Inducible biomolecular condensates play fundamental roles in cellular responses to intracellular and environmental cues. Knowledge about their composition is crucial to understand the functions that arise specifically from the assembly of condensates. This protocol combines an optogenetic and an efficient proximity labeling approach to analyze protein modifications driven by protein condensation in cultured cells. Low endogenous biotin level ensures sharp signals.
Protocol
An optogenetic proximity labeling approach to probe the composition of inducible biomolecular condensates in cultured cells

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SUMMARY
Inducible biomolecular condensates play fundamental roles in cellular responses to intracellular and environmental cues. Knowledge about their composition is crucial to understand the functions that arise specifically from the assembly of condensates. This protocol combines an optogenetic and an efficient proximity labeling approach to analyze protein modifications driven by protein condensation in cultured cells. Low endogenous biotin level ensures sharp signals. For complete details on the use and execution of this protocol, please refer to Frattini et al. (2021).

BEFORE YOU BEGIN
Sub-cellular compartments are essential for the organization of cell physiological processes. Compartments that are not surrounded by a membrane, called biomolecular condensates, can form via the self-assembly of multivalent protein scaffolds and nucleic acids (Banani et al., 2017; Hyman and Simons, 2012; Lyon et al., 2020; Shin and Brangwynne, 2017). An analysis of the composition of biomolecular condensates is necessary to understand their associated functions. Condensates, however, are labile and intrinsically difficult to isolate. If a major protein component of the condensate is known, then proximity labeling approaches can be used to identify the other components of the condensate (Branon et al., 2018; Lam et al., 2015; Roux et al., 2012; Youn et al., 2018). Many biomolecular condensates, however, form only transiently in response to environmental changes. Furthermore, methods are required to understand the functions that arise specifically from the assembly of biomolecular condensates. To tackle this problem, we present here a method that combines proximity labeling with optogenetics to control the formation of a biomolecular condensate in live cells (Bracha et al., 2019; Bracha et al., 2018). A protein scaffold with intrinsic capacity to undergo liquid phase separation is fused to a photoreceptor that oligomerizes upon exposure to a specific light wavelength. Light-induced oligomerisation of the photoreceptor facilitates the phase separation of the protein scaffold that forms a condensate, within minutes. In comparisons with analogous chemogenetics approaches, optogenetics offers a better timing resolution. In association with proximity labeling, the actuation of condensates enables to probe the functional consequences of protein condensation and to reveal the composition of the resulting compartment. In the method described below, a protein scaffold is fused to both cryptochrome 2 (Palayam et al., 2021), a light responsive photoreceptor from Arabidopsis thaliana and to TurboID (Branon et al., 2018), a highly efficient biotin ligase. We used this method to show that the activity of the master checkpoint kinase ATR is amplified within nuclear condensates formed by the multivalent protein scaffold TopBP1 (Frattini et al., 2021). The method is applicable to multi-modular protein scaffolds that have intrinsic capacity to self-assemble and phase separate to yield biomolecular condensates.
**Note**: Optogenetic-induction of biomolecular condensates on a cell-population scale requires an illumination device adapted to cell culture formats. Resources for the construction of optogenetics light boxes are available at [https://www.optobase.org/materials/](https://www.optobase.org/materials/).

