Measuring ATP levels within the cytosol of living cells in animals is important to understand how cellular activities are energetically supported, but is challenging because of tissue complexity and ATP sensor limitations. In this protocol, we describe how to quantify ATP levels using PercevalHR in *C. elegans* larvae during anchor cell invasion. PercevalHR is a fluorescent biosensor that reports the cytoplasmic ATP:ADP ratio. The protocol can be adapted to analyze the ATP:ADP ratios within other cell types in *C. elegans*.

Publisher’s note: Undertaking any experimental protocol requires adherence to local institutional guidelines for laboratory safety and ethics.
Protocol

Visualizing cytoplasmic ATP in C. elegans larvae using PercevalHR

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SUMMARY

Measuring ATP levels within the cytosol of living cells in animals is important to understand how cellular activities are energetically supported, but is challenging because of tissue complexity and ATP sensor limitations. In this protocol, we describe how to quantify ATP levels using PercevalHR in C. elegans larvae during anchor cell invasion. PercevalHR is a fluorescent biosensor that reports the cytoplasmic ATP:ADP ratio. The protocol can be adapted to analyze the ATP:ADP ratios within other cell types in C. elegans. For complete details on the use and execution of this protocol, please refer to Garde et al. (2022).

BEFORE YOU BEGIN

This protocol describes the use of a fluorescent ATP:ADP ratiometric biosensor PercevalHR to image the energy status of the anchor cell (AC) in C. elegans but can be used to image the ATP:ADP ratio of other worm cells.

Institutional permissions
Institutional permissions are not required for working with C. elegans.

PercevalHR basics

The ATP:ADP ratio is a key parameter for energy metabolism, as it measures the free energy available to drive energy consuming cellular processes (Tantama et al., 2013). Empirical studies have found that an increase in the ATP:ADP ratio usually indicates an increase in ATP levels within the cell (Zanotelli et al., 2018).

PercevalHR is comprised of the bacterial ATP-binding protein GlnK1 fused to a circularly permuted green fluorophore cpmVenus. ATP and ADP both compete for binding with the nucleotide binding pocket of GlnK1 and cause conformational changes in the sensor. Each nucleotide causes a unique change in the excitation spectra of PercevalHR—ATP binding increases its emitted fluorescence for 500 nm excitation while ADP binding increases its fluorescence for 420 nm excitation (Figure 1A) (Tantama et al., 2013). The ATP:ADP ratio is calculated as the ratio between the fluorescence intensities in the ATP bound state and the ADP bound state, making the ATP:ADP ratio independent of the concentration of sensor in the cell. This allows greater tolerance for variability in expression levels within cells and enables robust temporal analysis of changes in ATP:ADP ratios.

Anchor cell invasion

The AC is a specialized uterine epithelial cell that invades through its underlying basement membrane (BM) in the larval L3 stage, to establish a connection between the uterine and vulval tissues.
for egg laying. BM invasion by the AC is highly stereotyped and coordinated with the divisions of the underlying 1º-fated P6.p vulval precursor cells (VPCs) (Sherwood and Sternberg, 2003). The AC is specified prior to the L3 stage and grows during the P6.p 1-cell stage. The AC initially breaches...
the BM with small, F-actin rich membrane associated protrusions called invadopodia during the P6.p 2-cell stage. Following BM penetration, the AC generates a large invasive protrusion at the breach site to widen the BM gap during the P6.p 2-4-cell and early 4-cell stages. The AC retracts the protrusion and completes invasion by the P6.p late 4-cell stage and attaches to the central vulval cells to initiate uterine vulval connection (Figure 1B).

**Strain design considerations**
When designing the AC-specific PercevalHR strain, we considered the following parameters:

1. **Promoter**: Select a promoter that drives PercevalHR at high levels and with specific expression in the AC during AC invasion. In this study we used the lin-29 promoter (Figures 1C and 1D) to drive strong AC-specific expression of PercevalHR starting ~3 h prior to AC invasion and continuing ~5 h after invasion (Naegeli et al., 2017).

2. **Transgene generation method**: In C. elegans there are two primary approaches for transgene expression:
   a. generation of a multi-copy array with subsequent integration into the genome.
   b. generation of a single copy insertion via CRISPR-Cas9.

   **Note**: Due to the lower efficiency of mVenus fluorescence with 405 nm excitation, we found that PercevalHR expression and thus signal was highest in an integrated strain expressing a multi-copy insertion of the lin-29p::PercevalHR. For our experiments, we selected multi-copy array transgenic worms with robust signal in both fluorescent channels (refer to troubleshooting problem 2 for more details).

