecular machines (Couzens et al, 2013; Taipale et al, 2014; Arumughan et al, 2016). Mutation-dependent perturbations of PPIs play a crucial role in the development of diseases (Wang et al, 2012; Sahni et al, 2015). The systematic identification and characterization of interactions between human proteins is therefore of critical importance to better understand complex biological processes and molecular mechanisms of pathology (Stelzl et al, 2005; Bushell et al, 2008; Markson et al, 2009; Rolland et al, 2014; Hutlin et al, 2017).

Several genetic methods for the identification of PPIs in mammalian cells have been described (Buntru et al, 2016), including mammalian protein–protein interaction trap (MAPPIT; Lievens et al, 2016), fluorescence resonance energy transfer (FRET; Grünberg et al, 2013), single molecule pull-down (SiMPull; Jain et al, 2011), dual luminescence-based co-precipitation (DULIP; Trepte et al, 2015), avidity-based extracellular interaction screen (AVEXIS; Bushell et al, 2008), or mammalian membrane two-hybrid (MaMTH; Petschnigg et al, 2014), which all use different biophysical principles and terminal readouts for PPI detection. While in FRET-based assays, the association of proteins is monitored directly through the energy transfer of tagged fluorescent reporter proteins (Vogel et al, 2006), in MAPPIT or MaMTH, PPIs are detected indirectly through auxiliary reactions of reporter proteins (Petschnigg et al, 2014; Lievens et al, 2016). In some methods, for example, MaMTH, interactions are detected in vivo under steady state conditions, enabling the identification of transient and phosphorylation-dependent PPIs (Petschnigg et al, 2014). In others, mammalian cells expressing tagged hybrid proteins are lysed before binary interactions are detected in crude extracts. In

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DULIP or LUMIER (luminescence-based mammalian interactome mapping), for example, PPIs are monitored by luminescence in co-precipitations of protein complexes (Barrios-Rodiles et al., 2005; Trepte et al., 2015). In contrast to MaMTH, these methods preferentially detect PPIs with high binding affinities, while weak interactions are less efficiently identified (Trepte et al., 2015). Due to this diversity in procedures, underlying biophysical principles, and detection readouts, it is not surprising that largely complementary results are obtained when positive PPI reference sets are analyzed systematically (Braun et al., 2009; Venkatesan et al., 2009; Chen et al., 2010; Lievens et al., 2014). To create an improved technology for more comprehensive and sensitive identification of PPIs than currently possible, it seems advisable to combine different detection principles.

Here, we present LuTHy, a bioluminescence-based two-hybrid method that enables the detection of binary PPIs with high sensitivity and specificity in mammalian cells through the combination of two readouts in one experiment. Interactions between ProteinA-mCitrine- and NanoLuc-tagged hybrid proteins are first detected in vivo through quantification of BRET and subsequently ex vivo through a luminescence-based co-precipitation (LuC). We benchmarked LuTHy against known interactions and performed a targeted screen to identify interactions of the synaptic protein CSPα, which has a role in adult-onset neuronal ceroid lipofuscinosis (ANCL) and other neurodegenerative disorders (Nosková et al., 2011; Sharma et al., 2012; Burgoyne & Morgan, 2015). The disease relevance of our findings and further applications of LuTHy for systematic PPI mapping are discussed.

Results

Development of LuTHy

Our initial motivation in establishing LuTHy was to increase sensitivity of PPI detection in order to generate datasets of higher confidence and density than previously possible. To achieve this, we developed a method that can detect both weak and strong as well as direct and indirect PPIs in one procedure, using two different biophysical detection principles.

Direct interactions between proteins of interest (X and Y) are detectable in cells through quantification of bioluminescence resonance energy transfer (BRET; Pfleger & Eidne, 2006), when the tagged luciferase donors and fluorescent acceptors come into close proximity (<10 nm; Dacres et al., 2012). We first constructed plasmids for the co-production of NanoLuc luciferase (NL) and ProteinA-mCitrine (PA-mCit) hybrid fusion proteins in mammalian cells to measure BRET (Fig 1A and Appendix Fig S1A and B). We assumed that protein complex formation between NL-X and PA-mCit-Y should subsequently be detectable in lysed cells with a bioluminescence-based co-precipitation (LuC) assay (Fig 1A). Here, the PA-tag in the hybrid protein is used for bait precipitation and the NL-tag for the detection of the interacting prey (Fig 1A).

To prove our concept, we used the high-confidence interaction BAD/BCL2L1, shown with several PPI detection assays (Braun et al., 2009; Trepte et al., 2015). Plasmids encoding the hybrid proteins PA-mCit-BAD and NL-BCL2L1 as well as the control proteins PA-mCit, NL, PA-NL, and PA-mCit-NL were generated (Fig 1B and Dataset EV1), and recombinant protein production was confirmed by SDS–PAGE and immunoblotting (Appendix Fig S1C–E).

We first assessed the interaction between BAD and BCL2L1 hybrid proteins with in-cell BRET measurements. We co-expressed PA-mCit-BAD/NL-BCL2L1, PA-mCit-BAD/NL, PA-mCit/NL-BCL2L1, and PA-mCit/NL in HEK293 cells and quantified luminescence emission at 370–480 and 520–570 nm 48 h after transfection (Appendix Fig S2). Emission was also quantified with the positive control PA-mCit-NL and the control PA-NL, used to correct for donor luminescence bleed-through (Pfleger et al., 2006). BRET ratios were calculated (Appendix Fig S1F) and found to be significantly higher in cells co-producing PA-mCit-BAD/NL-BCL2L1 compared to the control interactions (Fig 1C), indicating that the BAD/BCL2L1 interaction can be detected through quantification of BRET. A high BRET ratio was also obtained for PA-mCit-NL (Fig 1C), where NL was directly linked to PA-mCit.

Next, we investigated PPI detection in LuC assays. We lysed the transiently transfected cells directly after BRET measurement and quantified fluorescence emission at 530 nm (mCitIN) as well as the total luminescence emission after substrate addition (NLIN) in crude protein extracts (Appendix Fig S3A) as well as the previously established reference sets hPRS (Homo sapiens positive) and hRRS (H. sapiens random; Braun et al., 2009), containing 81 and 80 PPIs, respectively (Appendix Fig S4A and B). Two expression plasmids were constructed for each protein to produce both NL- and PA-mCit-tagged fusions, allowing analysis of all proteins both as baits and preys.

LuTHy detects binary interactions with high specificity and sensitivity

To evaluate the performance of LuTHy more comprehensively, we examined the previously established reference sets hPRS (Homo sapiens positive) and hRRS (H. sapiens random; Braun et al., 2009), containing 81 and 80 PPIs, respectively (Appendix Fig S4A and B). Two expression plasmids were constructed for each protein to produce both NL- and PA-mCit-tagged fusions, allowing analysis of all proteins both as baits and preys.

In addition to potential interactions (NL-X/PA-mCit-Y), two control interactions (NL-X/PA-mCit and NL/PA-mCit-Y) were tested systematically, enabling calculation of corrected BRET (cBRET) and LuC (cLuC) ratios (Appendix Fig S1F and G) for the entire reference sets (Appendix Figs S4C and D, and S5A and B; Datasets EV2 and EV3). We obtained cBRET and cLuC ratios for 303 binary interactions from hPRS and hRRS, detecting PPIs with high reproducibility in independent experiments with both readouts (Fig 2A and B), indicating that LuTHy is suitable for larger scale screening. Receiver operating characteristic (ROC) analysis was performed to define the optimal cutoff for recovering positive interactions from hPRS.
“True” interactions in this PPI set were detected with cutoffs of \( \geq 0.01 \) and \( \geq 0.03 \) for the cBRET and cLuC ratios, respectively. Next, we also assessed whether the subcellular localization of the tested interaction pairs influences the success rate of PPI detection with LuTHy assays. Using the protein pairs in hPRS and hRRS, we performed ROC analysis for known cytoplasmic, nuclear, and membrane proteins to define the optimal cutoffs for PPI detection. We found that the initially defined cutoffs of \( \geq 0.01 \) and \( \geq 0.03 \) for cBRET and cLuC ratios, respectively, are well suited to define “true” PPIs where at least one of the tested proteins per interaction pair is a cytoplasmic, a nuclear, or a membrane protein (Appendix Fig S5C–E). However, for protein pairs where both tested proteins are known to be membrane-associated or contain membrane-spanning domains more stringent cutoffs of \( \geq 0.03 \) (cBRET) and \( \geq 0.05 \) (cLuC) are required to define “true” interactions (Appendix Fig S5F).

Thus, using two cutoffs, 40 binary interactions (49.4%) were finally recovered from hPRS, of which 36 (44.4%) were identified with BRET, 22 (27.2%) with LuC and 18 (22.5%) with both readouts (Fig 2D). When combining the results from both readouts, LuTHy detects PPIs in hPRS with higher success rates than KISS (Lievens et al, 2014), LUMIER (Barrios-Rodiles et al, 2005; Braun et al, 2009), Y2H (Venkatesan et al, 2009; Chen et al, 2010), and several other methods (Appendix Fig S6A and C), which have success rates between 25 and 35% (Braun et al, 2009; Lievens et al, 2014). Compared to the older methods, LuTHy detected both new and previously identified PPIs in hPRS, confirming that our approach to combine two independent readouts (BRET and LuC) is a valid strategy. Using the defined cutoffs, two interactions (2.5%) were recovered from hRRS, none of which was double-positive (0.0%), substantiating that the double-readout method detects PPIs in reference sets with very high specificity and sensitivity (Appendix Fig S6B and D).
LuTHy detects low- and high-affinity interactions

To investigate whether LuTHy is able to detect interactions with different binding affinities (K_Ds), we next screened an affinity-based interaction reference set, termed AIRS, including 71 interactions with known K_D values (Fig 3A and B and Dataset EV4). We assessed each protein in AIRS both as bait and prey in two independent experiments. Using the established cutoffs (Appendix Fig S5C–F), we found that both BRET and LuC assays in LuTHy detected AIRS PPIs with high reproducibility (Appendix Fig S7A and B). LuTHy recovered 35 (49.3%) of 71 binary interactions (Appendix Fig S7C), a similar success rate as with the hPRS (Fig 2D). We identified 31 (43.7%) interactions with the BRET read-out, 19 (26.8%) with LuC (Appendix Fig S7C–E), and 15 (21.1%) with both, confirming the two components of LuTHy to yield complementary but also overlapping results.

To assess the influence of interaction strength on the recovery of PPIs, groups of interactions with low, medium, and high binding affinities (Fig 3B) were analyzed. We found that in-cell BRET assays detect strong and weak interactions with similar success rates (Fig 3C), indicating binding-strength independence. LuC, in strong contrast, recovered high-affinity interactions with significantly higher success rates than low-affinity interactions (Fig 3C and D), as previously observed for co-precipitation-based methods (Trepte et al., 2015).

