Here, we present a protocol to characterize zebrafish LGP2 as a dual regulator of interferon (IFN) response. We detail in vivo assays using time-lapse comparison of IFN response between wild-type and lgp2 knockout zebrafish following spring viraemia of carp virus (SVCV) infection. We also describe in vitro assays including titration of infection duration in SVCV-infected fish cells to determine changes in IFN response. This protocol is effective to illuminate a regulatory switch of LGP2 in fish cells toward virus infection.

Publisher’s note: Undertaking any experimental protocol requires adherence to local institutional guidelines for laboratory safety and ethics.
Protocol
A protocol to characterize zebrafish LGP2 as a dual regulator of IFN response during viral infection

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SUMMARY
Here, we present a protocol to characterize zebrafish LGP2 as a dual regulator of interferon (IFN) response. We detail in vivo assays using time-lapse comparison of IFN response between wild-type and lgp2 knockout zebrafish following spring viraemia of carp virus (SVCV) infection. We also describe in vitro assays including titration of infection duration in SVCV-infected fish cells to determine changes in IFN response. This protocol is effective to illuminate a regulatory switch of LGP2 in fish cells toward virus infection.
For complete details on the use and execution of this protocol, please refer to Gong et al. (2022).1

BEFORE YOU BEGIN
LG2P (Laboratory of genetics and physiology 2) lacks the N-terminal CARDs (caspase activation and recruitment domains) that are required for signaling transduction in RIG-I (Retinoic acid inducible gene-I) and MDA5 (melanoma differentiation associated gene 5). In mammals, LGP2 is initially identified as a feedback inhibitor of RIG-I/MDA5 signaling but subsequently as an activator of MDA5 signaling and an inhibitor of RIG-I signaling.2 We previously established that zebrafish LGP2 is a dual regulator of fish IFN response toward viral infection.1,3 The protocol below describes the detailed steps in zebrafish larvae and adults as well as in cultured fish cells to determine LGP2’s dual regulation of fish IFN response. Before proceeding, it is imperative to generate lgp2-deficient zebrafish and validate a normal IFN response of cultured fish cells when infected with virus. The key point is to preestablish titration methods, including intensive titration of viral titer and infection duration, the doses of diverse stimuli, such as poly(I:C) and even LGP2 itself. Given the conservation and disparities of zebrafish and human LGP2, this protocol can be used in cultured mammalian cells, including HEK293T and COS7 cells, with a minor modification.4

Institutional permissions
Animal experiments and treatments were performed according to the Guide for Animal Care and Use Committee of Institute of Hydrobiology, Chinese Academy of Sciences (IHB, CAS, Protocol No. 2016–018).
Culture fish cell lines

Timing: 1 day

1. Grow epithelioma papulosum cyprini cells (EPC) in medium 199 (Gibco) supplemented with 10% fetal bovine serum (FBS) at 28°C in a humidified incubator containing 5% CO₂.

**CRITICAL:** Cell culture must be undertaken in microbiological safety cabinet using aseptic technique to ensure sterility. EPC cells are used here due to relatively high transfection efficiency and the fact that RLR-triggered IFN response can be easily detected when EPC cells respond to SVCV infection. EPC cells can be replaced by other fish cell lines with similar characteristics, such as grass carp C. idellus ovary cells (CO), and crucian carp (C. auratus) blastulae embryonic cells (CAB) in our lab.

Prepare SVCV solution

Timing: 7–10 days

2. Propagate SVCV.
   a. Seed EPC cells in 25 cm² cell culture flasks to reach monolayers after overnight (8–14 h) culture.
   b. Pour off medium carefully and wash cells with prewarmed serum-free M199 medium or PBS.
   c. Inoculate virus by pouring 0.5 mL $1 \times 10^{3}$ TCID₅₀/mL (tissue culture infective dose 50% per mL) SVCV to the culture flask for 20–30 min.
   d. Roll flask gently to ensure that all cells contact SVCV during inoculation.
   e. Pipette out SVCV and wash cells again with prewarmed serum-free M199 medium.
   f. Add 4–5 mL prewarmed M199 medium containing 0.5%–2% FBS, and place flasks in a 28°C incubator for at least 2–3 days.
   g. Freeze and thaw cells three times to thoroughly release virus when > 80% cytopathic effects (CPEs) appear.
   h. Move all lysates to 50 mL aseptic centrifuge tubes after mixing well by pipettes.
   i. Centrifugate at 13,000 g for 15 min.
   j. Sterilize supernatants through 0.22 μm membrane filter and store at –80°C.