   **Note**: Strains constructed using multi-copy insertion exhibit some variability in expression between animals. Due to the concentration-independent nature of the ATP:ADP ratio measured by PercevalHR, this does not affect the analysis.

**Designing and generating PercevalHR expressing strains**

© Timing: 4–6 weeks

3. Design the plasmid expressing PercevalHR under the promoter of choice (in our case lin-29p).
   a. PCR amplify PercevalHR sequence from the C. elegans codon optimized PercevalHR plasmid (pDONR221, obtained from Calico Life Sciences, key resources table).
   b. Use directional cloning or Gibson assembly to insert PercevalHR into a plasmid containing the cell-type specific promoter.

   **Note**: Sequence the entire PercevalHR construct to ensure that no mutations or deletions are introduced during amplification or assembly steps. The sequence of codon optimized PercevalHR contains repetitive regions that may result in deletions during amplification and assembly.

4. Microinject 50 ng/μL of the completed PercevalHR construct (lin-29p::PercevalHR), 50 ng/μL of unc-119 rescue DNA (key resources table), 50 ng/mL PBS SK(+), and 25 ng/mL cut salmon sperm DNA into gravid adult C. elegans unc-119 (ed4) mutant hermaphrodites, which are severely impaired in their locomotion. This allows for easy selection of transgenic animals, which will have wild-type locomotion, for future integration (Berkowitz et al., 2008).

5. Select 2–3 lines of wild-type animals exhibiting normal fertility and morphology, and expressing strong PercevalHR signal (observed via the GFP channel on a fluorescent dissecting or wide-field microscope) for integration via gamma irradiation. For additional detailed resources on generating integrated strains in C. elegans refer to Wormbook “Transformation and Microinjection” (Evans, 2006).
### KEY RESOURCES TABLE

| REAGENT or RESOURCE                  | SOURCE                        | IDENTIFIER |
|--------------------------------------|-------------------------------|------------|
| **Bacterial and virus strains**      |                               |            |
| E. coli OP50 standard food           | Caenorhabditis Genetics Center | OP50       |
| E. coli HT115 RNAi feeding strain    | Caenorhabditis Genetics Center | HT115 (DE3)|
| **Chemicals, peptides, and recombinant proteins** |                               |            |
| 100% white petrolatum                | Unilever                      | Vaseline   |
| Lanolin                              | BeanTown Chemical              | Cat # 144255|
| Paraffin wax; pellets                | Sigma-Aldrich                 | Cat # 18634|
| Cholesterol                         | Sigma-Aldrich                 | Cat # C8667|
| Bacto™ Peptone                       | Thermo Fisher Scientific      | Cat #211677|
| Agar A                               | Bio Basic                     | Cat #F80010|
| MgSO₄                                 | Macron Chemicals              | Cat # 6070-12|
| NaCl                                 | Macron Chemicals              | Cat #7581-12|
| KH₂PO₄                                | VWR chemicals                 | Cat # BDH9268|
| Na₂HPO₄                               | EMD Millipore Corp.           | Cat # 1063460500|
| DIFCO™ Noble agar                    | BD                            | Cat. # 214230|
| Rotenone                             | EMD Millipore Corp.           | Cat. # 557368|
| Polybead microspheres 0.10 μm        | Polysciences                  | Cat # 00875-15|
| Dimethyl sulfoxide (DMSO)            | VWR                           | Cat # MK494802|
| Levamisole hydrochloride             | MilliporeSigma                | Cat. # L9756|
| **Experimental models: Organisms/strains** |                               |            |
| C. elegans unc-119 injection strain; Genotype: unc-119 (ed4) III – gravid adult animals | Caenorhabditis Genetics Center | WB Strain |
| **Recombinant DNA**                  |                               |            |
| PercevalHR (C. elegans codon optimized) | (Bohnert and Kenyon, 2017)    | Calico Labs pDONR221|
| unc-119 rescue construct (pPK605)    | AddGene (Sieburth et al., 1998) | Cat #38148|
| Ultrapure Salmon Sperm DNA           | Thermo Fisher Scientific      | Cat # 15632011|
| pBlueScript II SK (+) vector         | Agilent                       | Cat # 212205|
| **Software and algorithms**          |                               |            |
| μManager                             | (Edelstein et al., 2014)      | Version 2.0.0 (https://micro-manager.org/Download_Micro-Manager_Latest_Release) |
| Fiji                                 | (Schneider et al., 2012)      | Version ImageJ 1.52p (https://imagej.net/software/fiji/downloads) |
| Zen Black 3.2                        | Carl Zeiss Microscopy         | Version 3.2 |
| **Other**                            |                               |            |
| Petri plates 60 mm (non-vented)      | Tritech Research              | T3308P     |
| Glass slides (25 × 75 × 1 mm)        | Globe Scientific Inc.         | Cat # 1301 |
| Cover glass (22 × 22 mm – No. 1.5)   | Fisher Brand                  | Cat # 12541B|
| PYREX™ Spot plate                    | Fisher Scientific             | Cat # 13-748B|
| Platinum wire (for worm pick)        | SPI supplies                  | Cat # 01703-AC|
| Open ended melted capillary KimBLE® KimAX® (for mouth pipette) | MilliporeSigma             | Cat # DWK34500-99 |
| 15” Aspirator Tube Assembly (for mouth pipette) | VWR                  | Cat # 53507-278|
| Zeiss AxioImager.A1 (discontinued) – Zeiss AxioImager 2 is comparable | Zeiss                  | n/a        |
| Yokogawa CSU-10 (discontinued) – CSU-W1 or CSU-X1 is comparable | Yokogawa Electric Corporation | n/a        |
| 10× EC Plan-Neofluor 10×/0.3 M27 (for locating worms) | Zeiss                  | Cat # 420340-9901-000 |