One of the high-affinity interactions (K_D = 0.3 fM; Johnson et al., 2007) from AIRS, between the ribonuclease RNASE1 and its inhibitor RNH1, is readily detectable with LuC but not with BRET (Appendix Fig S8A–D). This supports binding affinity as the critical parameter for detecting PPIs with LuC. With in-cell BRET, however, resonance-energy-transfer efficiency critically depends on the proximity and the orientation of donor and acceptor molecules (Dacres et al., 2012). As FRET (Vogel et al., 2006) and DULIP (Trepte et al., 2015) assays are based on similar detection principles as the BRET and LuC assays, respectively, we used them for the validation of this specific interaction. While no significant FRET efficiency could be measured (Appendix Fig S8E), we obtained high background-corrected normalized luminescence-based interaction ratios (cNI Rs) using the DULIP assay (Appendix Fig S8F–H).

BRET50 ratios provide relative binding strengths for binary interactions

Previous studies indicate that BRET assays can determine the relative binding strength between proteins when performed as donor saturation assays, in which increasing acceptor/donor ratios are
systematically analyzed (Appendix Fig S9A; Drinovec et al., 2012; Lavoie et al., 2013). We therefore produced saturation curves and calculated BRET50 values with high reproducibility (Appendix Fig S9B) for 25 of the 31 BRET-positive interactions in AIRS (Appendix Fig S9C). Next, we plotted these values against published KD values determined with cell-free biochemical methods (Wang et al., 2004; Kastritis et al., 2011) and observed a highly significant correlation between BRET50 and published KD values (Fig 3E). This shows that BRET50 values are indeed indicative of protein binding affinities and can provide valuable additional information about the strengths of NL- and PA-mCit-tagged hybrid-protein interactions.

PPIs detected with LuTHy can be confirmed by fluorescence and bioluminescence imaging (BLI)

In order to assess whether PPIs detected with LuTHy can be validated with imaging techniques, we examined the well-studied, functional interaction between the synaptic proteins Syntaxin-1 and Munc18 (Rizo & Südhof, 2012), including the Munc18 variants with single and double point mutations (K46E, E59K, and K46E/E59K). K46E and E59K significantly reduce the binding affinity of Munc18 to Syntaxin-1; the double mutation even prevents the interaction (Han et al., 2011). First, NL- and PA-mCit-tagged fusion protein expression was confirmed by SDS-PAGE and immunoblotting (Appendix Fig S10A). Systematic analysis with LuTHy revealed a strong BRET signal for the PA-mCit-Syntaxin-1/NL-Munc18-WT interaction, while lower BRET ratios were detected with NL-Munc18-K46E and -E59K (Appendix Fig S10B). No interaction was observed with the double mutant NL-Munc18-K46E/E59K, confirming published results (Han et al., 2011). Donor saturation curves and BRET50 analysis confirmed the impact of the point mutations on PPI detection (Appendix Fig S10C and D).

Finally, we assessed the influence of the mutations in Munc18 on the subcellular localization of Syntaxin-1 with fluorescence microscopy and BLI (Mezzanotte et al., 2017). Munc18 is critical for targeting Syntaxin-1 to the plasma membrane in mammalian cells (Han et al., 2011). We co-expressed PA-mCit-Syntaxin-1 with NL-Munc18-WT, -K46E, -E59K, or -K46E/E59K in HEK293 cells and
analyzed the Syntaxin-1 localization with fluorescence microscopy. PA-mCit and PA-mCit-NL were investigated as controls. They were distributed throughout the cytoplasm (Appendix Fig S10E), while PA-mCit-Syntaxin-1 was predominantly located in the perinuclear region in the absence of NL-Munc18-WT (Fig 4A). In its presence, PA-mCit-Syntaxin-1 was targeted to the plasma membrane, confirming published reports (Han et al., 2011). Next, the NanoLuc substrate was added and cells were assessed by BLI, with luminescence emission recorded at 460 and 535 nm using appropriate band-pass (BP) filters (Fig 4A and Appendix Fig S10E). Finally, BRET images were calculated by dividing the intensities recorded at the longer wavelength by those recorded at the shorter one. We observed a strong BRET signal at the plasma membrane in cells co-producing PA-mCit-Syntaxin-1 and NL-Munc18-WT, indicating a direct interaction (Fig 4A). In comparison, BRET signals at the plasma membrane were reduced or undetectable with the mutant variants, confirming the initial BRET measurements in microtiter plates (Appendix Fig S10B and C). Interestingly, in cells co-producing NL-Munc18-K46E or -E59K, an accumulation of PA-mCit-Syntaxin-1 in the perinuclear region was observed, probably attributable to the loss of interaction at the plasma membrane.

**Effects of chemical compounds and heat stress on binary interactions**

Targeting disease-relevant interactions with small molecules has great potential for the development of new medicines, but discovering drugs that potently and specifically stimulate or disrupt PPIs remains a challenge (Scott et al., 2016). To assess whether LuTHy can detect direct compound effects on PPIs, we first performed proof-of-principle experiments with the well-known immunosuppressant rapamycin, which was previously shown to stimulate the association of FKBP12 and the FRB domain of mTOR (Machleidt et al., 2015). We incubated HEK293 cells co-producing NL-FKBP12 and PA-mCit-FRB with different concentrations of rapamycin and quantified BRET and LuC ratios after 4 h. We observed a nanomolar, concentration-dependent stimulation of the interaction with BRET and LuC (Fig 4B), confirming earlier reports (Machleidt et al., 2015).

Next, we investigated the direct effect of Nutlin-3, an anti-cancer agent previously shown to disrupt the association between MDM2 and the tumor suppressor p53 (Lievens et al., 2014), on the NL-MDM2 and PA-mCit-p53 interaction. We incubated HEK293 cells with different concentrations of Nutlin-3 and monitored the association 6 h after compound addition. We observed that the compound potently inhibited (IC50 ~8 and 11 nM) the interaction in mammalian cells (Fig 4C), confirming previously published results (Lievens et al., 2014).

We hypothesized that besides direct also indirect compound effects on PPIs should be detectable with the LuTHy technology. To address this question, we next investigated the effects of the small molecules ganetespib and geldanamycin on the self-association of heat shock factor 1 (HSF1) in mammalian cells. This protein is known to trimerize under heat stress conditions, leading to transcriptional activation of heat shock genes, such as HSF70 and HSF40 (Morimoto, 1993; Gil et al., 2017). Furthermore, previous studies have demonstrated that a cellular response similar to a heat shock can be induced upon treatment of cells with the Hsp90 inhibitors ganetespib and geldanamycin (Stebbins et al., 1997; Lin et al., 2008), suggesting that these compounds might indirectly influence HSF1 oligomerization. To address this, we co-expressed NL-HSF1 and PA-mCit-HSF1 in HEK293 cells and, 24 h post-transfection, treated the cells with the chemical compounds ganetespib and geldanamycin. BRET ratios were quantified after 1, 3, 6, 12, 18, 24, and 48 h in compound-treated and untreated cells. In comparison with untreated cells, we observed a significant increase in the BRET signal after 3 h in compound-treated cells, which slowly decreased over time (Fig 4D), indicating that both compounds indirectly stimulate the self-association of HSF1.

Finally, we assessed whether heat shock conditions also stimulate the association of HSF1 in mammalian cells (Rabindran et al., 1993). We exposed HEK293 cells co-expressing the proteins NL-HSF1 and PA-mCit-HSF1 to heat shock (42°C) and monitored the association of the fusion proteins by quantification of BRET after 90, 180, or 360 min. In addition, BRET was quantified in cells before the heat shock and after a recovery phase of 18 h at 37°C. We observed a rapid increase of the BRET signal upon heat stimulation, which decreased again to normal levels during the recovery phase at 37°C (Fig 4E). Thus, the impact of heat stress on PPIs can be readily monitored using the LuTHy method.

**Figure 4. Effects of missense mutations and small molecules on PPI detection with LuTHy.**

A Live-cell bioluminescence imaging (BLI) of HEK293 cells co-producing the indicated fusion proteins. mCitrine was excited at 500 nm, and emitted fluorescence was detected at 535 nm. After substrate addition, short (460) and long (535) band-pass (BP) filters in a dual-view adapter were used to detect the emitted luminescence simultaneously at the respective wavelengths. BRET images were calculated by dividing the 535 BP by the 460 BP images using ImageJ. Scale bar = 20 μm.

B Quantification of the rapamycin-induced interaction between NL-FKBP12 and PA-mCit-FRB. Plasmids encoding the fusion proteins were co-transfected in HEK293 cells and 24 h later treated with the indicated concentrations of rapamycin. After an additional incubation for 4 h, BRET was quantified upon substrate addition. After cell lysis in buffer with indicated rapamycin concentrations, LuC ratios were determined. EC50 values were obtained with nonlinear curve fitting. Data are representative of at least two independent experiments. All values are mean ± s.d.

C Effects of Nutlin-3 on the interaction between NL-MDM2 and PA-mCit-p53. Same procedure as in (B) except for 6 h treatment with Nutlin-3. Data are representative of at least two independent experiments. All values are mean ± s.d.

D Time resolved HSF1 oligomerization by Hsp90 inhibition. HEK293 cells co-producing NL-HSF1 and PA-mCit-HSF1 for 24 h were transferred to 384-well plates containing ganetespib or geldanamycin to reach a final concentration of 1 μM. Luminescence was measured at the indicated time points and the calculated BRET ratios normalized to the respective untreated control. Data are mean of two biological replicates. All values are mean ± s.e.m. Significance was calculated by two-way ANOVA followed by Dunnett’s multiple comparison’s post hoc test. *P < 0.05, **P < 0.005, ***P < 0.001.

E Schematic depiction of a heat shock experiment. HEK293 cells co-producing NL-HSF1/PA-mCit-HSF1 for 48 h were subjected to a heat shock at 42°C for 6 h, after which the cells were recovered at 37°C for 24 h. Luminescence was measured at the indicated time points: immediately before the heat shock, during the heat shock at 90, 180, and 360 min, and after a recovery phase of 24 h. Calculated BRET ratios were normalized to a control plate that was not heat-treated. All values are mean ± s.e.m. from three independent experiments. Significance was calculated by one-way ANOVA followed by Dunnett’s multiple comparisons post hoc test; *P < 0.05.
Figure 4.

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The positioning of fusion tags in hybrid proteins influences the success rate of PPI detection

Previous studies indicate that the positioning of fusion tags either at the N- or C-terminus of proteins of interest influences the success of PPI detection with two-hybrid methods (Stellberger et al., 2010). To address this question systematically, we generated expression plasmids to produce the protein VCP (valosin-containing protein) and 10 ubiquitin regulator X (UBX) domain-containing proteins (p37, p47, UBXD1-6, UBXD8, and UBXD9) as N- and C-terminally tagged NL- and PA-mCit-fusions in mammalian cells (Fig 5A, and Datasets EV1 and EV5). Previously, interactions between VCP and all of these 10 UBX-domain-containing proteins have been reported (Raman et al., 2015), suggesting that they should also be detectable with the LuTHy technology. We systematically tested each interaction (e.g., between VCP and p37) in the eight possible hybrid-protein combinations, determining cBRET and cLuC ratios for 80 PPIs in HEK293 cells. We found that all reported PPIs are detectable with both BRET and LuC.
and LuC measurements in LuTHy assays (Fig 5B and C), confirming the previous results (Raman et al., 2015). However, it is important to note that the success of PPI detection with both readouts is influenced by the site of tagging (Dataset EV5). We found that 46 (57.5%) and 41 (51.3%) of the 80 tested binary interactions scored positive in BRET and LuC assays, respectively, indicating that multiple combinations of hybrid proteins reveal positive PPIs in systematic LuTHy experiments.