### Key Resources Table

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies** | | |
| pATR (T1989) | GeneTex | Cat# GTX128145; RRID: AB_2687562 |
| Chk1 | Santa Cruz Biotechnology | Cat# sc-8408; RRID: AB_627257 |
| pChk1 (Ser345) | Cell Signaling Technology | Cat# 2348; RRID: AB_331212 |
| TopBP1 | Bethyl | Cat# A300-111A; RRID: AB_2272050 |
| pTopBP1 (Ser1138) | Interchim | Cat# orb140434 |
| Alexa Fluor 546 goat anti-mouse IgG | Molecular Probes | Cat# A-11030, RRID: AB_144695 |
| Alexa Fluor 488 goat anti-rabbit IgG | Molecular Probes | Cat# A-11010, RRID: AB_2534077 |
| Alexa Fluor 488 goat anti-mouse IgG2b | Molecular Probes | Cat# A-21141, RRID: AB_141626 |
| Anti-mouse IgG, HRP linked Antibody | Cell Signaling Technology | Cat# 7076, RRID: AB_330924 |
| Anti-rabbit IgG, HRP linked Antibody | Cell Signaling Technology | Cat# 7074; RRID: AB_2099233 |
| ATR | Bethyl/Euromedex | Cat# A300-137A; RRID: AB_185544 |
| FancJ/BRIP1 | Novus | Cat# NB100-416; RRID: AB_2066307 |
| BRCA1(C-20) | Santa Cruz Biotechnology | Cat# sc-642; RRID: AB_630944 |
| **Bacterial and virus strains** | | |
| S-alpha Competent E. coli (High Efficiency) | NEB | Cat# C2987 |
| **Chemicals, peptides, and recombinant proteins** | | |
| Biotin | Sigma-Aldrich | Cat# B4501; CAS: 58-85-5 |
| Doxycline | Clontech | Cat# 631311; CAS: 10592-13-9 |
| Blasticidin | InvivoGen | Cat# anti-bl |
| Hygromycin | Sigma-Aldrich | Cat# H3274; CAS: 31282-04-9 |
| Zeocin | ThermoFisher Scientific | Cat# R25001 |
| Penicillin Streptomycin | Sigma-Aldrich | Cat# P0781; ID 329820056 |
| Ampicillin | Sigma-Aldrich | Cat# A9518; CAS: 69-52-3 |
| cOmplete, EDTA free | Roche | Cat# 4693159001 |
| Halt phosphatase inhibitor cocktail | ThermoFisher Scientific | Cat# 78427 |
| Benzonase Nuclease | Sigma-Aldrich | Cat# E1014; CAS: 9025-65-4 |
| Ethidium bromide solution | Sigma-Aldrich | Cat# E1510; CAS: 1239-45-8 |
| Streptavidin-Agarose | Sigma-Aldrich | Cat# S1638; MDL: MCFD00082035 |
| Dulbecco’s Modified Eagle’s Medium - high glucose | Sigma-Aldrich | Cat# D5796 |
| BioWest - Fetal Bovine Serum | Eurobio Scientific | Cat# 51810 |
| Glycerol ≥95% | WR Chemicals | Cat# 24388-295; CAS: 56-81-5 |
| Bromophenol Blue | Ethylenediaminetetraacetic acid | Cat# B0126; CAS: 115-39-9 |
| Ethylenediaminetetraacetic acid | Ethylenediaminetetraacetic acid | Cat# EDS; CAS: 60-00-4 |
| HEPES | Sigma-Aldrich | Cat# H3375; CAS: 7365-45-9 |
| Sodium Chloride | WR Chemicals | Cat# 27810-295; CAS: 7647-14-5 |
| Ethylene glycol-bis(2-aminoethyl)-N,N,N',N'-tetraacetic acid | Sigma-Aldrich | Cat# E4378; CAS: 67-42-5 |
| Sodium deoxycholate ≥97% | Sigma-Aldrich | Cat# D6750; CAS: 302-95-4 |
| Triton X-100 | Sigma-Aldrich | Cat# T8787; CAS: 9002-93-1 |
| Tergitol Solution type NP-40 | Sigma-Aldrich | Cat# NP405; MDL: MFCDD01779855 |
| Sodium dodecyl sulfate 20% | Biosolve | Cat# 198123; CAS:151-21-3 |
| Tris base | Euromedex | Cat# 200923-A; CAS: 77-86-1 |
| Dulbecco’s Phosphate Buffered Saline | Sigma-Aldrich | Cat# D8537; MDL: MFCDD0131855 |
| Water | Sigma-Aldrich | Cat# W3500; CAS: 7732-18-5 |
| LiCl | Sigma-Aldrich | Cat# L9650; CAS: 7447-41-8 |
| **Critical commercial assays** | | |
| Clarity Western ECL Substrate | Bio-Rad | Cat# 170-5061 |
| Clarity Max Western ECL Substrate | Bio-Rad | Cat# 1705062 |

(Continued on next page)
Continued

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|----------------------|--------|------------|
| Criterion TGX stain free gel 7.5% | Bio-Rad | Cat# 5678024 |
| Criterion TGX Stain-Free Gel 4–15% | Bio-Rad | Cat# 5678084 |
| Criterion TGX Stain-Free Gel 10% | Bio-Rad | Cat# 5671034 |
| Mini-PROTEAN TGX Stain-Free Gels, 7.5% | Bio-Rad | Cat# 4568023 |
| Mini-PROTEAN TGX Stain-Free Gels, 4–15% | Bio-Rad | Cat# 4568083 |
| Mini-PROTEAN TGX Stain-Free Gels, 10% | Bio-Rad | Cat# 4568033 |
| Trans-Blot Turbo Transfer Pack 0.2μm Nitrocellulose Midi, 10 pack | Bio-Rad | Cat# 1704159 |
| Trans-Blot Turbo Transfer Pack 0.2μm Nitrocellulose Mini, 10 pack | Bio-Rad | Cat# 1704158 |
| Color Prestained Protein Standard, Broad Range | BioLabs | Cat# P7712S |
| Lipofectamine 2000 | ThermoFisher Scientific | Cat# 11668-019 |
| Quick Start™ Bradford 1X Dye Reagent | Bio-Rad | Cat# 500-0205 |
| Filter Syringe Clearline 30mm AC 0.2 μm | Dominique Dutscher | Cat# DSR146560 |
| TERUMO Syringe Without Needle SML | Dominique Dutscher | Cat# 050006D |
| TERUMO Syringe Without Needle 10ML | Dominique Dutscher | Cat# 050008 |
| Cell Spatula | TPP - Techno Plastic Products | Cat# 99010 |
| Amersham Hyperfilm ECL (8 x 10^8) | Dominique Dutscher | Cat# 28906839 |
| Raw data | This paper | https://doi.org/10.17632/zb35ntwpjx.1 |