3. Titrate SVCV virus.
   a. Put EPC cells in 96-well plates (100 μL/well) overnight (8–14 h) to reach a monolayer.
   b. 10-fold serially dilute SVCV virus in 96-well plates or microcentrifuge tubes. Make sure you re-suspend the dilutions very well before going to the next dilution.
   c. Pour off medium carefully, and wash cells with prewarmed serum-free M199 medium or PBS.
   d. Transfer virus dilutions to 96-well plates, with 8 wells each gradient and 100 μL each well.
   e. Leave the plates in a 28°C incubator for 5–7 days to ensure viral propagation.
   f. Check daily under an inverted microscope.
   g. Stain cells by dropping 1% crystal violet solution for 1–2 h.
   h. Discard the dye solution and calculate TCID₅₀/mL.

**CRITICAL:** A work with virus is biological safety level II. Accordingly, safety equipment, such as laboratory gloves, coats, gowns, shoe covers, boots, respirators, face shields, safety glasses, or goggles must be worn for this genetic safety level. However, SVCV is an aquatic virus that cannot infect human cells. SVCV solution with a titer of >10⁸ TCID₅₀/mL is conductive to the follow-up infection experiments.

Breed WT (lgp2+/+) or lgp2⁻/⁻ zebrafish

Timing: 1 week
4. Keep zebrafish in a circulation system, which continuously filters and aerates the system water.
   a. Store aeration water in the large water tank to supplement the loss of system water.
   b. Maintain the room temperature (RT) to be between 27°C–28°C.
   c. Set the system water temperature to be between 28°C–28.5°C, and the lighting conditions at 14:10 h (light:dark).

5. Transfer mature WT and lgp2−/− zebrafish into two 3-L tanks for breeding at the night before embryo collection. The ratio of females to males is 2:1 or 1:1.

△ CRITICAL: Zebrafish husbandry, maintenance and breeding are performed according to “Zebrafish Methods and Protocols 3.1.” It is recommended to feed the fish an extra meal 1 h before transferring them to breeding tanks. Transfer right after eating probably increases waste products in the water, which will decrease breeding efficiency.

6. Turn on the lights in the next morning, and you can see that females are chased by males for breeding under the stimulation of light.

△ CRITICAL: To facilitate stimulation, it is recommended to place a partition in the middle of tanks to separate females from males the night before embryo collection, and pull it out the next morning.

7. Collect and mix all embryos from similar clutches within 3 h post mating.
8. Wash embryos with system water to remove debris.
9. Transfer about 200 embryos to a 10-cm dish filled with fresh system water in a 28°C incubator. The incubator provides physiological light/dark cycle to ensure normal development.

△ CRITICAL: About 1,000 embryos are enough for further experiments. In our lab, sometimes we add methylene blue (10 mg/L) to limit fungal growth when incubating embryos, and wash the larvae by system water to remove methylene blue before experiments.

10. Remove dead embryos daily to keep the system water clean. The embryos develop into larvae 3 days post fertilization (dpf).

△ CRITICAL: Dead embryos are easily identified by their opaque or cloudy appearance. Generally, the unfertilized embryos are stained by methylene due to the changes in membrane permeability.

11. Perform viral-infection assays with zebrafish larvae or adults.

Prepare buffers

⊙ Timing: 1–2 h

12. Make 1% crystal violet solution.
13. Make 1× Tris-Gly working buffer (inside and outside inner modules), TBST, transfer buffer, 5% milk according to the recipe provided below.
14. Make 8% native-PAGE gels according to the recipe provided below.