It is known that UBX proteins bind with their C-terminal UBX-domain to the N-domain of VCP (Fig 5A). Accordingly, the highest cBRET ratios were obtained when VCP was tagged at its N-terminus and the UBX proteins at their C-terminus, respectively (Fig 5D). Exceptions were UBXD1, which binds to the C-terminus of VCP and UBXd9, which is known to remodel VCP into a heterotetrameric structure (Arumughan et al., 2016). This suggests that for BRET experiments, fusion tags should be added in close proximity to the interaction domain. In contrast, the LuC results showed no clear preference for fusion-tag orientation. However, it seems that the highest cLuC ratios are obtained when VCP was used as the bait (Fig 5D), indicating that it is crucial for PPI detection with LuC, which of the two proteins of interest contains the PA-tag.

Finally, we investigated whether the LuTHy technology also detects PPIs when the hybrid proteins are expressed in cells at similar-to-endogenous protein levels. To address this question, we focused on the interaction between VCP and UBXd9 that was repeatedly detected in BRET and LuC assays (Fig 5B–D). We first performed transfection experiments with different concentrations of plasmid DNA and determined that concentrations of 1 and 25 ng of the plasmids pcDNA3.1-NL-VCP and -UBXd9-mCit, respectively, are sufficient to detect luminescence and fluorescence activities in HEK293 cells (Appendix Fig S11A and B). Analysis of cell extracts by SDS–PAGE and immunoblotting revealed that under these experimental conditions, the recombinant proteins are expressed at endogenous protein levels or even below (Appendix Fig S11C and D). Next, we co-transfected the plasmids into HEK293 cells (1 and 25 ng) and assessed their association by quantification of BRET. In comparison with control interactions (NL-VCP/mCit or UBXd9-mCit/NL), we detected a significant BRET signal in NL-VCP/UBXd9-mCit co-expressing cells (Appendix Fig S11E), indicating that the LuTHy technology detects PPIs, even when the hybrid proteins are expressed at very low levels in cells.

### Identification of CSPα-interaction partners

To assess whether LuTHy can be applied for systematic protein interaction screening, we generated a target library of expression plasmids encoding 125 NL-tagged presynaptic proteins that play a functional role in synaptic transmission. We selected various components of the exocytosis/endocytosis machinery as well as regulators of this process (Appendix Fig S12A and Dataset EV6). In addition, we created a plasmid for the expression of PA-mCit-tagged CSPα, a molecular chaperone in the presynapse (Sharma et al., 2011). CSPα maintains protein homeostasis in the presynapse and likely associates with multiple proteins that play a role in synaptic transmission (Zhang et al., 2012; Donnelier & Braun, 2014). To this day, however, only a few direct binding partners such as SNAP25 or dynamin-1 have been identified (Sharma et al., 2011; Zhang et al., 2012), suggesting that additional PPIs can be found by targeted interaction screening.

Using the established cutoffs, we systematically screened 125 protein pairs in two independent experiments (Appendix Fig S12B and C), identifying a total of 42 interactions with CSPα-mCit-PA (Fig 6A, Appendix Fig S13A and B and Dataset EV6); 36 PPIs were identified with the BRET and seven with the LuC readout (Appendix Fig S13A and B and Dataset EV6). Three (7.1%) of the 42 identified PPIs were already known (Dataset EV7), while over 90% of PPIs detected with LuTHy have not been reported. A large fraction of the detected interactors are SNARE complex members and proteins involved in vesicle fusion processes (Dataset EV8), confirming a functional role of CSPα in synaptic vesicle exocytosis (Sharma et al., 2011).

Our analysis revealed that a relatively large fraction (22 of 42, 52%) of the identified CSPα interaction partners are membrane-associated or membrane domain-containing proteins (Appendix Fig S14A, and Datasets EV6 and EV8). This is not unexpected due to the fact that CSPα in cells gets palmitoylated (Greaves et al., 2012) and probably is attached to membranes, when it interacts with NL-tagged partner proteins. To assess whether CSPα interacts specifically with membrane proteins, we performed an additional focused PPI screen with the protein PICK1 (Thomas et al., 2013), which, similar to CSPα, is palmitoylated in cells and attaches to membranes (Appendix Fig S14B). Quantification of cBRET and cLuC ratios confirmed the interactions between CSPα-mCit-PA and the 22 tested NL-tagged membrane-associated or membrane domain-containing proteins (Appendix Fig S14C and D, Dataset EV9). In contrast, no or very weak interactions were detected with PICK1, substantiating our result that CSPα interacts specifically with membrane proteins.

Finally, we investigated whether the interactions between CSPα and membrane-associated or membrane domain-containing proteins can be also detected in primary hippocampal neurons under endogenous conditions. We selected five positive and one negative interaction from the PPI dataset and performed in situ proximity ligation assays (PLAs) using target-specific antibodies. With this approach, all LuTHy positive PPIs such as the association of CSPα with SLC17A7, SYP, or SYT1 were confirmed in neuronal cells, while the LuTHy negative interaction with STX4 could not be detected (Appendix Fig S14E). This indicates that biologically relevant interactions with membrane proteins are identified with the LuTHy technology.

### CSPα preferentially associates with synaptic proteins

The relatively high success rate of PPI detection (42 of 125, 34%) in our initial interaction screen suggests that CSPα preferentially interacts with synaptic proteins involved in endo-/exocticotic processes (Dataset EV6). As confirmation, we performed an additional interaction screen with CSPα-mCit-PA and a target library of 131 mostly non-synaptic NL-tagged proteins. With this protein set, we identified a total of 18 PPIs (14%), of which 12 interactions were identified with the BRET and eight with the LuC readout (Appendix Fig S14F, Dataset EV10). Interestingly, the success rate of PPI detection with CSPα was significantly lower among non-synaptic than among synaptic proteins (Appendix Fig S14G), supporting our hypothesis that CSPα has a preference of interacting with synaptic proteins. This is in agreement with previous observations that CSPα is a...
presynaptic co-chaperone that is critical for synapse maintenance (Sharma et al., 2011).

Finally, our analysis revealed that six of the 18 (33%) identified CSPα interacting partners in the non-synaptic protein set are membrane-associated or contain membrane-spanning domains (Appendix Fig S14H), suggesting that the chaperone has a bias for such proteins. However, statistical analysis revealed that CSPα has no significant preference for interacting with membrane or membrane-associated proteins in both tested protein sets (synaptic and non-synaptic; Appendix Fig S14H).

Disease-causing mutations influence the association of CSPα with partner proteins

Two mutations in CSPα (AL116 and L115R, Fig 6A) cause adult-onset neuronal ceroid lipofuscinosis (ANCL), a rare neurological disease characterized by dementia and impairment of muscle coordination (Josephson et al., 2001; Nosková et al., 2011). To address whether these mutations influence the association of CSPα with partner proteins, we tested whether the mutant fusion proteins CSPα(AL116)-mCit-PA and CSPα(L115R)-mCit-PA show altered interactions with NL-tagged partner proteins in comparison with wild-type CSPα-mCit-PA. Interactions with the mutant variants in comparison with wild-type CSPα showed many altered cBRET and cLuC ratios (Fig 6B and C, Appendix Fig S15A and B and Dataset EV11), indicating that the mutations influence CSPα PPIs. In general, more interactions with membrane proteins were affected than with non-membrane proteins (Appendix Fig S15C) and in most cases the cBRET or the cLuC signals for membrane proteins were decreased (11 of 13, 85%). In comparison, many non-membrane proteins showed enhanced binding to the mutant variants of CSPα (5 of 7, 71%; Appendix Fig S15A–C and Dataset EV11).
To validate the impact of the mutations on PPIs exemplarily, we performed donor saturation assays with synaptotagmin-1 (SYT1), a calcium sensor for neurotransmitter release (Südhof & Rizo, 2011). Significantly higher BRET$_{50}$ values were obtained with the mutant variants of CSPα compared to the wild-type control protein (Appendix Fig S15D and E), confirming weaker binding to SYT1. In addition to SYT1, many proteins involved in vesicle fusion and neurotransmitter release (LIN7A, SCAMP1, STX3, SYP, SYT5, VAMP2-4) showed altered binding to mutant CSPα, suggesting that the disease-causing mutations might influence synaptic transmission in brains of ANCL patients. Using LuTHy, we found that ~50% of all interactions detected with wild-type CSPα-mCit-PA are disturbed by the two disease mutations (Fig 6D and Dataset EV11).

The disease-causing mutations ΔL116 and L115R promote the association between CSPα and the palmitoyltransferase ZDHHC17

Using LuTHy, a strong interaction between CSPα and the palmitoyltransferase ZDHHC17 was detected with both BRET and LuC readouts, confirming previously published results (Greaves et al., 2012). Palmitoylation of CSPα in presynaptic terminals was reported to be critical for normal protein function (Lopez-Ortega et al., 2017). It occurs in a cysteine-string domain (Fig 6A), which is altered in its amino acid sequence by the disease-causing mutations ΔL116 and L115R, suggesting they might influence palmitoylation efficiency and/or the interaction with the palmitoylating enzymes. We found that the disease-causing mutations promote the association between CSPα and ZDHHC17 (Appendix Fig S15A and B), a result that was also confirmed in independent experiments by applying LuTHy in primary hippocampal neurons showing significantly increased eBRET and LuC ratios with the mutant variants compared to wild-type CSPα (Fig 6E).

SDS–PAGE and immunoblotting of the tagged mutant CSPα fusions revealed that SDS-stable, high-molecular-weight species are detectable in cell lysates besides soluble protein migrating at ~63 kDa (Fig 6F). Such structures were not observed in cells expressing wild-type CSPα-mCit-PA, indicating that at least a fraction of the mutant fusion proteins is abnormally aggregated in mammalian cells. The formation of protein aggregates in the perinuclear region was confirmed by immunofluorescence microscopy (Fig 6G). Furthermore, we observed co-localization of ZDHHC17 with CSPα AL116 and L115R aggregates in cells, suggesting that the stronger interactions observed with mutant CSPα proteins in BRET and LuC assays are caused by protein co-aggregation. This view is supported by recent observations that disease-associated aggregates can recruit a variety of cellular proteins and promote their co-assembly into higher molecular weight structures (Kim et al., 2016).