**Experimental models: Cell lines**

| Oligonucleotides | Invitrogen | Cat# R78007, RRID: CVCL_U427 |
|------------------|------------|-----------------------------|
| TurboID Nter: catcaacgtgttttgac | Eurofins MWG | N/A |
| TurboID Cter: aggcctggctcatact | Eurofins MWG | N/A |
| CRY2 Cter: gcgcgcctcagtcacgcaggt | Eurofins MWG | N/A |
| CRY2 Nter and mCherry Cter: ggtcagatccaagagcttc | Eurofins MWG | N/A |
| mCherry Nter: agggtcgcccctcgcctt | Eurofins MWG | N/A |

**Recombinant DNA**

| Recombinant DNA | Addgene, Cat# 166504 |
|-----------------|-----------------------|
| pCDNA_FRT-TO_TurboID-mCherry-Cry2 | Frattini et al, 2021 |
| pCDNA_FRT-TO_TurboID-TopBP1WT-mCherry-Cry2 | Frattini et al, 2021 |
| pOG44 Flp-Recombinase Expression Vector | Thermo Fisher Scientific |
| | Cat# V600520 |

**Software and algorithms**

| Software and algorithms | OME Remote Objects software | https://www.openmicroscopy.org/ |
|-------------------------|-------------------------------|---------------------------------|
| Cell Profiler 2.2.0 | Cell image analysis software | https://cellprofiler.org/ |
| Image Lab™ Software (Version 5.2.1) | Bio-Rad | http://www.bio-rad.com/fr-fr/product/image-lab-software?ID=KRE6FSE8Z |
| Biorender Software | Science Suite Inc. | RRID: SCR_018361 |

**Other**

| Other | Binder | Cat# 9040-0078 |
|-------|--------|----------------|
| CO2 Incubator C150 | Binder | Cat# 9040-0078 |
| Sonicator, VibraCell- 72405 | BioBlock scientific | N/A |
| KNF LABOPORT Mini Diaphragm Vacuum Pump N 811 in Pumps, Compressors | Dominique Dutscher | Cat# KNF_28002 |
| Centrifuge Hettich Mikro 200 | Grosser | N/A |
| Mini-PROTEAN Tetra Vertical Electrophoresis Cell | Bio-Rad | Cat# 1658004 |
| PowerPac™ HC High-Current Power Supply | Bio-Rad | Cat# 1645052 |
| Trans-Blot Turbo Transfer System | Bio-Rad | Cat# 1704150 |

**MATERIALS AND EQUIPMENT**

**Equipment**

We use a custom-made illumination box containing an array of 24 Light Emitting Diode (LEDs) (488 nm) delivering 10 mW/cm² (light intensity measured using a ThorLabs-PM16-121-power meter). The 24 LEDs are made of Cree® XLamp® XR-E LED (Figure 2). The details on how to build the light box along with the code used for programming is described here: https://github.com/jvrana/OptogeneticsLightBox. Alternatively, a service for tailored made optogenetic devices is
Preparation of buffers

| RIPA/SDS Lysis buffer |  |  |  |
|-----------------------|-----------------|-----------------|-----------------|
| Stock concentration  | Final concentration | Amount |
| Tris-HCl pH 7.5  | 1 M | 50 mM | 2.5 mL |
| NaCl | 5 M | 150 mM | 1.5 mL |
| EDTA | 0.5 mM | 1 mM | 100 μL |
| EGTA | 0.5 mM | 1 mM | 100 μL |
| NP-40 | 100% | 1% | 500 μL |
| SDS | 20% | 0.1% | 250 μL |
| Sodium deoxycholate | n/a | 0.5% | 250 mg |
| ddH2O | n/a | n/a | Up to 50 mL |
| Total | n/a | n/a | 50 mL |

△ CRITICAL: For SDS usage, wear personal protective equipment/face protection.

Store the buffer at 4°C for three to six weeks.

| Wash Buffer 1 |  |  |  |
|----------------|-----------------|-----------------|-----------------|
| Stock concentration | Final concentration | Amount |
| SDS | 20% | 2% | 5 mL |
| ddH2O | n/a | n/a | 45 mL |
| Total | n/a | n/a | 50 mL |

△ CRITICAL: For SDS usage, wear personal protective equipment/face protection.

Store the buffer at 18°C–25°C.

| Wash Buffer 2 |  |  |  |
|----------------|-----------------|-----------------|-----------------|
| Stock concentration | Final concentration | Amount |
| HEPEs pH 7.5 | 0.5 M | 50 mM | 5 mL |
| NaCl | 5 M | 500 mM | 5 mL |
| EDTA | 0.5 mM | 1 mM | 100 μL |
| Triton X-100 | 20% | 1% | 2.5 mL |
| Sodium deoxycholate | n/a | 0.2% | 100 mg |
| ddH2O | n/a | n/a | Up to 50 mL |
| Total | n/a | n/a | 50 mL |

△ CRITICAL: Store the buffer at 4°C for three to six weeks.
The day of the experiment, add 50 μL of 2-mercaptoethanol (14.3 M) per 950 μL of 2X Laemmli Sample buffer (Total volume 1 mL).