(Continued on next page)
MATERIALS AND EQUIPMENT

**M9 buffer**

| Reagent      | Final concentration | Amount |
|--------------|---------------------|--------|
| Na₂HPO₄      | 42.2 mM             | 6 g    |
| KH₂PO₄       | 22 mM               | 3 g    |
| NaCl         | 85.5 mM             | 5 g    |
| 1 M MgSO₄    | 1 mM                | 1 mL   |
| Deionized water diH₂O | n/a    | 999 mL |
| Total        | n/a                 | 1 L    |

Note: Sterilize by autoclaving and aliquot into 100 mL bottles. Unopened stocks can be stored at room temperature (~20°C–25°C) for 3 months.

**Nematode growth medium (NGM agar plates)**

| Reagent        | Final concentration | Amount |
|----------------|---------------------|--------|
| Agar A         | 17 g/L              | 34 g   |
| Peptone        | 2.5 g/L             | 5 g    |
| NaCl           | 25.66 mM            | 3 g    |
| Cholesterol (5 mg/mL) | 12.92 μM | 2 mL   |
| diH₂O          | n/a                 | 1.95 L |
| Total          | n/a                 | 2 L    |

Note: Sterilize by autoclaving for 60 min and then cool to 55°C in a water bath before adding 50 mL of 1 M KPO₄ buffer (pH 6.0), 2 mL of 1 M MgSO₄ and 2 mL of 1 M CaCl₂. Pour 8 mL of warm NGM per 60 mm non-vented plastic petri dish using a peristaltic pump and sterile technique and allow to cool overnight. Store plates upside down at 4°C for up to 3 months. Warm to room temperature (~20°C–25°C) before seeding with OP50 or HT115 E. coli bacteria for feeding and worm husbandry.

**Levamisole stock solution (anesthetic)**

© Timing: 5–10 min

- Prepare 1.5 mL of a 200 mM stock solution of levamisole in sterile water.
• Aliquot the stock solution into Eppendorf tubes in 150 μL increments for storage at −20°C.

⚠ CRITICAL: The efficacy of levamisole solutions declines over time and with repeated freeze-thaw cycles. We recommend preparing fresh levamisole stock solution every 2–3 weeks for optimal immobilization during image acquisition.

5% (weight/volume) noble agar

◎ Timing: 5–10 min

• Melt 5% (weight/volume) noble agar suspended in water in the microwave until dissolved.
• Aliquot 1.5 mL of the solution into disposable glass tubes and cover with a plastic cap. Store at room temperature (~20°C–25°C) for up to 3 months.
• Before use, melt the noble agar in the glass tube over a Bunsen burner and store in a 70°C heat block to keep the solution liquid.

VALAP

◎ Timing: 15–20 min

• Combine equal parts of Vaseline, lanolin, and paraffin wax by weight for a total weight of 45 g in a glass beaker.
• Heat the beaker over a hot plate on low heat (close to minimum setting) and stir occasionally with a metal spatula until the mixture has completely melted and the liquid appears a golden yellow color and homogenous. Take care not to overheat this mixture and warm only until all the components liquefy.
• Aliquot 1.5 mL into Eppendorf tubes and store them at room temperature (~20°C–25°C). Stocks will last indefinitely.
• When ready to use, warm the Eppendorf tube in a 70°C heat block to melt the mixture for painting on to slides with a fine paintbrush.