Discussion

LuTHy is a new double-readout bioluminescence-based two-hybrid method for systematic protein interaction mapping. Performing in-cell BRET, followed by a cell lysis step and an ex vivo luminescence co-precipitation assay with crude protein extract (Fig 1A), it was possible for the first time to obtain two quantitative measurements for tested protein pairs in one procedure. PPIs detected with two readouts provide more information, are potentially of higher confidence, and require less further validation (Braun et al., 2009) than PPIs identified with only one readout. In contrast to earlier methods (Trepte et al., 2015), LuTHy, with its BRET component, identifies low-affinity interactions before cell lysis, circumventing disruption of weak interactions that only occur in the intact cellular environment. In this way, LuTHy increases sensitivity and coverage considerably. Benchmarking the method with well-defined reference sets (Braun et al., 2009; Venkatesan et al., 2009), we found similar reproducibility and specificity than with previously reported single-readout methods. With a sensitivity of ~50% (Fig 2D); however, LuTHy outperforms single-readout methods like MAPPIT or LUMIER that only detect ~35% of PPIs (Braun et al., 2009).

Positive BRET suggests direct binding, as interactors need to come into close proximity (Pfleger & Eidne, 2006). LuC, like all precipitation approaches, does not exclude the possibility that bridging proteins mediate complex formation (De Las Rivas & Fontanillo, 2010) and exhibits a preference for high-affinity PPIs (Fig 3C). Interacting proteins detected with both readouts are likely to associate directly and strongly. However, it is important to note that potential false-positive LuTHy results could emerge from crowding effects due to high hybrid-protein expression or when both tagged proteins of interest are targeted to membrane surfaces. To circumvent this, we generally use low plasmid DNA concentrations to produce recombinant proteins in cells (Appendix Fig S11) and have also implemented a more stringent cutoff specifically for the investigation of membrane protein interactions (Appendix Fig S5F).

Quantitative PPI maps have recently been created using LUMIER with BACON (Taipale et al., 2012), revealing functional chaperone complexes and cellular proteostasis pathways (Taipale et al., 2014). Through its double readout, LuTHy should strongly advance hierarchical clustering and improve the predictions of functional protein modules (Braun et al., 2009).

For validation and also upscaling, we performed targeted PPI screens with libraries of >120 synaptic or non-synaptic control proteins against the presynaptic chaperone CSPα (Donnelier & Braun, 2014). Identification of 42 synaptic proteins, with >90% not yet described as CSPα partners, indicates that only few of the CSPα-interacting proteins in the presynapse are currently known and confirms the protein’s functional role in various synaptic processes (Südhof & Rizo, 2011).

LuTHy is also highly powerful to detect stimulating and disrupting effects of chemical compounds or external stimuli on interactions. It may be a valuable tool to address this challenging area in drug discovery. Arrays of binary interactions, for example, for specific signaling cascades like MAPK (Bandypadhyay et al., 2010) or ErbB (Petschnigg et al., 2014), could be used as quantitative sensors for systematic drug profiling with LuTHy.

Likewise, LuTHy facilitates improved characterization of disease-associated mutations, providing reliable data on increased as well as decreased binding of proteins when mutated. The impact of missense mutations has been systematically assessed with Y2H or LUMIER, with ~60% found to perturb PPIs and ~30% to cause complete loss of interactions (Sahni et al., 2015). Quantitative assessment with LuTHy for the disease mutations in CSPα confirmed this observation (Fig 6D).

Besides its potential to increase coverage and quality, LuTHy can be used in the future to study the dynamics of interactions induced...
by specific stimuli in mammalian cells. Finally, it has an application
in investigating the impact of posttranslational modifications on
PPIs and signaling cascades.

In summary, LuTHy is a powerful method for improved, quan-
titative interactomics studies. Calculation of dimensionless BRET/
LuC ratios improves comparability between labs, a challenge for
all binary PPI detection methods to date. LuTHy detects PPIs
without a bias for or against specific protein classes and is trans-
ferrable to a variety of cell lines. Greater depth of information
per interaction in shorter time and amenability to automation will
make large-scale screening campaigns considerably more time-
and cost-efficient.

Materials and Methods

Reagents and Tools table

| Reagent/resource | Reference or source | Identifier or catalog number |
|------------------|---------------------|-----------------------------|
| **Experimental models** | | |
| HEK-293 cells (H. sapiens) | ATCC | CRL-1573 |
| SHEP cells (H. sapiens) | Muth el al (2010) (Cancer Research) | Prof. Dr. Manfred Schwab, DKFZ, Germany |
| CS7Bl6/ (M. musculus) | Charles River/JAXTM | N/A |
| CS7Bl6/ (M. musculus) | FEM Charité | N/A |
| **Recombinant DNA** | | |
| pmTq2-DEVD-mCit | Grünberg et al (2013) | N/A |
| FR-20-Che-Sp1 | Grünberg et al (2013) | N/A |
| FK-20-Cit(K)-WW | Grünberg et al (2013) | N/A |
| pPA-RL-GW | Trepte et al (2015) | N/A |
| pGW-RL-PA | Trepte et al (2015) | N/A |
| pGW-FL-V5 | Trepte et al (2015) | N/A |
| pNL1.1 | Promega | #N1001 |
| pcDNA3.1 (+) | ThermoFisher | V79020 |
| FUGW | Addgene | #14883 |
| pDONR221 | ThermoFisher | 12536017 |
| pCMV-SPORT6-DNAJS | Open Biosystems | EHS1001-737176 |
| pDONR223-SYT1 | The CCSB Human Orfeome Collection | 12023 |
| pDONR223-RNASE1 | The CCSB Human Orfeome Collection | 5532 |
| pDONR223-RN41 | The CCSB Human Orfeome Collection | 3255 |
| pDONR221-mCherry | Trepte et al (2015) | N/A |
| pDONR221-Munc18 wt | Trepte et al (2015) | N/A |
| pDONR221-Munc18 K46E | Trepte et al (2015) | N/A |
| pDONR221-Munc18 E59K | Trepte et al (2015) | N/A |
| pDONR221-Munc18 K46E/E59K | Trepte et al (2015) | N/A |
| pDS_X-mCherry | ATCC | MBA-303 |
| pDONR221-NL | This study | N/A |
| pDONR221-FR8 | This study | N/A |
| pDONR221-FKBP12 | This study | N/A |
| pDONR221-DNAJC5 | This study | N/A |
| pDONR221-SYT1YTM | This study | N/A |
| pDONR221-VCP | This study | N/A |
| pPA-mCit-mChe | This study | N/A |
| **Additional plasmids and more information** | This study | Dataset EV1 |
| **Antibodies** | | |
| Sheep gamma globulin | Jackson ImmunoResearch | 013-000-002 |
| Reagent/resource                      | Reference or source          | Identifier or catalog number |
|--------------------------------------|------------------------------|------------------------------|
| Rabbit anti-sheep IgGs               | Jackson ImmunoResearch       | 313-005-003                  |
| Goat anti-rabbit HRP                 | Sigma                        | A0545                        |
| Goat anti-mouse HRP                  | Sigma                        | A0168                        |
| Rabbit anti-NanoLuc                  | Promega                      | N/A                          |
| Rabbit anti-GFP                      | Abcam                        | ab290                        |
| Rabbit anti-CSPα                    | Synaptic Systems             | 154 003                      |
| Rabbit anti-H3                       | Abcam                        | ab1791                       |
| Mouse anti-VCP                      | Progen                       | 65278                        |
| Rabbit anti-UBXD9                    | LifeSpan Biosciences         | LS-C156546/EPR8616           |
| Mouse anti-tubulin                   | Sigma                        | 76074                        |
| Mouse anti-VGLUT1                    | Synaptic Systems             | 135 311                      |
| Mouse anti-VGAT                      | Synaptic Systems             | 131 011                      |
| Mouse anti-STX4                      | Synaptic Systems             | 110 041                      |
| Mouse anti-SYP                       | Sigma                        | SVP-38                       |
| Mouse anti-SYT1                      | SantaCruz                    | sc-135574                    |
| Mouse anti-VAMP2                     | Synaptic Systems             | 104 211                      |
| Mouse IgG control                    | ThermoFisher                 | 02-6502                      |

### Oligonucleotides and sequence-based reagents

| PCR primers                          | This study                   | Dataset EV12                 |
|--------------------------------------|------------------------------|------------------------------|

### Chemicals, enzymes and other reagents

| Phusion High-Fidelity DNA Polymerase | ThermoFisher                 | FS30L                        |
|-------------------------------------|------------------------------|------------------------------|
| Gateway™ LR Clonase™ II Enzyme mix  | ThermoFisher                 | 11791100                     |
| DMEM low glucose                    | ThermoFisher                 | 31885049                     |
| DMEM high glucose                   | ThermoFisher                 | 41965062                     |
| DMEM high glucose, HEPES, no phenol red | ThermoFisher             | 21060329                     |
| Fetal bovine serum (FBS)            | ThermoFisher                 | 10270106                     |
| Linear polyethylenimine (PEI), 25 kDa | Polysciences               | 23966-2                      |
| Cell culture microplate, 96-well white | Greiner                   | 659583                       |
| High binding microplate, 384-well white, small volume | Greiner | 784074                       |
| Coelenterazin-h                     | NanoLight                    | 301                          |
| Coelenterazin-h                     | pik                          | 102182                       |
| DPBS                                | ThermoFisher                 | 14190169                     |
| NanoGlo                             | Promega                      | N1120                        |
| Nutlin-3                            | Sigma                        | N6287                        |
| Rapamycin                           | SantaCruz                    | 3504A                        |
| FuGENE® HD                          | Promega                      | E2311                        |
| Geldanamycin                        | Selleckchem                  | S2713                        |
| Ganetesib                           | Selleckchem                  | S1159                        |
| Dual-Clo®                           | Promega                      | E2980                        |
| NuPACE™ NOVEX™ 4-12% Bis-Tris       | ThermoFisher                 | N032380X                     |
| Nitrocellulose membrane             | GE Healthcare                | 10401197                     |
| WesternBright Quantum               | Advansta                     | 12042-020                    |
| ibidi® μ-Slide 8-well               | ibidi                        | 80827                        |
| X-tremeGENE 9                       | Sigma-Aldrich                | 000000000636579001           |
| Duolink® In Situ Detection Reagents Orange | Sigma-Aldrich            | DU092007                     |
Methods and Protocols

**Plasmid construction**

Destination vectors encoding N- and C-terminally tagged fusion proteins were generated based on the pcDNA3.1 (+) vector (Invitrogen). For the acceptor vectors pcDNA3.1-PA-mCit-GW, pcDNA3.1-GW-mCit-PA, and pcDNA3.1-PA-mCit, the coding sequences for mCitrine (mCit) and ProteinA (PA) were amplified from pmTq2-DEVd-mCit (kindly provided by Dr. Raik Gru¨nberg, Center for Genomic Regulation, Barcelona) and pPA-RL-GW (Trepte et al, 2015), respectively. For N-terminal fusions, cDNAs encoding PA and mCit were amplified using the primers #1, to #4, respectively. For generation of plasmids encoding C-terminal fusions, cDNA fragments encoding PA and mCit were amplified using the primers #5 and #6 as well as #7 and #8, respectively. For the control vector pPA-mCit, which contains a stop codon after the mCit coding sequence, the primers #1 and #2 were used to amplify the PA-tag and the primers #3 and #9 to amplify the mCit tag. For the donor vectors pcDNA3.1-NL-GW, pcDNA3.1-GW-NL, and pcDNA3.1-NL, the NanoLuc coding sequence was amplified from pNL1.1 (Promega). For plasmids encoding N-terminal and C-terminal fusions, cDNAs were amplified using the primers #10 and #11 as well as #12 and #13, respectively. For the control vector pNL, the primers #10 and #14 were used to amplify a fragment encoding cmyc-NanoLuc. For all destination vectors, the resulting PCR fragments were cloned into pcDNA3.1 (Invitrogen) via the restriction sites NheI/HindIII. To generate the C-terminal fusion vectors, the restriction sites NheI/HindIII were used. To generate the vector pcDNA3.1-PA-NL, the cDNA encoding the PA-tag was amplified using the primers #19 and #20; the cDNA fragment encoding NanoLuc was amplified using the primers #21 and #22. The resulting PCR products were cloned simultaneously in pcDNA3.1 (+) (Invitrogen) via the restriction sites NheI/EcoRI/HindIII. The vector pmCherryGW (Invitrogen) was digested using the restriction sites Xhol and SgrAI. Then, it was ligated with the self-annealed primer #23 to obtain the plasmid pmCherry lacking the Gateway cassette. Lentiviral LuTHy plasmids were generated by the Viral Core Facility (Charité, Universitätsmedizin Berlin, Germany) based on the FUGW plasmid (FUGW was a gift from David Baltimore, Addgene plasmid #14883). The respective reporter genes and Gateway cassettes were cloned from the pcDNA3.1-based LuTHy vectors to generate pLenti PA-mCit-GW, pLenti-GW-mCit-PA, pLenti-NL-GW, and pLenti-GW-NL.