To obtain 1X Laemmli Sample buffer, dilute 1 mL of 2X Laemmli Sample buffer with 1 mL distilled water.

⚠ CRITICAL: For SDS usage, wear personal protective equipment/face protection.

Store the buffer at 18°C–25°C for up to 2 years without adding 2-mercaptoethanol.

## Stock solutions

### Biotin Stock Solution

| Reagent   | Final concentration | Amount |
|-----------|---------------------|--------|
| Biotin    | 250 mM              | 610 mg |
| DMSO      | n/a                 | Up to 10 mL |
| **Total** | n/a                 | 10 mL  |

Note: Filter with 0.22 μm filter, prepare 100 μL aliquots of the stock solution and store at –20°C for up to 1 year. Avoid repeated freeze-thaw cycles.
Use the 10 mg/mL stock solution of doxycycline to further dilute 1:5000 to obtain 2 \( \mu \)g/mL working solution in the DMEM medium supplemented with 10% FBS.

**Note:** Filter with 0.22 \( \mu \)m filter, prepare 100 \( \mu \)L aliquots of the stock solution and store at -20°C for up to 1 year. Avoid repeated freeze-thaw cycles.

**STEP-BY-STEP METHOD DETAILS**

The first major step is to create a stable cell line for the inducible expression of a protein scaffold that forms a biomolecular condensate, tagged with TurboID and Cry2.

**Cloning procedure**

© Timing: 2 weeks

1. Introduce your gene of interest (GOI), which encodes a major phase separation component of a condensate, into the pCDNA5_FRT-TO_TurboID-mCherry-Cry2 (Opto-module). We have deposited this plasmid with Addgene (ID number 166504).

   Sub-clone the cDNA encoding your protein of interest into pCDNA5_FRT-TO_TurboID-mCherry-Cry2 using standard molecular biology procedures. You can use a variety of cloning strategies. For example, the cDNA can be sub-cloned in the KpnI site of the vector (Figure 1) using the In-Fusion HD cloning system (Takara Bio, USA). For a detailed protocol and instructions on the design of primers, follow the manufacturer’s guidelines (https://www.takarabio.com/documents/User%20Manual/In/In-Fusion%20HD%20Cloning%20Kit%20User%20Manual_102518.pdf).

2. Verify by DNA sequencing that the cDNA has been inserted in frame and that no mutations have been introduced in the construct.
   a. Design multiple primers set to cover the entire gene of interest.
   b. Use the primers indicated in the Key Resources for the sequencing of the TurboID, mCherry and CRY2 genes.

This construct will allow you to generate stable Flp-In\textsuperscript{TM}-Rex\textsuperscript{TM}293 cell lines to express your protein of interest fused to TurboID and mcherry-CRY2, in an inducible manner. Before the production of stable cell lines, we recommend that you transfect the construct transiently to verify the expression and the localization of your protein of interest, as described below.

**Experimental validation of your construct**

3. Thaw the Flp-In\textsuperscript{TM}-Rex\textsuperscript{TM}293 cell line several days before the transfection day and maintain the cell line in medium containing Zeocin selective reagent (100 \( \mu \)g/mL) and blasticidin (15 \( \mu \)g/mL).
Refer to the manufacturer’s instructions for culturing and handling the Flp-In/C228 T-REx/C228 293 cell line: (https://assets.thermofisher.com/TFS-Assets/LSG/manuals/flpinotrexcells_man.pdf).

4. Transfect Flp-In/C228 T-REx/C228 293 using Lipofectamine 2000 Reagent as described in the manufacturer’s instructions (https://assets.thermofisher.com/TFS-Assets/LSG/manuals/Lipofectamine_2000_Reag_protocol.pdf).

5. Verify by Western blotting that your protein of interest is expressed using either an anti-mCherry antibody or an antibody directed against your protein of interest.

6. Using standard fluorescence microscopy, verify the intracellular localization of your protein of interest.

Generation of stable cell lines expressing your protein of interest

© Timing: 6–8 weeks

7. Generate stable Flp-In™-TREx™293 expression cell lines by co-transfection of the pCDNAS_FRT-TO_TurboID-GOI-mCherry-Cry2 expression construct and the pOG44 recombinase plasmid (Figure 1).

Note: We used Flp-In™-TREx™293 cell lines but it is also possible to use Flp-In™-TREx™ U2OS/HeLa cell lines. We noticed however that the expression level of recombinant proteins was heterogeneous in U2OS/HeLa Flp-In™-TREx™ cells.