STEP-BY-STEP METHOD DETAILS
Preparing *C. elegans* expressing PercevalHR for imaging

◎ Timing: 1–2 weeks

ATP synthesis enzymes and levels of ATP in whole body assays are altered by starvation (Chin et al., 2014). To avoid starvation related changes in ATP levels, maintain animals expressing PercevalHR as follows:

1. Grow animals on the standard OP50 bacteria feeding strain for 2–3 generations without starvation before imaging. Additional resources on *C. elegans* culture, handling, and maintenance can be found in Wormbook “Maintenance of *C. elegans*” (Stiernagle, 2006).
2. Use plate-level morphological characteristics to select animals at the appropriate stage of development for PercevalHR imaging from an asynchronous, well-fed population. To select animals to examine AC invasion, assess vulval morphology using a dissecting microscope to identify mid L3 larval staged worms.
3. Pick mid-L3 larval worms from the plate with a platinum wire and mount onto an agar pad on a microscope slide.
4. Use differential interference contrast (DIC) microscopy and high magnification on a compound microscope to identify the AC and its stage of invasion (e.g., pre-, during, post-; see Figure 1B and “Imaging ACs expressing PercevalHR using point-scanning or spinning disk confocal microscopy” below).
Note: If synchronized populations worms are required for the experiment, we recommend using sequential egg-laying to achieve a developmentally synchronized population instead of bleach synchronization and subsequent starvation induced L1 arrest, as L1 arrest introduces metabolic adaptations that could persist into larval development (Jobson et al., 2015).

△ CRITICAL: When conducting RNAi experiments using strains expressing PercevalHR, animals should be grown on RNAi bacteria (HT115) expressing empty vector control for 2–3 generations without starvation before transferring onto RNAi conditions for synchronized egg-laying. This prevents OP50 contamination on RNAi plates. Animals can then be transferred to blank NGM plates with no bacteria for 45–60 min to clear HT115 empty vector bacteria, before transferring to experimental RNAi plates.

Microscopy settings

© Timing: 15–20 min

PercevalHR can be imaged using confocal microscopy. This step describes how to use either a Zeiss LSM 880 and Zen Black 3.2 software as an example of a point scanning confocal or a Zeiss Axiomager microscope equipped with a Yokogawa CSU-10 spinning disc confocal, VersaLase(4) laser system and an ImqagEM EMCCD camera and Micromanager software as an example of a spinning disk confocal. These instructions can be modified for use with other acquisition software.

5. Point-scanning confocal setup using a Zeiss 63× Plan-Achromat 1.4NA oil immersion objective: set up the following channels in Zen Black (Figure 2).
   a. Channel 1: Excitation 488 nm.
   b. Channel 2: Excitation 405 nm.
   c. Emission: 490–553 nm.
   d. Pinhole size: 2.8 Airy units.

Note: The recommended setup will result in a 2 μm thick central slice of signal from the AC in both the ATP-bound (488 nm) and ADP-bound (405 nm) channels. We opted to forgo optical sectioning due to the homogeneity of the cytoplasmic ATP:ADP ratio in the Z-axis and the ability to control pinhole size on a point-scanning confocal microscope.

6. Spinning disk confocal setup using a Zeiss 100× Plan-Achromat 1.4NA oil immersion objective: set up the following acquisition settings in Micromanager (Figure 3).
   a. Channel 1 (Channel 488): Excitation laser 488 nm, Emission filter 525 nm.
   b. Channel 2 (Channel 405 Perceval): Excitation laser 405 nm, Emission filter 525 nm.
   c. Z-step: 0.5 μm.

Note: The recommended setup will result in ~0.3 μm thick slices of signal from the AC in both the ATP-bound (488 nm laser) and ADP-bound (405 nm laser) channels. To collect sufficient signal from the central 2 μm of the AC, we imaged five central slices of the AC that were then summed into a projection for analysis.

Note: Laser power settings will vary across image acquisition systems and laser power should be optimized for each channel. Image acquisition should result in at least ~1.5–2 fold greater average cytoplasmic signal compared to average background signal in both acquisition channels for the most accurate ATP/ADP ratio determination. Care should be taken when choosing the laser power for the 405 nm laser to balance the best signal collection with the potential negative problem of photobleaching during acquisition (refer to troubleshooting problem 4 for more details). We have found that if laser power is not optimized, there can be insufficient
signal compared to background in the ADP bound channel. This can lead to aberrantly high ATP:ADP ratio values after image processing and it can limit the detection of a further reduction in ADP levels (refer to troubleshooting problem 3 for more details).