To generate the entry plasmids pDONR221-NL, pDONR221-FRB, pDONR221-FKBP12, pDONR221-DNAJC5 (CSPα) pDONR221-SYT1 ATM, and pDONR221-VCP the sequences encoding the proteins of interest were PCR amplified and the resulting fragments were shuttled into pDONR221 (Invitrogen) using the BP clonase (Invitrogen). The cDNA encoding NanoLuc was amplified using the primers #24 and #25 from vector pNL1.1 (Promega). The FRB fragment encoding the amino acids 2,021–2,113aa of mTOR was amplified using the primers #26 and #27 from the vector FR-20-Che-Sp1. FKBP12 was amplified using the primers #28 and #29 from the vector FK-20-Cit(K)-WW, which was kindly provided by Dr. Raik Grünberg (Center for Genomic Regulation, Barcelona). The coding sequence of CSPα/DNAJC5 was amplified using the primers #30 and #31 from the pCMV-SPORT6-DNAJC5 vector (Open Biosystems, EHS1001-7373176). To create the SYT1, the entry plasmid lacking the vesicular and transmembrane domain, base pairs 241–1,266 (amino acids 81–422) of SYT1 (CCSB-12023) was amplified using the primers #32 and #33. The coding sequence of VCP lacking the stop codon was amplified from the entry clone IOH52832 using the primers #34 and #35. To generate the CSPα deletion mutation...
AL116 and the single point mutation L115R (Nosková et al., 2011), the vector pDONR221-DNAJC5 was PCR amplified with the phosphorylated primers #36 and #37 (AL116) or #38 and #39 (L115R), respectively. The PCR products were ligated and transformed into Mach1 competent Escherichia coli cells. The DULIP expression plasmids and the entry vectors pDONR221-mCherry, pDONR221-Munc18-wt, -K46E, -E59K, and -K46E/E59K were generated previously (Trepte et al., 2015).

For LuTHy experiments, all cDNAs were shuttled from the entry vectors into the LuTHy destination vectors (pcDNA3.1 or pLenti) using the LR clonase technology according to the manufacturer’s instructions (Invitrogen). Similarly, for DULIP and FRET assays, the cDNAs from the entry vectors pDONR223-RNASE1 (CCSB-5532) and pDONR223-RNH1 (CCSB-3255) were shuttled into the destination vectors pGW-RL-PA, pGW-FL-V5, or pDS-X-mCherry (ATCC® MBA-303™). The plasmid encoding the PA-mCit-mChere fusion protein was generated by LR recombination using the pDONR221-mCherry and pPA-mCit-GW vectors. All used primers can be found in Dataset S1.

**Cell culture, transfection, and lentivirus preparation**

The human embryonic kidney cell line 293 (HEK293) was grown in low-glucose (1 g/l) DMEM (Gibco®), ThermoFisher) for DULIP and FRET experiments and in high glucose (4.5 g/l) for LuTHy assays. In both cases, media were supplemented with 10% heat inactivated fetal bovine serum (Gibco®, ThermoFisher) and cells were grown at 37°C and 5% CO₂. Cells were subcultured every 3–4 days and transfected with linear polyethyleneimine (25 kDa, Polysciences) using the reverse transfection method according to the manufacturer’s instructions. For LuTHy transfections, cells were seeded in phenol-red-free, high-glucose DMEM media (Gibco®, ThermoFisher) supplemented with 10% heat inactivated fetal bovine serum. Transfections were performed with a total DNA amount of ~200 ng per well in a 96-well plate. If expression plasmid concentration was below 200 ng/well, pcDNA3.1 (+) was used as carrier DNA to obtain the total DNA amount of 200 ng.

For LuTHy experiments in hippocampal neurons, all procedures involving animals were approved by the animal welfare committee of Charité Medical University and the Berlin state government. First, hippocampi were dissected from WT mice P0-2 brains in cold Hanks’ salt solution (Millipore), followed by a 30-min incubation in enzyme solution [DMEM (Gibco, ThermoFisher Scientific), 3.3 mM cysteine, 2 mM CaCl₂, 1 mM EDTA, 20 U/ml papain ( Worthington)] at 37°C. Papain reaction was inhibited by the incubation of hippocampi in inhibitor solution DMEM, 10% fetal calf serum (ThermoFisher Scientific), 100 mM BSA (Sigma-Aldrich), and 95 mM trypsin inhibitor (Sigma-Aldrich) for 5 min. Afterward, cells were triturated in NBA (neurobasal-A medium, 2% B27, 1% glutamax, 0.2% P/S, ThermoFisher Scientific) by gentle pipetting up and down.

All lentiviral particles were provided by the Viral Core Facility of the Charité Berlin and prepared as described previously (Lois et al., 2002). Briefly, HEK293T cells were co-transfected with 10 μg of shuttle vector, 5 μg of helper plasmids pCMVdR8.9, and 5 μg of pVSV.G using X-tremeGENE 9 DNA transfection reagent (Roche Diagnostic). Virus containing cell culture supernatant was collected after 72 h and filtered for purification. Aliquots were flash-frozen in liquid nitrogen and stored at −80°C.

**Creation of reference sets**

The CCSB reference sets hPRS and hRRS were previously described (Braun et al., 2009). The previously described affinity-based reference set (AIRS) was extended by 14 PPIs (Trepte et al., 2015), by selecting 18 PPIs from PDBbind (Wang et al., 2004) and 53 PPIs from the Protein-Protein Interaction Affinity Database 2.0 (Kastritis et al., 2011). The cDNAs encoding 71, 81, and 80 protein pairs in AIRS, hPRS, and hRRS were shuttled into LuTHy expression plasmids for systematic interaction testing.

**LuTHy experiments**

1) Reverse transfect HEK293 cells in white 96-well microtiter plates (Greiner, 655983) at a density of 4.0–4.5 × 10⁴ cells per well. Plasmids encoding donor and acceptor proteins are transfected at a 1:10 to 1:20 ratio, with 5–10 ng DNA for the donor and 100 ng for the acceptor. Primary hippocampal neurons are plated on poly-l-lysine coated white 96-well microtiter plates (Greiner, 655983) at a density of 1 × 10⁶ cells per well. Infect neurons with 5–35 μl of the viral solution (0.5–1 × 10⁶ IU/ml) per well 2–4 days post-plating and culture in NBA at 37°C and 5%, for a total of 13–15 days (days in vitro, DIV) before starting measurements.

2) Measure mCitrine fluorescence 48 h after transfection in intact cells (Ex/Em: 500 nm/530 nm).

3) Add coelenterazine-h (NanoLight, 301 or pik, 102182) to a final concentration of 5 μM. Incubate cells for an additional 15 min and measure the total, short-WL and long-WL luminescence. Here, fluorescence and luminescence were measured using the Infinite® microplate readers M200, M1000, or M1000Pro (Tecan) using the BLUE1 (370–480 nm) and the GREEN1 (520–570 nm) filters at 1,000 ms integration time.

4) After luminescence measurements in intact cells, the luminescence-based co-precipitation (LuC) is performed. Lyse cells in 50–100 μl HEPES-phospo-lysis buffer (50 mM HEPES, 150 mM NaCl, 10% glycerol, 1% NP-40, 0.5% deoxycholate, 20 mM NaF, 1.5 mM MgCl₂, 1 mM EDTA, 1 mM DTT, 1 U Benzonase, protease inhibitor cocktail (Roche, EDTA-free), 1 mM PMSF, 25 mM glycerol-2-phosphate, 1 mM sodium orthovanadate, 2 mM sodium pyrophosphate) for 30 min at 4°C.

5) Production of PA-mCit- and NL-tagged fusion proteins is monitored by measuring fluorescence (mCitN) and luciferase activity (NLN) in crude cell lysates in white, small-volume 384-well microtiter plates (Greiner, 784074). Add 5 μl coelenterazine-h to 5 μl of cell lysates to a final concentration of 10 μM and measure the luminescence activity as before in a microplate reader with 100 ms integration time.

6) Coat small-volume 384-well microtiter plates (Greiner, 784074) the day before use with sheep gamma globulin (Jackson ImmunoResearch, 013-000-002) in carbonate buffer (70 mM NaHCO₃, 30 mM Na₂CO₃, pH 9.6) for 3 h at room temperature. Block with 1% BSA in carbonate buffer before incubating with rabbit anti-sheep IgGs in carbonate buffer (Jackson ImmunoResearch, 313-005-003) overnight at 4°C. Equilibrate all wells with lysis buffer immediately before use. Do not store IgG-coated plates longer than 24 h.
7) Incubate 15 μl of cell lysate for 3 h at 4°C in the IgG-coated 384-well plates. Wash all wells three times with lysis buffer and measure the mCitrine fluorescence (mCh₉₅₇) as above. Finally, 15 μl of PBS buffer containing 10 μM coelenterazine-h is added to each well and the luminescence activity (N₉₅₇) is measured as described above.

**LuTHy donor saturation experiments**

1) For donor saturation experiments, plasmid DNAs (between 0.1 and 20.0 ng) encoding NL fusion proteins are co-transfected with increasing amounts of plasmid DNAs encoding the PA-mCit-tagged constructs.

2) BRET measurements are performed as described before, 72 h after transfection using NanoGlo® (Promega) at a final concentration of 1:500.

3) Calculate the fluorescence to luminescence ratio for the PA-mCit-NL tandem construct. In this fusion protein, the stoichiometric ratio of mCitrine and NL is 1:1. Next, the fluorescence to luminescence ratios for the studied interactions of interest can be estimated by normalizing the calculated values to the value obtained with the tandem construct.