Note: Flp-In™-TREx™293 are generally amenable to transfection using standard methods. However, we recommend the transfection by Lipofectamine™ 2000 Reagent. Refer to the manufacturer’s instructions for the transfection procedure (https://assets.thermofisher.com/TFS-Assets/LSG/manuals/Lipofectamine_2000_Reag_protocol.pdf).

8. Select stable transfectants using 100–200 μg/mL hygromycin B.

9. After 6–8 weeks of selection, pool all the hygromycin resistant foci.

10. Grow the cells until 70–80% confluence, harvest and freeze multiple vials of early-passage cells.

Pause point: Place the cell vials in liquid nitrogen for long-term cryopreservation.
Experimental validation of the stable cell lines

11. Induce the expression of your gene of interest with doxycycline to a final concentration of 2 μg/mL.

   Note: Optimize the concentration of doxycycline for your specific application.

   Note: For immunofluorescence analysis, culture cells on coverslips.

12. Incubate the cells for 16 h to 24 h at 37°C.
13. The next day, place the plates in the illumination box in a CO2 incubator for optogenetic activation (Figure 2.2).

   Apply multiple light-dark cycles. For induction of TopBP1 condensation, we use cycles of 4 s light followed by 10 s dark for 3 min (Figure 2.3).

   Note: Induction of biomolecular condensates usually occurs within minutes. The efficacy of optogenetic induction of condensates depends on protein concentration and protein sequence, hence the time of exposure to blue-light depends on your specific application.

14. Process the samples for whole cell extraction or fixation and immunostaining using standard procedures.
   a. Evaluate the expression level of the doxycycline-inducible protein by immunoblot analysis (Figure 3A).
   b. Verify by standard fluorescence microscopy the ability of your recombinant protein to form biomolecular condensates upon light activation (Figure 3B).

Biotin - labeling of proteins within light-induced condensates

15. Timing: 2 days

   In this second major step, you will control the formation of biomolecular condensates with 488 nm light and label component proteins with biotin (Figure 4).

   The required number of cells depends on your specific application (e.g., immunoblotting, mass spectrometry). This protocol describes the analysis of TopBP1 proximal proteins by immunoblotting.
using $4 \times 10^6$ Flp-In$^{TM}$-REx$^{TM}$293 cells stably transfected with the doxycycline-inducible TurboID-TopBP1WT-mCherry-Cry2 construct.

Day 1 Morning:

15. Prewarm cell media, PBS, and trypsin at 37°C in a water bath.
16. Seed $4 \times 10^6$ Flp-In$^{TM}$-REx$^{TM}$293 cells/per 10 cm dish in 5 mL of DMEM medium supplemented with 10% fetal bovine serum (FBS) without any selection antibiotics. Use three dishes for a typical experiment:
   a. with doxycycline/ Light OFF
   b. without doxycycline/ Light ON
   c. with doxycycline/ Light ON

Day 1 Evening:

17. Add 4 µg/mL of doxycycline in a falcon tube containing 5 mL of DMEM supplemented with 10% FBS to reach a final concentration of 2 µg/mL doxycycline, as described in step 18.
18. Add the 5 mL DMEM supplemented with 10% FBS and 4 µg/mL doxycycline to the 5 mL cells culture from step 16. The final concentration of doxycycline is 2 µg/mL. Allow protein expression for 16–24 h.

Day 2:

19. Prewarm the DMEM supplemented with 10% FBS at 37°C.
20. Thaw the biotin stock solution.
21. Prepare in 15 mL Falcon tube 10 mL of DMEM supplemented with 500 μM biotin and keep at 37°C.
22. Place the illumination box in a humidified 37°C incubator with 5% CO₂ (Figure 2).
23. Replace the cell culture medium with 10 mL of DMEM media supplemented with 500 μM biotin and 2 μg/mL doxycycline.

⚠️ CRITICAL: Add the medium from the wall of the cell culture dish and not directly on the cells to avoid cell detachment.

24. Transfer rapidly the cell culture dish without cover into the custom-made illumination box and turn ON the device.

Note: At this stage, it is not essential to manipulate cells in a sterile environment because the cells will be collected and lysed for biochemical analyses immediately after optogenetic induction of condensates.

25. Expose the cells to blue light for 10 min of light-dark cycles (4 s light followed by 30 s dark).

Note: In principle, high doses of 488 nm light could generate reactive oxygen species. However, in comparison with dental photo-polymerization sources, for example, the light intensity used here (10 mW/cm²) is weak. Yet, it would be wise to minimize exposure times. For example, 3 minutes of light dark cycles (4 s light followed by 10 s resting in the dark) induce robust TopBP1 condensation. After optogenetic activation, condensates will dissolve progressively. Five minutes after light activation, approximately 40% of TopBP1 condensates have dissolved (Frattini et al. 2021). The light-dark cycle parameters required for the maintenance of condensates during proximity labeling depends on the cellular concentration and on the sequence of the protein that holds condensates together via multiple weak and cooperative interactions. The parameters of optogenetic activation need to be determined on a case-by-case basis.