Figure 2. Microscopy settings for Zeiss LSM 880 (point-scanning confocal) in Zen Black, related to step 5
(A and B) This figure displays the two acquisition channels for PercevalHR image acquisition. The excitation wavelength, laser power, gain and pinhole size settings for the ATP-bound state are highlighted in green-dashed boxes (488 LSM), and the excitation wavelength, laser power, gain and pinhole size settings for the ADP-bound state are highlighted in cyan-dashed boxes (405 LSM). The wavelengths for the emission range are highlighted in orange-dashed boxes.
Anesthetizing and mounting *C. elegans* for imaging

### Timing: 30 min

Animals expressing PercevalHR must be immobilized for accurate alignment between the ATP and ADP channels to generate a ratiometric image during analysis. This section will provide detailed steps for anesthetizing and mounting L3 larvae immersed in anesthetic solution onto a 5% noble agar pad for subsequent imaging (Figure 4). For additional detailed instructions and videos on transferring and anesthetizing worms and making agar microscopy pads, see Nature Protocols article, “Live-cell confocal microscopy and quantitative 4D image analysis of anchor-cell invasion through the basement membrane in *Caenorhabditis elegans*” (Kelley et al., 2017).

7. Anesthetize *C. elegans* in levamisole.
   a. In one well of a glass spot plate, dilute 200 mM levamisole stock to a 5 mM working stock in 100 μL of M9 buffer (see materials and equipment).
   b. Using a platinum wire pick, transfer 20–30 animals into the well of the glass spot plate. Limit the transfer of *E. coli* from the culture plate as bacteria will interfere with anesthetizing and later imaging. Swirling the pick gently and moving it in and out of the buffer slowly will help move the animals off the pick without introducing excess *E. coli* into the buffer (Figure 4A).
If large chunks of E. coli are being introduced into the anesthetic, refer to troubleshooting problem 1 for alternative transfer techniques.

c. Allow animals to anesthetize at room temperature for 15–20 min. This is sufficient to immobilize animals for up to 1.5 h if maintained in the anesthetic solution on the agar pad while imaging, since AC invasion occurs between two developmental starvation-induced developmental arrest checkpoints (Schindler et al., 2014).

8. Mount C. elegans onto agar pad.
   a. Prepare a 5% (weight/volume) noble agar pad (see materials and equipment) that is sized so that it is slightly smaller than a 22 mm × 22 mm coverslip (Figure 4B).
   b. Transfer 20–30 anesthetized animals onto the prepared agar pad using a mouth pipette. Remove excess M9 buffer using the edge of a Kimwipe or mouth pipette. Gently place the coverslip over the animals on the agar pad starting with one edge and then slowly lowering the coverslip until the opposite edge rests on top of the pad. Then, seal the edges of the coverslip using a fine paintbrush and VALAP, leaving two small gaps in the VALAP seal to allow gas-exchange (Figures 4C and 4D).

Note: To image PercevalHR in the AC (or any other cell type), animals should not be anesthetized in sodium azide. Although sodium azide is a common anesthetic used to immobilize C. elegans for microscopy, it is a potent inhibitor of mitochondrial ATP metabolism and will impact cellular ATP levels and PercevalHR readout (Harvey et al., 1999; Tsubaki and Yoshihara, 1993). Instead, we use 5 mM levamisole with and without polystyrene beads. Refer to troubleshooting problem 5 for more details.
**Imaging ACs expressing PercevalHR using point-scanning or spinning disk confocal microscopy**

**Timing:** 2 h

After mounting onto a slide, each animal expressing PercevalHR is imaged once at a single developmental time-point.

9. Place the slide from above with anesthetized L3 staged animals expressing PercevalHR onto the microscope stage and bring the AC into focus.
   a. Find L3 staged worms using a low magnification (10× or 20× air objective) and mark X-Y coordinates in Zen Black or Micromanager.
   b. Switch to higher magnification (63× or 100× oil-immersion objective) and using DIC microscopy, determine the precise developmental stage of the AC using the number of P6.p vulval precursor cell descendants as shown previously (Kelley et al., 2017) and schematic in Figure 1B.
   c. Using the 488 nm laser bring the central plane of the AC into focus. In this plane, the nucleus, which lacks PercevalHR signal, should appear at its largest (Figures 1C and 1D, asterisk).