**LuTHy data analysis**

The BRET and LuC ratios from *in vivo* BRET and *ex vivo* co-precipitation measurements are calculated as follows:

\[
BRET = \frac{LWL}{SWL} - Cf
\]

with LWL and SWL being the detected luminescence at the long (520–570 nm) and the short (370–480 nm) wavelengths, respectively. Cf is the correction factor for the donor bleed-through, which is the ratio of the luminescence measured at LWL and SWL in cells expressing the PA-NL construct and calculated as:

\[
Cf = \frac{LWL_{PA-NL}}{SWL_{PA-NL}}
\]

2) The corrected BRET (cBRET) ratio is calculated by subtracting either the BRET ratio of control 1 (NL/PA-mCit-Y) or of control 2 (NL-X/PA-mCit) from the BRET ratio of the studied interaction of interest (NL-X/PA-mCit-Y). The calculated LuC ratios obtained for the controls 1 and 2 are always compared with each other and the higher value is used to correct the LuC ratio of the interaction of interest (see Appendix Fig S1F).

3) For the LuC readout, the obtained luminescence precipitation ratio (PIR) of the control protein PA-NL (PIR₉₅₇) is used for data normalization, which is calculated as follows:

\[
PIR_{PA-NL} = \frac{N_{OUT}}{3 \times N_{IN}}
\]

with N₉₅₇ being the total luminescence measured after co-IP and N₉₅₇ the luminescence measured in cell extracts directly after lysis.

4) Subsequently, LuC ratios are calculated for all interactions of interest and normalized to the PIR₉₅₇ ratio:

\[
LuC = \frac{NL_{OUT}/3 \times NL_{IN}}{PIR_{PA-NL}}
\]

5) Finally, a corrected LuC (cLuC) ratio is calculated by subtracting either the LuC ratio of control 1 (NL/PA-mCit-Y) or of control 2 (NL-X/PA-mCit) from the LuC ratio of the studied interaction of interest (NL-X/PA-mCit-Y). The calculated LuC ratios obtained for the controls 1 and 2 are always compared with each other and the higher value used to correct the LuC ratio of the interaction of interest (see Appendix Fig S1G).

A receiver operating characteristics (ROC) curve was obtained for PPIs of the hPRS and hRRS using corrected cBRET and cLuC ratios. Binary interactions in all LuTHy screens (hPRS, hRRS, AIRS, UBX-domain-containing, synaptic, non-synaptic) were scored positive with cBRET and cLuC cutoffs ≥ 0.01 and ≥ 0.03, respectively. Interactions between two integral-membrane or membrane-associated proteins were scored positive with cBRET and cLuC cutoffs ≥ 0.03 and ≥ 0.05, respectively.

Gene ontology (GO) analysis was performed using the database for annotation, visualization, and integrated discovery (DAVID) (Huang et al, 2009), https://david.ncifcrf.gov.

**Investigating the effects of small molecules on PPIs with LuTHy**

For the investigation of the effects of small molecules on PPIs with LuTHy, HEK293 cells were co-transfected with plasmids encoding the proteins NL-MDM2 and PA-mCit-p53 or FL-FKBP12 and PA-mCit-FRB in white 96-well microtiter plates (Greiner, 655983). Increasing concentrations of Nutlin-3 (Sigma, N6287) or rapamycin (Santa Cruz, 3504A) were added to cells 24 h after transfection. Luminescence measurements in intact cells were performed after an additional incubation for 6 (Nutlin-3) or 4 h (rapamycin) upon addition of 5 μM coelenterazine-h. After the measurement of BRET in intact cells, the cells were lysed with HEPES-phospho-lysis buffer that contained the respective concentration of compounds. Finally, the compound effects on PPIs were assessed by LuC as described above.

For Hsp90 inhibition, cells were transfected in a 10-cm dish using FuGENE (Promega) according to manufacturer’s instructions. Geldanamycin (Selleckchem S2713) and ganetespib (Selleckchem S1159) were added to a 384-well plate 24 h after transfection and cells seeded on top at a density of 15,000 cells per well. BRET measurements were performed in a Tecan Spark reader using filter settings at 360–485 nm and 520–650 nm at the indicated time points by coelenterazine-h injection. For heat shock experiments, HEK293 cells were transferred 48 h after transfection to 42°C for 6 h followed by a recovery phase at 37°C for an additional 24 h. Control treatment was conducted with identically transfected cells, which were constantly kept at 37°C. BRET measurements at the depicted time points (Appendix Fig S10D) were performed as described above.

**DULIP assay**

The DULIP assay and data analysis were performed as described before (Trepte et al, 2015). Briefly, HEK293 cells were reversely transfected and lysed after 48 h. Production of PA-RL- and FL-tagged fusion proteins was monitored by measuring the respective luciferase activities in crude cell lysates. In parallel, cell lysates
were incubated for 3 h at 4°C in IgG-coated 384-well plates. Measurement of firefly and Renilla luminescence activities was performed using an Infinite® M1000 (Tecan) plate reader and the Dual-Glo® Stop & Glow® reagents (Promega). NIR and cNIR values were determined as described previously (Trepte et al., 2015).

**FRET assay**

FRET assays were performed as described before (Trepte et al., 2015). Briefly, HEK293 cells were reversibly transfected using 100 ng plasmid DNAs encoding PA-mCitrine- and mCherry-tagged fusion proteins in black 96-well microtiter plates at a density of 4.5 × 10^4 cells per well. 72 h after transfection, cells were fixed with 4% paraformaldehyde in PBS and washed twice with PBS. Fluorescence signals were detected with an Infinite® M1000Pro (Tecan) plate reader: donor channel [excitation (Ex)/emission (Em): 500/530 nm], acceptor channel (Ex/Em: 580/610 nm) and FRET channel (Ex/Em: 500/610 nm). In brief, FRET efficiencies (E_{app} in %) were calculated according to the sensitized emission formula as follows:

\[
E_{app} = \frac{(DA - cD \times DD - cA \times AA)}{AA}
\]

with \( DD \) = donor channel signal; \( AA \) = acceptor channel signal; \( DA \) = FRET channel signal; \( cD \) = donor bleed-through using donor only sample (pPA-mCit); \( cA \) = acceptor cross-excitation using acceptor only sample (pmCherry).

**Western blotting**

For Western blotting experiments, cells were lysed for 30 min at 4°C in HEPES-phospho-lysis buffer and the protein concentrations determined using the Pierce™ BCA assay (Thermo Scientific). Then, the same amounts of protein lysates were loaded onto NuPAGE™ Novex™ 4–12% Bis-Tris precast polyacrylamide gels (Thermo Fisher) according to the manufacturer’s instructions. Following, proteins were transferred onto a nitrocellulose membrane (GE, 10401197) using a BioRad™ wet blotting system. The membrane was blocked in 3%-milk PBS-T (phosphate-buffered saline, 0.05% Tween) for 60 min at room temperature and incubated with primary antibody in 3%-milk PBS-T overnight at 4°C. Bound antibodies were detected with an appropriate HRP (horse-radish-peroxidase)-coupled secondary anti-rabbit (1:2,000, Sigma A0545) and anti-mouse (1:2,000, Sigma A0168) antibody by measuring chemiluminescence in a Fujifilm LAS-3000 after the addition of WesternBright Quantum (Advansta, 12042-D20). The primary antibodies used were as follows: anti-NanoLuc (rabbit, 1:5,000, kindly provided by Promega), anti-GFP (rabbit, 1:5,000, abcam ab290), anti-CSP (rabbit, 1:1,000, Synaptic Systems, 154 003), anti-Histone H3 (rabbit, 1:5,000, abcam ab1791), anti-VCP (mouse, 1:1,000, Progen, 65278), anti-UBXD9 (rabbit, 1:1,000, LifeSpan Biosciences, LS-C156546, EPR8616), and anti-tubulin (mouse, 1:4,000, Sigma, T6074).

**Bioluminescence imaging (BLI)**

HEK293 cells were seeded at a low density with 3 × 10^4 cells per well in an ibidi® μ-Slide 8-well tissue culture dish and transfected using X-tremeGENE (Roche®) according to the manufacturer’s instructions. Forty-eight hours after transfection, imaging was performed using an Olympus® inverted microscope IX83 with an Andor Zyla camera and a 60x objective. A Dual-View™ adapter (Optical Insights®) between microscope und camera was equipped with a short (460/50, Chroma®) and long (535/50, Chroma®) wavelength filter. Imaging was performed in tyrode’s buffer (100 mM NaCl, 5 mM KCl, 2 mM HEPES, 2 mM CaCl2, 4 mM MgCl2, 33 mM glucose) containing 1:50 NanoGlo® (Promega) substrate. Before substrate addition, mCitrine was excited with a LED excitation system (CoolLED® pE, 2%) using a YFP-excitation filter (Olympus®, AHP F48-003 ET-Set); the emitted fluorescence was detected with an exposure time of 100 ms. Bioluminescence imaging started immediately after substrate addition with an exposure time of 3,000 ms. Images were processed with ImageJ (http://imagej.nih.gov/ij/). Fluorescence images were background-corrected with the contrast adjusted for each image individually as the fluorescence signal was only used to study the localization of mCit-tagged proteins. The bioluminescence images, which contain the short (460 BP) and long wavelength (530 BP) signals, were split into two separate images. These two images were stacked and aligned using the StackReg plugin in ImageJ (Thévenaz, 1998). Next, the images were smoothed by median filtering, background-subtracted, and thresholded on luminescence intensities. To calculate the BRET ratio on a pixel-by-pixel basis, the 530 BP image was divided by the 460 BP image using the image calculator and presented in pseudo-color for better visualization.