26. Turn OFF the device.
27. Remove medium completely by aspiration and wash the cells carefully at 18°C–25°C with 5 mL/ of 1× PBS per dish.
28. Gently scrape all the cells off the plate with a cell scraper and place the cell suspension in a 15 mL Falcon tube. Keep on ice.
29. Centrifuge the cells at 400 \( \times \) g for 3 min.
30. Aspirate the medium and freeze the pellet at \(-80^\circ\)C.
31. Proceed to the next step.

**Pause point:** You can store the cell pellets at \(-80^\circ\)C for several weeks before proceeding to the next step.

### Affinity purification of biotinylated proteins

**Timing:** 2 days

The third major step is to isolate biotinylated proteins using streptavidin coated beads and identify these proteins by immunoblotting or proteomic analyses (Figure 5).

32. Prepare all the buffers listed in the Materials section (You can prepare them several days before the experiment).

**Note:** To reduce keratin contamination, use DNase/RNase-free tubes that have not previously been opened and wear gloves.

33. The day of the experiment, add 1X of complete protease and phosphatase inhibitors to 10 mL of RIPA/SDS Lysis buffer.
34. Thaw gently the frozen cell pellets.
35. Resuspend the cell pellet in 500 μL of RIPA/SDS Lysis buffer supplemented with 1X complete protease and phosphatase inhibitors and 250 U benzonase and transfer the lysate into 1.5 mL Eppendorf tube.
36. Incubate the pellet on a rotating wheel for 1 h at 4°C.
37. Sonicate the lysate on ice using a sonicator (40% amplitude, 3 cycles 10 s sonication- 2 s resting).

△ CRITICAL: Clean the sonicator probe between samples to avoid cross-contamination.

38. Spin down cellular debris at 16000 × g for 30 min at 4°C.

Note: During the centrifugation step, prepare the required amount of agarose-streptavidin (50 μL of agarose streptavidin beads slurry per condition). Resuspend gently the stock of streptavidin beads by tapping/flicking the bottom of the tube. Wash the beads twice with RIPA/SDS lysis buffer and resuspend in RIPA/SDS buffer.

39. Transfer the cleared supernatant to a new 1.5 mL Eppendorf tube.
40. Determine total protein concentration using Bradford protein assay.
41. To isolate biotinylated proteins from cell lysates, incubate up to 1 mg of cell lysates with the pre-washed agarose-streptavidin beads for 3 h on a rotating wheel at 4°C.
42. Spin the bead suspensions for 1 min at 400 × g, at 18°C–25°C, and carefully remove the supernatant by pipetting.
43. Wash the beads via resuspension and incubation on a rotating wheel for 3 min as indicated below. Between each step, pull down the beads by centrifugation for 1 min at 400 × g at 18°C–25°C and remove carefully the supernatant:
   a. Wash with 1 mL of RIPA/SDS Lysis buffer
   b. Wash with 1 mL Wash buffer 1
   c. Wash with 1 mL Wash buffer 2
   d. Wash with 1 mL Wash buffer 3
   e. Wash with 1 mL Wash buffer 4 (twice)

△ CRITICAL: Avoid pipetting the beads excessively, as they can bind to the pipet tips, leading to sample loss.

44. Add 60 μL of 2X Laemmli Sample buffer to each sample and flick to mix.
45. Heat samples for 10 min at 95°C to elute the proteins.

Pause point: The samples can be stored at –20°C or –80°C for at least 1 year.

△ CRITICAL: Avoid multiple boiling and freezing cycles of your samples.

46. For western blot analysis, resolve proteins by SDS-polyacrylamide gel electrophoresis (SDS-PAGE). We recommend loading 2–4 μg of proteins in the “Input” control well.
47. For Mass spectrometry analyses, use up to 2 mg of proteins isolated by streptavidin pull down. Analyze raw mass spectrometry data using the MaxQuant software (version 1.5.5.1) with standard settings. Use Perseus (version 1.6.1.1) for graphical representation and statistical analysis. Perform a standard t test to evaluate differences of protein abundance between samples.

EXPECTED OUTCOMES
The protocol allows probing and identifying proteins and their post-translational modifications in proximity of a protein scaffold before and after its self-assembly into a biomolecular condensate. An example is shown in (Frattini et al., 2021) where phospho-signals that indicate ATR kinase
activity are probed. In this experiment, phospho Ser1138 TopBP1 and phosho Thr1989 ATR signals are detected in proximity of TopBP1 specifically after induction of TopBP1 condensation by optogenetic activation (Figure 6A, lane 9). By contrast, phospho Ser1138 TopBP1 and phosho Thr1989 ATR are not detected in cells expressing the condensation defective mutant W1145R TopBP1 (Figure 6A, lane 12). In this particular example, the condensation of TopBP1 acts as a molecular switch for the activation of the ATR signaling pathway. The method can be exploited also to monitor the enrichment of proximal proteins as a result of their partitioning within condensates. For example, Figure 6B shows an immunoblot of FANCJ and BRCA1, two TopBP1 partner proteins detected by proximity labeling. Upon optogenetic induction of TopBP1 condensates, the BRCA1 signal increases (Figure 6B, lane 8), suggesting that BRCA1 concentrates within TopBP1 condensates.