Prepare sampling tools

⊙ Timing: 4–5 h

15. Bake animal dissection forceps, animal anatomical scissors and ceramic beads in an oven at 240°C for 4 h to inactivate the residue RNase.
# KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| CaIRF3 Rabbit mAb  (1:5000) | Sun et al. ¹ | N/A |
| Zebrafish LGP2 Rabbit mAb (1:5000) | Gong et al.³ | N/A |
| β-Actin Rabbit mAb (1:5000) | Cell Signaling Technology | Cat#: B457; RRID: AB_10950489 |
| HRP Goat Anti-Rabbit IgG (H+L) (secondary antibody) (1:5000) | ABclonal, China | Cat#: A5014, RRID: AB_2769854 |
| **Bacterial and virus strains** | | |
| Spring viraemia of carp virus (SVCV) | Gong et al.³ | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| Protease inhibitor cocktail | Bimake | Cat#: B14002 |
| Phosphatase inhibitor cocktail | Bimake | Cat#: B15001 |
| poly(l) C | SIGMA | Cat#: P0913; CAS: 42424-50-0 |
| RNA later RNA stabilization reagent | QIAGEN | Cat#: 76106 |
| Methylene blue | Solarbio, China | Cat#: M8030 |
| MS-222 | Alfa | Cat#: 1809123 |
| Crystal violet stain solution (1%) | Solarbio, China | Cat#: G1062 |
| Penicillin-streptomycin (100X), sterile | MedChemExpress | Cat#: HY-K1006 |
| RIPA lysis buffer (weak) | Beyotime, China | Cat#: P0013D |
| Non-denaturing PAGE loading buffer (5X) | Beyotime, China | Cat#: P0016 |
| Sodium deoxycholate | Macklin, China | Cat#: D6128; CAS: 302-95-4 |
| Ammonium persulfate | Macklin, China | Cat#: A801034; CAS: 7277-54-0 |
| 1 M Tris-HCl (pH=8.8) | Biosharp, China | Cat#: BLS15B |
| 1 M Tris-HCl (pH=6.8) | Biosharp, China | Cat#: BLS14A |
| 30% Acrylamide-bisacrylamide (29:1) | Biosharp, China | Cat#: BLS13B |
| N, N’, N’-Tetramethylethylenediamine (TEMED) | Beyotime, China | Cat#: ST7278 |
| Glycin | Genview | Cat#: FG149 |
| Tris-base | Biofroxx | Cat#: 1115KG001; CAS: 77-86-1 |
| Polyethyleneimine, linear (MW 25,000) (PEI) | Sigma-Aldrich | Cat#: 23966-1 |
| PBS | Monad | Cat#: CR10601S |
| Tween-20 | DING GUO, China | Cat#: DH358-4 |
| Clarity Western ECL substrate | BIO-RAD | Cat#: 1705061 |
| Methanol | SCR, China | Cat#: 40064260; CAS: 67-66-1 |
| Non-fat powdered milk | BBI, China | Cat#: A600669 |
| medium 199 (Gibco) | HyClone | Cat#: SH30253.01 |
| Fetal bovine serum (FBS) | WISENT | Cat#: 086-150 |
| Opti-mem | Invitrogen | Cat#: 31985-070 |
| **Critical commercial assays** | | |
| Endo-free Plasmid Mini Kit II | OMEGA | Cat#: D6950-02 |
| SV Total RNA isolation | Promega | Cat#: Z3100 |
| TRizol Plus RNA Purification kit | Invitrogen | Cat#:12183555 |
| GoScript™ Reverse Transcription System | Promega | Cat#: A5001 |
| Universal Blue qPCR SYBR Green Master Mix | YEASEN, China | Cat#: 11201508 |
| Dual-Luciferase® Reporter Assay System | Promega | Cat#: P0913 |
| **Deposited data** | | |
| Raw and analyzed data | This paper | Lead contact, Yibing Zhang (ybzhang@ihb.ac.cn) |
| Experimental models: Cell lines | | |
| Epithelioma papulosum cyprini cells (EPC) | ATCC (CRL-2872) | N/A |
| Ovary cells of grass carp (CO) | Kept in IHB, CAS | N/A |
| Crucian carp (C. auratus L.) blastulae embryonic cells (CAB) | Chen et al.⁷ | N/A |

(Continued on next page)
MATERIALS AND EQUIPMENT

### Reagent or Resource

| Experimental models: Organisms/strains |
|----------------------------------------|
| Zebrafish (Danio rerio) strain AB: larvae (6 pdf) and adults (60 dpf, male: female = 1:1) | China Zebrafish Resource Center | N/A |

| Recombinant DNA |
|-----------------|
| pcDNA3.1(+) | Invitrogen | Cat# V79020 |
| DrIFN-1pro-luc (expression plasmid) | Sun et al. | N/A |
| DrLGP2 (expression plasmid) | Gong et al. | N/A |
| pRL-TK (expression plasmid) | Promega | Cat# E2241 |

| Software and algorithms |
|-------------------------|
| GraphPad Prism 8.0 | Dotmatics | https://www.graphpad.com/ |