**Proximity ligation assay**

All animals were sacrificed according to the permit given by the Office for Health Protection and Technical Safety of the regional government of Berlin (LaGeSo, X9009/14) and in compliance with regulations laid down in the European Community Council Directive. One day before preparation of primary neurons, all coverslips containing culture plates were coated with poly- DL ornithine hydrobromide (PO, 0.005% PO in H2O) over night at 37°C. On the preparation day, the coating solution was removed and plates were incubated with DMEM containing 10% FCS for several hours. The pregnant mice were sacrificed by cervical dislocation and the embryos were removed and placed in ice-cold standard salt solution (SSS, 135 mM NaCl, 5 mM KCl, 1.25 mM CaCl2, 0.5 mM MgCl2, 0.5 mM MgSO4, 1 mM NaHCO3, 10 mM HEPES, 25 mM glucose, 1× AB/AM). Brains of all embryos were isolated and collected in SSS on ice. Next, the meninges were removed and hippocampi of all embryos were isolated. The tissue was cut in smaller pieces and cultured in basal medium (neurobasal medium, 25 mM glucose) containing 1:100 NanoGlo® reagents (Promega). NIR and cNIR imaging was performed using a Dual-View™ adapter (Optical Insights®) between microscope und camera was equipped with a short (460/50, Chroma®) and long (535/50, Chroma®) wavelength filter. Imaging was performed in tyrode’s buffer (100 mM NaCl, 5 mM KCl, 2 mM HEPES, 2 mM CaCl2, 4 mM MgCl2, 33 mM glucose) containing 1:50 NanoGlo® (Promega) substrate. Before substrate addition, mCitrine was excited with a LED excitation system (CoolLED® pE, 2%) using a YFP-excitation filter (Olympus®, AHP F48-003 ET-Set); the emitted fluorescence was detected with an exposure time of 100 ms. Bioluminescence imaging started immediately after substrate addition with an exposure time of 3,000 ms. Images were processed with ImageJ (http://imagej.nih.gov/ij/). Fluorescence images were background-corrected with the contrast adjusted for each image individually as the fluorescence signal was only used to study the localization of mCit-tagged proteins. The bioluminescence images, which contain the short (460 BP) and long wavelength (530 BP) signals, were split into two separate images. These two images were stacked and aligned using the StackReg plugin in ImageJ (Thévenaz, 1998). Next, the images were smoothed by median filtering, background-subtracted, and thresholded on luminescence intensities. To calculate the BRET ratio on a pixel-by-pixel basis, the 530 BP image was divided by the 460 BP image using the image calculator and presented in pseudo-color for better visualization.
6.9, 1 mM EGTA, 1 mM MgCl₂, 90% MeOH). Following, cells were blocked in 0.1% saponin (Sigma-Aldrich), 1% BSA (VWR), 4% goat serum (Dako), 50 ng/µl RNase A (Promega), 11 ng/µl poly (A) (P9403 Sigma-Aldrich), and 2.5 ng/µl salmon sperm DNA (Invitrogen) in PBS for 30 min at room temperature. Proximity ligation assays (PLAs) were performed according to manufacturer’s protocol (Duolink® In Situ Detection Reagents Orange, Sigma-Aldrich). Primary antibodies were prepared in antibody diluent PBS containing (0.1% saponin, 3% BSA, and 1:1000 salmon sperm DNA) and incubated with slides (50 µl/sample) for 1 h at 37°C in a humidity chamber. Unbound primary antibodies were removed by washing the slides 3× in PBS with 0.1% saponin. Thereafter, both PLA probes (Duolink® Anti-Rabbit PLUS and Anti-Mouse MINUS) were diluted 1:5 in antibody diluent and allowed to sit for 20 min before being added to cover slips (40 µl/sample) for 1 h at 37°C in a humidity chamber. Slides were washed for 2 × 5 min in wash buffer A (0.01 M Tris, 0.15 M NaCl, 0.05% Tween) under gentle agitation to remove unbound PLA probes. For the ligation, cover slips were incubated with Duolink® ligation reagent (Ligation stock 1:5 in antibody diluent plus ligase, 1:40; 40 µl/sample) for 30 min at 37°C in a humidity chamber. Washing was performed 2 × 2 min with wash buffer A under gentle agitation. For signal amplification, Duolink® Amplification stock was diluted 1:5 in high-purity water and polymerase (1:80) was added before slides were incubated (40 µl/sample) for 100 min at 37°C in a humidity chamber. Final washing steps were performed with 2 × 10 min 1× wash buffer B (0.2 M Tris, 0.1 M NaCl) and 1 × 1 min 0.01× wash buffer B. Coverslips were mounted in fluorescent mount medium (Dako) and images acquired on a Leica TCS SP5 confocal microscope in the Advanced Light Microscopy (ALM) technology platform at the Max Delbrück Center for Molecular Medicine. Images were processed with ImageJ.

**Immunofluorescence imaging**

SHEP cells (Muth et al, 2010) were seeded at low density (3 × 10⁵ cells) onto fibronectin (10 µg/ml)-coated coverslips in a 6-well plate and transfected using GeneJet™ (SignaGen, SL100488) according to the manufacturer’s instructions. Forty-eight hours after transfection, the cells were fixed with 4% paraformaldehyde in PBS with 1:5,000 Hoechst (ThermoFisher, 33342). Cells were permeabilized with 0.1% saponin in PBS for 10 min and blocked with 3% BSA, 0.1% saponin in PBS for 30 min at RT. Primary anti-NanoLuc (rabbit, 1:1,000, Promega) and secondary Cy5-coupled anti-rabbit (1:500) antibodies were diluted in 3% BSA, 0.1% saponin in PBS and applied successively to the coverslips for 1 h at 37°C with three washing steps in between. Coverslips were mounted in fluorescent mount medium (Dako) and images acquired on a Leica TCS SP5 confocal microscope in the Advanced Light Microscopy (ALM) technology platform at the Max Delbrück Center for Molecular Medicine. Images were processed with ImageJ.

**Data availability**

The protein interactions from this publication have been submitted to the IMEx (http://www.imexconsortium.org) consortium through IntAct (Orchard et al, 2014) and assigned the identifier IM-26441.

**Expanded View** for this article is available online.

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**Author contributions**

PT and EEW conceived the study, and PT designed all experiments and analyzed the data. SKr, AT, SKo, MZ, KR, and SG generated the expression vectors, and SKr performed most LuTHy experiments with technical assistance by MZ, AT, KR, and SG. Donor saturation experiments were performed by SKr, immunoblotting by SKr and SKo and the compound studies by SKr and SG. Heat shock studies with HSF1 were designed and performed by SKo and PT. PLA was established by LD and performed by SG. Immunofluorescence imaging was performed by AB, AT, and CS. The FRET experiment was performed by SKo. Under supervision by CCG, SH performed bioluminescence imaging studies and LuTHy experiments in hippocampal neurons. Primary neuronal cultures were prepared by AS for PLA experiments and by SH for LuTHy assays. PT, AB, and EEW conceptually designed the assay. SKr, AB, SKo, AT, and KK assisted in the design, analysis, and interpretation of experiments. All authors discussed the results and EEW, SS, and PT wrote the manuscript with comments from all authors.

**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**

Arumughan A, Roske Y, Barth C, Forero LL, Bravo-Rodriquez K, Redel A, Kostova S, McShane E, Opitz R, Faelber K, Rau K, Mielke T, Daumke O, Selbach M, Sanchez-Garcia E, Rocks O, Panáková D, Heinemann U, Wanker EE (2016) Quantitative interaction mapping reveals an extended UBX domain in ASPL that disrupts functional p97 hexamers. Nat Commun 7: 13047

Bandopadhyay S, Chiang C-Y, Sivestava J, Gersten M, White S, Bell R, Kurschner C, Martin CH, Smoot M, Sahasrabudhe S, Barber DL, Chanda SK, Ideker T (2010) A human MAP kinase interactome. Nat Methods 7: 801 – 805
Barrios-Rodiles M, Brown KR, Ozdaman B, Bose R, Liu Z, Donovan RS, Shinjo F, Liu Y, Dembowy J, Taylor IW, Luga V, Przulj N, Robinson M, Suzuki H, Hayashizaki Y, Jurisica I, Wrana JL (2005) High-throughput mapping of a dynamic signaling network in mammalian cells. Science 307: 1621 – 1625

Braun P, Tasan M, Drezè M, Barrios-Rodiles M, Lemmens I, Yu H, Sahalie JM, Murray RR, Roncari L, de Smet A-S, Venkatesan K, Rual J-F, Vandenhaute J, Cusick ME, Pawson T, Hill DE, Tavernier J, Wrana JL, Roth FP, Vidal M (2009) An experimentally derived confidence score for binary protein-protein interactions. Nat Methods 6: 91 – 97

Buntru A, Trepte P, Klockmeier K, Schnoegel S, Wanker EE (2016) Current approaches toward quantitative mapping of the interactome. Front Genet 7: 74

Burgoyne RD, Morgan A (2015) Cysteine string protein (CSP) and its role in preventing neurodegeneration. Semin Cell Dev Biol 40: 153 – 159

Bushell KM, Sillén C, Schuster-Boeckler B, Bateman A, Wright G (2008) Large-scale screening for novel low-affinity extracellular protein interactions. Genome Res 18: 622 – 630

Cafarelli TM, Desbuleux A, Wang Y, Choi SG, De Ridder D, Vidal M (2017) Mapping, modeling, and characterization of protein–protein interactions on a proteomic scale. Curr Opin Struct Biol 44: 201 – 210

Chen Y-C, Rajagopala SV, Stellberger T, Uetz P (2010) Exhaustive benchmarking of the yeast two-hybrid system. Nat Methods 7: 667 – 668. Author reply 668

Couzens AL, Knight JR, Kean MJ, Teo G, Weiss A, Dunham WH, Lin Z-Y, Bagshaw RD, Sicheri F, Pawson T, Wrana JL, Choi H, Gingras A-C (2013) Protein interaction network of the mammalian hippo pathway reveals mechanisms of kinase-phosphatase interactions. Sci Signal 6: rs15

Dacres H, Michie M, Wang J, Pfleger KDC, Trowell SC (2012) Effect of enhanced Renilla luciferase and fluorescent protein variants on the Förster distance of Bioluminescence resonance energy transfer (BRET). Biochem Biophys Res Comm 425: 625 – 629

De Las Rivas J, Fontanillo C (2010) Protein–protein interactions essentials: key concepts to building and analyzing interactome networks. PLoS Comput Biol 6: e1000807

Donnelier J, Braun JEA (2014) CSPα-chaperoning presynaptic proteins. Front Cell Neurosci 8: 116

Drinovec L, Kubale V, Nøhr Larsen J, Vrecl M (2017) Mathematical models for quantitative assessment of bioluminescence resonance energy transfer: application to seven transmembrane receptors oligomerization. Front Endocrinol 3: 104

Gil RS, Ooi I, Yerbury JJ, Ecroyd H (2017) The heat shock response in neurons and astrogliosis and its role in neurodegenerative diseases. Mol Neurodegener 12: 65

Greaves J, Lemenidis K, Corleku OA, Cruchaga C, Grefen C, Chamberlain LH (2012) Palmitoylation-induced aggregation of cysteine-string protein mutants that cause neuronal ceroid lipofuscinosis. J Biol Chem 287: 37330 – 37339

Grünberg R, Burnier JF, Ferrar T, Beltran-Sastre V, Stricher F, Van Der Sloop AM, Garcia-Olivas R, Mallabiaurrena A, Sanjuan X, Zimmermann T, Serrano L (2013) Engineering of weak helper interactions for high-efficiency FRET probes. Nat Methods 10: 1021 – 1027

Han GA, Malintan NT, Saw NMM, Li H, Han L, Meunier FA, Collins BM, Sugita S (2011) Munc18-1 domain-1 controls vesicle docking and secretion by interacting with syntaxin-1 and chaperoning it to the plasma membrane. Mol Biol Cell 22: 4134 – 4149

Huang DW, Sherman BT, Lempicki RA (2009) Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. Nat Protoc 4: 44 – 57

Huttlin EL, Bruckner RJ, Paula JR, Cannon JR, Ting L, Baltier K, Colby G, Gebreab F, Gygi MP, Parzen H, Szpyt J, Tam S, Zarraga C, Pontano-Valetes L, Swarup S, White AE, Schwepe DK, Rad R, Erickson BK, Obar RA et al (2017) Architecture of the human interactome defines protein communities and disease networks. Nature 150: 1068