LIMITATIONS

The fusion of a biotin ligase and a cryptochrome to a protein can potentially affect its stability, expression level and localization. These parameters should be verified beforehand to exclude potential alterations of protein properties and functions. Biotinylation of the TurboID fusion protein may interfere with protein activity and exert cytotoxicity (Branon et al., 2018).

In addition, the conjugation biotin to a protein can hinder modification sites and modify its charge. These parameters can potentially alter attractive interactions involved in protein condensation. Thus, we recommend to verify that the fusion of the protein scaffold to TurboID does not alter its condensation by optogenetic activation.

Target proteins that lack or have hindered primary amine groups will remain undetected. Furthermore, as the labeling radius is about 10 nm, steric hindrances may challenge the labeling of

Figure 6. Illustration of expected outcomes

Flip-In™T-REx™293 cell line expressing TurboID-TopBP1-mCherry-Cry2 were incubated with 500 μM of biotin and exposed to blue light for 10 min of light-dark cycles (4 s light followed by 30 s dark). TopBP1 partner proteins labeled with biotin were pulled-down using streptavidin-coated beads.

(A) Ser1138 phospho TopBP1, TopBP1, Thr1989 phospho ATR and ATR were detected by immunoblotting. This figure is reprinted with permission from Frattini et al., 2021.

(B) Immunoblot of TopBP1, BRCA1 and FANCJ isolated with streptavidin-coated beads.
proximal proteins when TurboID is fused to high molecular weight protein scaffolds. To overcome this limitation, it is possible to increase the size of the linker between the ligase and the scaffold. This will increase the labeling radius and maximize the capture of partner proteins (Kim et al., 2016).

Endogenous biotin levels are elevated in some cell culture conditions. If this is the case, then addition of exogenous biotin will not induce further protein biotinylation (May et al., 2020), and hinder the identification of proteins enriched within biomolecular condensates. Furthermore, high levels of endogenous biotin will promote the labeling of proximal proteins as the scaffold fused to TurboID journeys from the cytoplasm to the nucleus or through mitosis. This will confound the identifications of client proteins that partition within biomolecular condensates.

**TROUBLESHOOTING**

**Problem 1**
Poor expression of the scaffold fused to TurboID and Cry2 (step 5)

**Potential solution**
Verify the integrity of the construct as described in the cloning procedure. If necessary, verify by sequence a different clone, or correct mutations using standard molecular biology approaches. If no mutation is detected, increase the concentration of doxycycline and the time of induction.

**Problem 2**
The method does not allow delivering the condensates to specific locations (step 6).

**Potential solution**
The recombinant scaffold may include a sequence of amino-acids that functions as a targeting signal to transport the protein to its appropriate destination. For example, if the recombinant scaffold protein has a nucleo-localization signal, it should localize to the nucleus. Furthermore, the recombinant scaffold may associate with endogenous proteins to form condensates at specific nucleation sites. Alternatively, light-induced hetero-dimerization of CRY2 and CIB1 may be used to control the localization of a protein of interest through protein-protein interaction (Duan et al., 2017). For chromatin-associated condensates, the method may be coupled to a CRISPR-Cas9-based technology to anchor the condensate to a specific locus (Shin et al., 2018).

**Problem 3**
Chronic biotinylation of the TurboID fusion protein exerts cytotoxicity (step 8).

**Potential solution**
Use dialyzed FBS to reduce the level of endogenous biotin.

Reduce the time of induction of protein expression with doxycycline.

Reduce the time of incubation with exogenous biotin.

**Problem 4**
High level of endogenous biotin (step 16).

**Potential solution**
Use dialyzed FBS to minimize the abundance of endogenous biotin and verify that protein biotinylation is induced by exogenous biotin.
Problem 5
Biotinylation of proximal proteins occurs before condensation (step 23).

Potential solution
The protein scaffold fused to TurboID biotinylates proximal proteins before and after optogenetic induction of biomolecular condensates. The molecular events that arise specifically from the process of condensation, such as posttranslational modifications of proteins, are inferred by comparison with samples prepared before optogenetic activation. The method can be coupled to quantitative mass spectrometry approaches, such as stable isotope labeling with amino acids in cell culture (SILAC), or isobaric tagging reagents for quantitative proteomic analysis (iTRAQ) for high resolution quantitative analyses of condensation-driven protein modifications or protein compartmentalization.

Problem 6
Poor detection of proximal proteins or post-translational modifications (step 46).

Potential solution
Increase the time of incubation with exogenous biotin during optogenetic activation from 15 to 60 min.

Scale up the isolation of biotinylated proteins on streptavidin beads. Increase the amount of proteins loaded on SDS-PAGE gel.