△ **CRITICAL:** Low laser power should be used when selecting and bringing ACs into focus to prevent photobleaching. We also recommend using the 488 nm laser, which has a longer wavelength and results in less phototoxicity and photobleaching of PercevalHR signal compared to the 405 nm laser.

10. Acquire PercevalHR images.
    a. **Point-scanning confocal microscope:**
        i. Set both lasers to their required laser power. We found 22% for the 488 nm laser and 8% for the 405 nm laser to be optimal laser power settings on a Zeiss LSM 880.
        ii. Acquire fluorescence signal from the ATP-bound and ADP-bound channels simultaneously through the emission filter stated in microscopy settings.
    b. **Spinning-disk confocal microscope.**
        i. Set both channels to their required laser powers and exposure settings. We found 18% laser power and 250 ms exposure for the ATP-bound channel and 488 nm laser, and 35% laser power and 400 ms exposure for the ADP-bound channel and 405 nm laser to be optimal settings on a Zeiss AxioImager equipped with an ImagEM EMCCD camera and VersaLase(4) laser system.
        ii. Acquire confocal z-stack in both channels.

**PercevalHR image processing**

**Timing:** 1–2 h

The ATP:ADP ratio within the cell or tissue is determined by dividing the fluorescence intensity of the pixels in the ATP-bound channel (488 nm channel) by the intensity of the pixels in the ADP-bound channel (405 nm channel). For a video tutorial on PercevalHR image processing and analysis, refer to Methods video S1.

11. Preparing ratiometric PercevalHR image.
    a. Open .tiff files directly in Fiji. When using proprietary (Zeiss, Nikon, Andor etc.) file formats such as .czi, import images using the Bio-Formats Importer plugin “Plugins > Bio-Formats > Bio-Formats Importer. The Bio-Formats plugin package can be downloaded from https://www.openmicroscopy.org/bio-formats/.
b. If the two channel image files are combined, split the 488 nm and 405 nm channels apart using “Image > Color > Split Channels”. Ensure that the two images have different names or unique identifiers for future steps (e.g., C1 and C2 at the end of the file name).

c. If using data in the form of a single 2 μm thick slice from a point-scanning confocal as in step 1, proceed to part d. below. If the data were collected as z-stacks, it is important to create sum projections of each channel separately to include all the pixel values in the core of the cell. This can be done by “Image > Stacks > Z Project...” and choosing “Sum Slices” from the “Projection type” dropdown menu.

d. To obtain the ATP:ADP ratio, the pixel values in the 488 nm channel must be divided by the pixel values in the 405 nm channel using the Image Calculator function “Process > Image Calculator”. Within the Image Calculator window, use the “Operation” dropdown menu to select “Divide”. Select the 488 nm image as Image 1 and the 405 nm image as Image 2. Check the boxes for “Create new window” and “32-bit (float) result”.

e. This is the final ratiometric ATP:ADP image. Each pixel value in the image represents the ATP:ADP ratio of that pixel of the cell.

Optional: When the range of pixel values in the ratiometric image does not extend through the full bit-depth of the detector, images should be contrast-stretched (the displayed minimum and maximum pixel value adjusted) to display only the range of pixel intensities within the cell being imaged. These pixel values can set using “Image > Adjust > Brightness and Contrast > Set”. For publication, the same displayed minimum and maximum pixel values should be set for all images within an experiment to facilitate accurate comparison. To better visualize and represent heterogeneities in cellular distribution of ATP:ADP ratio, this image can also be represented as a spectral heatmap using “Image > Lookup Tables > 16 color” (See Figure 1B). Additional resources about displaying fluorescence images can be found in (Johnson, 2012).

12. Quantifying PercevalHR images.

a. To quantify the total ATP:ADP ratio within the cell, draw a region of interest (ROI) using the freehand selection tool around the whole cell and measure the mean pixel values within the selection (“Analyze > Measure” or “m” key shortcut). Use the same ROI to measure the mean background pixel values (within the worm but away from the PercevalHR expressing cell). The final ATP:ADP ratio value for each cell can be represented as: [mean intracellular pixel intensity] – [mean background pixel intensity].

b. To quantify local enrichment of ATP:ADP ratio within a cell, draw ROIs around the regions to be compared (e.g., basal vs. apical) and measure the mean pixel values. Compute the background corrected ATP:ADP values for each region using the background value as determined above. The relative enrichment of one region compared to the other can then be represented as a ratio of their final ATP:ADP ratio values.