Jain A, Liu R, Ramani B, Arauz E, Ishitsuka Y, Raganathan K, Park J, Chen J, Xiang YK, Ha T (2011) Probing cellular protein complexes using single-molecule pulse-down. Nature 473: 484 – 488

Johnson RJ, McCoy JG, Bingman CA, Phillips GN Jr, Raines RT (2007) Inhibition of human pancreatic ribonuclease by the human ribonuclease inhibitor protein. J Mol Biol 368: 434 – 449

Josephson SA, Schmidt RE, Millsap P, McManus DQ, Morris JC (2001) Autosomal dominant Kufs’ disease: a cause of early onset dementia. J Neurol Sci 188: 51 – 60

Kastritis PL, Moal IH, Hwang H, Weng Z, Bates PA, Bonvin AMJJ, Janin J (2011) A structure-based benchmark for protein-protein binding affinity. Protein Sci 20: 482 – 491

Kim YE, Hosp F, Frottin F, Ge H, Mann M, Hayer-Hartl M, Hartl FU (2016) Soluble oligomers of PolyQ-expanded huntingtin target a multiplicity of key cellular factors. Mol Cell 63: 951 – 964

Lavioe H, Li JJ, Padgeganah A, Guiral S, Duchaine J, Mao DYL, Bouvier M (2013) Inhibitors that stabilize a closed RAF kinase domain conformation induce dimerization. Nat Chem Biol 9: 428 – 436

Lievens S, Gerlo S, Lemmens I, De Clercq DJH, Risseeuw MDP, Vanderroost N, de Smet A-S, Ryusinskii E, Chevet E, Van Calenbergh S, Tavernier J (2014) Kinase Substrate Sensor (KISS), a mammalian in situ protein interaction sensor. Mol Cell Proteomics 13: 3332 – 3342

Lievens S, Van der Heyden J, Masscheele D, De Ceuninck L, Petta I, Gupta S, De Puysseleyr V, Vauthier V, Lemmens I, De Clercq DJH, Defever D, Vanderroost N, de Smet A-S, Eckerman S, Van Calenbergh S, Martens L, De Bosscher K, Libert C, Hill DE, Vidal M et al (2016) Proteome-scale binary interactomics in human cells. Mol Cell Proteomics 15: 3624 – 3639

Lin T-Y, Bear M, Du Z, Foley KP, Ying W, Barsoum J, London C (2008) The novel HSP90 inhibitor STA-9090 exhibits activity against Kit-dependent and -independent malignant mast cell tumors. Exp Hematol 36: 1266 – 1277

Lois C, Hong EJ, Pease S, Brown EJ, Baltimore D (2002) Germline transcription and tissue-specific expression of transgenes delivered by lentiviral vectors. Science 295: 868 – 872

Lopez-Ortega E, Ruiz R, Tabares L (2017) CSPα, a molecular co-chaperone essential for short and long-term synaptic maintenance. Front Neurosci 11: 1323

Macleidt T, Woodroofe CC, Schwinn MK, Méndez J, Robers MB, Zimmerman K, Otto P, Daniels DL, Kirkland TA, Wood KV (2015) NanoBRET—a novel BRET platform for the analysis of protein–protein interactions. ACS Chem Biol 10: 1797 – 1804

Markson G, Kiel C, Hyde R, Brown S, Charalabous P, Bremm A, Semple J, Woodsmith D, Juley S, Salehi-Ashtiani K, Vidal M, Komander D, Serrano L, Lehner P, Sanderson CM (2009) Analysis of the human E2 ubiquitin conjugating enzyme protein interaction network. Genome Res 19: 1905 – 1911

Mezzanotte L, van’t Rooth M, Karatas H, Goun EA, Löwik CWGM (2017) In vivo molecular bioluminescence imaging: new tools and applications. Trends Biotechnol 35: 640 – 652

Morimoto RI (1993) Cells in stress: transcriptional activation of heat shock genes. Science 259: 1409 – 1410

Muth D, Ghazaryan S, Eckerle I, Beckett E, Pöhler C, Bätzler J, Beisel C, Gogolin S, Fischer M, Henrich K-O, Ehemann V, Gillespie P, Schwab M,
Westermann F (2010) Transcriptional repression of SKP2 is impaired in MYCN-amplified neuroblastoma. Cancer Res 70: 3791 – 3802

Nosklov L, Stránecký V, Hartmannová H, Pístoupilová A, Baresová V, Ivánk R, Hulcová H, Jahnová H, van der Zee J, Staropoli JF, Sims KB, Tynjälä J, Van Broeckhoven C, Nijsen PCG, Male SE, Elf er M, Kmoch S (2011) Mutations in DNAJC5, encoding cysteine-string protein alpha, cause autosomal-dominant adult-onset neuronal ceroid lipofuscinosis. Am J Hum Genet 89: 241 – 252

Orchard S, Ammar M, Aranda B, Breuza L, Briganti L, Broackes-Carter F, Campbell NH, Chavali G, Chen C, del-Toro N, Duesbury M, Dumousseau M, Galeota E, Hinz U, Iannuccelli M, Jagannathan S, Jimenez R, Khadake J, Lagreid A, Licata I et al (2014) The MintAct project – IntAct as a common curation platform for 11 molecular interaction databases. Nucleic Acids Res 42: D358 – D363

Pfleger KDG, Eidne KA (2006) Illuminating insights into protein-protein interactions using bioluminescence resonance energy transfer (BRET). Nat Methods 3: 165 – 174

Pfleger KDG, Seeber RM, Eidne KA (2006) Bioluminescence resonance energy transfer (BRET) for the real-time detection of protein-protein interactions. Nat Protoc 1: 337 – 345

Rabindran SK, Haroun RI, Clos J, Wisniewski J, Wu C (1993) Regulation of heat shock factor trimer formation: role of a conserved leucine zipper. Science 259: 230 – 234

Raman M, Sergeev M, Garnaas M, Lydeard JR, Huttlin EL, Goessling W, Shah JV, Harper JW (2015) Systematic proteomes of the VCP-UBXN adaptor network identifies a role for UBXN10 in regulating ciliogenesis. Nat Cell Biol 17: 1356 – 1369

Rizo J, Südhof TC (2012) The membrane fusion enigma: SNAREs, Sec1/Munc18 proteins, and their accomplices–guilty as charged? Annu Rev Cell Dev Biol 28: 279 – 308

Rolland T, Anouti M, Charlieux B, Pevzner SJ, Zholq Q, Sahni N, Ys S, Lemmens I, Fontantillo C, Mosca R, Kamburov A, Chissian SD, Yang X, Ghamsari L, Balcha D, Begg BE, Brain P, Brehme M, Broly MP, Carvunis AR et al (2014) A proteome-scale map of the human interactome network. Cell 159: 1212 – 1226

Sahni N, Yi S, Taipele M, Fuxman Bass JI, Coulombe-Huntington J, Yang F, Peng J, Weile J, Karras GI, Wang Y, Kovácés IA, Kamburov A, Krykbaeva I, Lam MH, Tucker L, Kherana V, Sharma A, Liu Y, Yachie N, Zhong Q et al (2015) Widespread macromolecular interaction perturbations in human genetic disorders. Cell 161: 647 – 660

Scott DE, Bayly AR, Abell C, Skidmore J (2016) Small molecules, big targets: drug discovery faces the protein-protein interaction challenge. Nat Rev Drug Discov 15: 533 – 550

Sharma M, Burré J, Südhof TC (2011) CSPα promotes SNARE-complex assembly by chaperoning SNAP-25 during synaptic activity. Nat Cell Biol 13: 30 – 39

Sharma M, Burrré J, Bronk P, Zhang Y, Xu W, Südhof TC (2012) CSPα knockout causes neurodegeneration by impairing SNAP-25 function. EMBO J 31: 829 – 841

Stebbins CE, Russo AA, Schneider C, Rosen N, Hartl FU, Pavletich NP (1997) Crystal structure of an Hsp90-geldanamycin complex: targeting of a protein chaperone by an antitumor agent. Cell 89: 239 – 250

Stellberger T, Häuser R, Baiker A, Pothineni VR, Haas J, Uetz P (2010) Improving the yeast two-hybrid system with permuted fusions proteins: the Varicella Zoster Virus interactome. Proteome Sci 8: 8

Stelzl U, Worm U, Lalowski M, Haenig C, Brembeck FH, Goehler H, Strodzieck M, Zanker M, Schoenherr A, Koeppen S, Timm J, Mintzaffl S, Abraham C, Bock N, Kietzmann S, Goedde A, Tokszé L, Drooge A, Krobitsch S, Kom B et al (2005) A human protein-protein interaction network: a resource for annotating the proteome. Cell 122: 957 – 968

Südhof TC, Rizo J (2011) Synaptic vesicle exocytosis. Cold Spring Harb Symp Quant Biol 86: a005637

Taipale M, Krykbaeva I, Koeva M, Kayatekin C, Westover KD, Karras GI, Lindquist S (2012) Quantitative analysis of HSP90-client interactions reveals principles of substrate recognition. Cell 150: 987 – 1001

Taipele M, Tucker G, Peng J, Krykbaeva I, Lin Z-Y, Larsen B, Choi H, Berger B, Gingers A-C, Lindquist S (2014) A quantitative chaperone interaction network reveals the architecture of cellular protein homeostasis pathways. Cell 158: 434 – 448

Thévenaz P (1998) A pyramid approach to subpixel registration based on intensity. IEEE Trans Image Process 7: 27 – 41

Thomas GM, Hayashi T, Huganir RL, Linden DJ (2013) DHHC8-dependent PICK1 palmitoylation is required for induction of cerebellar long-term synaptic depression. J Neurosci 33: 15401 – 15407

Trepte P, Buntru A, Klockeier K, Willmore L, Arumughaan A, Secker C, Zanker M, Brusendorf L, Rauh A, Redel A, Wanker EE (2015) DULIP: a dual luminescence-based co-immunoprecipitation assay for interactome mapping in mammalian cells. J Mol Biol 427: 3375 – 3388

Venketesan K, Rual J-F, Vazquez A, Stelzl U, Lemmens I, Hirozane-Kishikawa T, Hao T, Zanker M, Xin X, Goh K-L, Yildirim MA, Simonis N, Heinzmann K, Gebreab F, Sahalie JM, Cevik S, Simon D, de Smet A-S, Dann E, Smolyar A et al (2009) An empirical framework for binary interactome mapping. Nat Methods 6: 83 – 90

Vogel SS, Thaler C, Koushik SV (2006) Fanciful FRET. Sci STKE: re2

Wang X, Wei X, Thijssen B, Das J, Lipkin SM, Yu H (2012) Three-dimensional reconstruction of protein networks provides insight into human genetic disease. Nat Biotechnol 30: 159 – 164

Zhang Y-Q, Henderson MX, Colangelo CM, Ginsberg SD, Bruce C, Wu T, Chandra SS (2012) Identification of CSPα clients reveals a role in dynamin 1 regulation. Neuron 74: 136 – 150

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