For immunoblotting, use precast polyacrylamide gels, fresh preparations of primary antibody, increase antibody concentration and time of incubation.

RESOURCE AVAILABILITY
Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Angelos Constantinou (angelos.constantinou@igh.cnrs.fr)

Materials availability
The plasmid generated in this study; pCDNA5_FRT_TO_TurboID-mCherry-Cry2 has been deposited with Addgene under the ID number 166504.

Data and code availability
Original western blot images have been deposited to Mendeley data, https://doi.org/10.17632/zb35ntwpnj.1, and are publicly available as of the date of publication. Microscopy data reported in this paper will be shared by the lead contact upon request.

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AUTHOR CONTRIBUTIONS
Conceptualization, J.B. and A.C.; methodology, J.B.; validation, J.B. and A.C.; investigation, J.B. and E.A.; writing original draft, J.B. and E.A.; writing-review & editing, A.C., J.B., and E.A.; visualization, J.B. and E.A.; supervision, J.B. and A.C.; project administration, A.C.; funding Acquisition, A.C.
DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES

Banani, S.F., Lee, H.O., Hyman, A.A., and Rosen, M.K. (2017). Biomolecular condensates: organizers of cellular biochemistry. Nat Rev Mol. Cell. Biol. 18, 285–298.

Bracha, D., Wallis, M.T., and Brangwynne, C.P. (2019). Probing and engineering liquid-phase organelles. Nat. Biotechnol. 37, 1435–1445.

Bracha, D., Wallis, M.T., Wei, M.T., Zhu, L., Kurian, M., Avalos, J.L., Toettcher, J.E., and Brangwynne, C.P. (2018). Mapping local and global liquid phase behavior in living cells using photo-oligomerizable seeds. Cell 175, 1467–1480.e13.

Branon, T.C., Bosch, J.A., Sanchez, A.D., Udeshi, N.D., Svinina, T., Carr, S.A., Feldman, J.L., Perrimon, N., and Ting, A.Y. (2018). Efficient proximity labeling in living cells and organisms with TurboID. Nat. Biotechnol. 36, 880–887.

Duan, L., Hope, J., Ong, Q., Lou, H.Y., Kim, N., McCarthy, C., Acero, V., Lin, M.Z., and Cui, B. (2017). Understanding CRY2 interactions for optical control of intracellular signaling. Nat. Commun. 8, 547.

Frattini, C., Promonet, A., Alghoul, E., Vidal-Eychene, S., Lamarque, M., Blanchard, M.P., Urbach, S., Basbous, J., and Constantinou, A. (2021). TopBP1 assembles nuclear condensates to switch on ATR signaling. Mol. Cell 81, 1231–1245.e6.

Hyman, A.A., and Simons, K. (2012). Cell biology. Beyond oil and water-phase transitions in cells. Science 337, 1047–1049.

Kim, D.I., Jensen, S.C., Noble, K.A., Kc, B., Roux, K.H., Motamedchiaboki, K., and Roux, K.J. (2016). An improved smaller biotin ligase for BioID proximity labeling. Mol. Biol. Cell 27, 1188–1196.

Lam, S.S., Martell, J.D., Kamer, K.J., Deerinck, T.J., Ellisman, M.H., Moota, V.K., and Ting, A.Y. (2015). Directed evolution of APEX2 for electron microscopy and proximity labeling. Nat. Methods 12, 51–54.

Lyon, A.S., Peeples, W.B., and Rosen, M.K. (2020). A framework for understanding the functions of biomolecular condensates across scales. Nat. Rev. Mol. Cell Biol. 22, 215–235.

May, D.G., Scott, K.L., Campos, A.R., and Roux, K.J. (2020). Comparative application of BioID and TurboID for protein-proximity biotinylation. Cells 9, 1070.

Palayam, M., Ganapathy, J., Guercio, A.M., Tal, L., Deck, S.L., and Shabek, N. (2021). Structural insights into photoactivation of plant Cryptochrome-2. Commun. Biol. 4, 28.

Roux, K.J., Kim, D.I., Raida, M., and Burke, B. (2012). A promiscuous biotin ligase fusion protein identifies proximal and interacting proteins in mammalian cells. J. Cell Biol. 196, 801–810.

Shin, Y., and Brangwynne, C.P. (2017). Liquid phase condensation in cell physiology and disease. Science 357, eaaf4382.

Shin, Y., Chang, Y.C., Lee, D.S.W., Berry, J., Sanders, D.W., Ronceray, P., Wingreen, N.S., Haataja, M., and Brangwynne, C.P. (2018). Liquid nuclear condensates mechanically sense and restructure the genome. Cell 175, 1481–1491.e13.

You, J.Y., Dunham, W.H., Hong, S.J., Knight, J.D.R., Bashkurov, M., Chen, G.I., Bagci, H., Rathod, B., MacLeod, G., Eng, S.W.M., et al. (2018). High-density proximity mapping reveals the subcellular organization of mRNA-associated granules and bodies. Mol. Cell 69, 517–532.e1.