EXPECTED OUTCOMES

The ATP:ADP ratio within mammalian cells is in the range of 1 to >100 (Tantama et al., 2013). It can also be spatially heterogeneous and exhibit temporal dynamics. Using PercevalHR to visualize the ATP:ADP ratio within intact cells in C. elegans can allow for detailed characterization of ATP:ADP ratio distribution within the cell as well as the temporal dynamics of ATP production during biological processes. In combination with genetic mutants, RNAi mediated knockdown, and pharmacological inhibition, it can also be used to determine the genes and metabolic processes driving ATP production. For example, visualizing PercevalHR in the AC allowed us to determine that the AC has a 75% higher ATP:ADP ratio within the basal region at the site invasion compared to the apical domain (Figure 5A). Further, imaging PercevalHR over developmental time revealed that the AC exhibits a dynamic and transient 2.5-fold burst in ATP:ADP ratio levels during BM breaching (Figure 5A). Finally, by imaging ACs expressing PercevalHR after treatment with a mitochondrial...
oxidative phosphorylation (OXPHOS) inhibitor rotenone, we were able to detect a large reduction in the ATP:ADP ratio, which demonstrates that ATP levels in the AC are dependent on mitochondrial OXPHOS (Figure 5B).

QUANTIFICATION AND STATISTICAL ANALYSIS

The ATP:ATP ratio values calculated for each cell via PercevalHR processing and analysis can be plotted on box-and-whisker plots or violin plots to accurately display the spread of the data and help with data comprehension and comparison. Statistical comparisons of the total PercevalHR ATP:ADP ratio values between data sets – over developmental time, between treatment and control groups etc. – can be conducted using a standard statistical analysis i.e., t-test when comparing two groups and one-way ANOVA with post-hoc Tukey's test when comparing 3 or more groups.

LIMITATIONS

While this protocol is useful for imaging PercevalHR to measure the ATP:ADP ratio within the AC during invasion, it has several limitations. First, the image acquisition settings we describe are optimized to image PercevalHR in the AC at a single time-point. We found that these settings cannot be used to measure ATP:ADP ratio dynamics with repeated imaging, which is required for time-lapse analysis as repetitive imaging caused significant photobleaching. This limitation might be overcome in other cells if a high fluorescence signal can be achieved with PercevalHR using lower laser power in both channels to reduce photobleaching over time. We found this was not possible in the AC using the lin-29 promoter. Another potential limitation is that this protocol relies on the presence of a strong cell-type specific promoter to drive sensor expression. Thus, adaption of this protocol for other cells and tissues in C. elegans will require selection of high expression promoters. PercevalHR (like many fluorescent biosensors) is sensitive to low pH, thus it is critical to ensure that changes in distribution or magnitude of ATP:ADP ratio detected in cells are not because of
variations in intracellular pH. In (Garde et al., 2022), we used the pH sensor Rosella (Rosado et al., 2008) to independently verify that there were no pH gradients in the AC. However, the fluorescence spectra of Rosella (emission at ~508 nm and 587 nm) overlap with that of PercevalHR, precluding simultaneous analyses of both the ATP:ADP ratio and pH in the AC. To our knowledge, a single-wavelength pH sensor based on a red fluorophore such as pHRed (to prevent spectral overlap with PercevalHR) has not been optimized for use in C. elegans. A final limitation is that the image processing pipeline we describe requires manual intervention when selecting ROIs for analysis, which may limit high-throughput studies. However, for large-scale screens, an additional cell-type specific cytoplasmic or membrane marker with distinct fluorescent excitation and emission properties from PercevalHR (e.g., a red fluorophore) could be used for image segmentation and ROI generation, followed by semi-automated analysis using macros.

**TROUBLESHOOTING**

**Problem 1**
Excess *E. coli* introduced into levamisole anesthetic buffer when transferring animals from *E. coli* seeded NGM plates (step 7).

**Potential solution**
To reduce the transfer of *E. coli* into the well of the spot plate containing levamisole, animals can first be transferred from *E. coli* containing plates onto blank unseeded NGM plates for 5 min. Animals can then be transferred from blank plates into the levamisole anesthetic solution in the spot plate. If transferring animals from a blank plate with a pick is challenging, wash animals off the blank plate with 1 mL of M9 buffer (using a 1 mL pipette tip with the end cut off to widen it and prevent damage to the animals) into a 1.5 mL Eppendorf tube and gently pellet them in a benchtop centrifuge at ~200 x g. Discard the supernatant and transfer 100 μL of worm pellet into an empty well of the glass spot plate. Add 2.5 μL of the 200 mM levamisole stock solution to the 100 μL of liquid in the well, resulting in a 5 mM working concentration of levamisole to anesthetize the animals.

**Problem 2**
Low fluorescence signal in both channels of PercevalHR acquisition (before you begin – strain design considerations).

**Potential solution**
Low fluorescence intensity (less than 1.5-fold signal-to-background) in both acquisition channels at the time of imaging could indicate that the PercevalHR transgene selected for the experiment is not highly expressed in the cell. This could suggest either a low-copy integrant or a weak promoter being chosen to drive expression. Try opting for another integrated line with higher levels of PercevalHR expression for imaging and analysis or changing the promoter used to drive PercevalHR expression.

**Problem 3**
Low fluorescence intensity in ADP-bound channel of PercevalHR. Insufficient signal compared to background fluorescence in the ADP bound channel can result in artificially high ATP:ADP ratio values after image processing (step 10).

**Potential solution**
We have found that the fluorescence intensity of mVenus when excited by 405 nm laser (ADP-bound form) can be significantly lower than that of mVenus excited by 488 nm wavelengths (ATP bound) on a spinning disk confocal when both lasers are set to similar laser powers. Furthermore, *C. elegans* exhibit more autofluorescence when excited with 405 nm wavelength compared to 488 nm wavelength. To ensure that sufficient signal is being collected in the ADP bound channel, the signal intensity in the ADP-bound channel can be raised by increasing either the laser power, gain, or exposure time. We have found that optimizing an acquisition setting that results in at least a 1.5-fold
ADP-bound channel signal compared to animal background fluorescence results in consistent ATP:ADP ratios after image processing (Figure 6).

**Problem 4**
PercevalHR signal is photobleaching during image acquisition of the z-stack using a spinning-disk confocal (step 10).

**Potential solution**
405 nm light can be phototoxic for cells (Wäldchen et al., 2015) and cause photobleaching in both PercevalHR acquisition channels. To reduce possible phototoxicity and the photobleaching effects of image acquisition we have two suggestions:

- Reduce the laser power of the 405 nm laser if it is possible to do so while still collecting sufficient signal intensity (see problem 3).
- If the 405 nm laser power cannot be reduced further, we recommend acquiring images first in the 488 nm channel and then in the 405 nm channel to reduce the impact of the 405 nm laser on the 488 nm acquisition.

**Problem 5**
Animals move during imaging (steps 7 and 10).

**Potential solution**
Levamisole alone usually results in sufficient animal immobilization to image PercevalHR in the AC allowing precise alignment between the ATP-bound and ADP-bound channels. However, if animals are moving during image acquisition, additional immobilization is necessary. Physical fixation with polystyrene beads can be used in combination with levamisole to achieve more complete immobilization. It is crucial to ensure that the polystyrene bead preparation does not contain sodium azide as a preservative, as this will impact cellular ATP:ADP ratios and alter PercevalHR readouts. If animals still move after treatment with levamisole, we recommend using 1 μL of Polysciences Polybead.

Figure 6. Examples of ideal and insufficient ADP-bound channel signal, related to troubleshooting problem 3 (A) Strong PercevalHR fluorescence signal in the ADP-bound (405 nm laser) channel of ≥1.5-fold signal-to-background intensity results in accurate ATP:ADP ratio determination in the AC. (B) Weak PercevalHR fluorescence signal in the ADP-bound (405 nm laser) channel of ≤1.5-fold signal-to-background intensity can result in artificially high ATP:ADP ratios in the AC. Scale bars represent 5 μm.
microspheres 0.10 μm (catalog #00875-15) on the agar pad before adding the worms. This bead preparation does not contain sodium azide, and we have validated that it does not impact PercevalHR readouts compared to levamisole anesthetic alone. Alternatively, cyanoacrylate glue may be used to immobilize animals without anesthetic (Goodman et al., 1998), however we have not yet tested it with animals expressing PercevalHR in the AC.

RESOURCE AVAILABILITY

Lead contact
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Materials availability
There are restrictions to the availability of C. elegans codon-optimized PercevalHR due to the reagent being licensed by Calico Life Sciences LLC. The plasmid encoding C. elegans codon-optimized PercevalHR can be obtained directly from Calico Life Sciences.

Data and code availability
This study did not generate/analyze datasets or code.

SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.xpro.2022.101429.

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AUTHOR CONTRIBUTIONS
Conceptualization, A.G. and D.R.S.; methodology, A.G.; formal analysis, A.G.; investigation, A.G.; writing – original draft, A.G. and D.R.S.; writing – review and editing, A.G. and D.R.S.; visualization, A.G.; funding acquisition, D.R.S.; resources, D.R.S.; supervision, D.R.S.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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