| Other |
|-------|
| Mini-PROTEAN® tetra system | BIO-RAD | Cat# 1658004 |
| Mini Trans-Blot® Electrophoretic Transfer Cell | BIO-RAD | Cat# 170-3930 |
| Mini-PROTEAN® system casting stand | BIO-RAD | Cat# 1658050 |
| LUMINESCENT IMAGE ANALYZER | LISTED | Model: ImageQuant LAS 4000 mini |
| 24-well plate | LABSELECT, China | Cat# 11320 |
| RNase-free microcentrifuge tubes | BBI, China | Cat# F611541-0001 |
| Animal dissection forceps | Fine Science Tools | Cat# 11295-10 |
| Animal anatomical scissors | Easybio, China | CAT#BE6621 |
| Ceramic beads, 1.4 mm (325 g) | QIAGEN | CAT#13113-325 |

### 10× tris-glycine (Tris-Gly) stock buffer

| Reagent | Final concentration | Amount |
|---------|---------------------|--------|
| Tris-base | 250 mM | 30.2 g |
| Glycine | 2 M | 144 g |
| ddH₂O | N/A | Add to 1 L |
| Total | N/A | 1 L |

Store at room temperature (20°C–25°C) for at least half year.

### 1× Tris-Gly working buffer (inside tank)

| Reagent | Final concentration | Amount |
|---------|---------------------|--------|
| 10× Tris-Gly stock buffer | 1× | 100 mL |
| Sodium deoxycholate | 0.2% | 2 g |
| ddH₂O | N/A | Add to 1 L |
| Total | N/A | 1 L |

Store at room temperature for at least half year.

### 1× Tris-Gly working buffer (outside tank)

| Reagent | Final concentration | Amount |
|---------|---------------------|--------|
| 10× Tris-Gly stock buffer | 1× | 100 mL |
| ddH₂O | N/A | Add to 1 L |
| Total | N/A | 1 L |

Store at room temperature for at least half year.
### Native-PAGE gel (8% separation gel)

| Reagent                               | Final concentration | Amount  |
|---------------------------------------|----------------------|---------|
| ddH₂O                                 | N/A                  | 2.3 mL  |
| 30% acrylamide-bisacrylamide (29:1)   | 8.1%                 | 1.35 mL |
| 1 M Tris-HCl (pH=8.8)                 | 250 mM               | 1.25 mL |
| 10% Ammonium Persulfate (APS)*        | 0.1%                 | 50 μL   |
| TEMED                                 | 0.06%                | 3 μL    |
| Total                                 | N/A                  | About 5 mL |

Mix well, pour into the rubber plate quickly and avoid bubbles. Add ddH₂O slowly for liquid sealing. Store in fume hood for 30–60 min. Carry out the entire process in fume hood. *10% APS: 1 g Ammonium persulfate in 10 mL ddH₂O. Store 10% APS at −20°C for three months, or at 4°C for a week.

### Native-PAGE gel (5% stacking gel)

| Reagent                               | Final concentration | Amount |
|---------------------------------------|----------------------|--------|
| ddH₂O                                 | N/A                  | 1.35 mL |
| 30% acrylamide-bisacrylamide (29:1)   | 5%                   | 335 μL |
| 1 M Tris-HCl (pH=6.8)                 | 125 mM               | 250 μL |
| 10% APS                               | 0.1%                 | 20 μL  |
| TEMED                                 | 0.1%                 | 2 μL   |
| Total                                 | N/A                  | About 2 mL |

Mix well, pour into the rubber plate quickly and avoid bubbles. Insert the comb quickly. Store in fume hood for 30–60 min. Carry out the entire process in fume hood.

### Transfer buffer

| Reagent     | Final concentration | Amount |
|-------------|---------------------|--------|
| Tris-base   | 26 mM               | 3.13 g |
| Glycine     | 200 mM              | 14.4 g |
| Methanol    | 20%                 | 200 mL |
| ddH₂O       | N/A                 | Add to 1 L |
| Total       | N/A                 | 1 L    |

Mix well, store at room temperature for at least half year.

### 5% milk

| Reagent     | Final concentration | Amount  |
|-------------|---------------------|---------|
| Non-fat powdered milk | 5%     | 5 g     |
| TBST        | N/A                 | Add to 100 mL |
| Total       | N/A                 | 100 mL  |

Mix well, store at 4°C for <12 h.

### TBST buffer

| Reagent     | Final concentration | Amount |
|-------------|---------------------|--------|
| Tris-base   | 25 mM               | 3.03 g |
| NaCl        | 137 mM              | 8 g    |
| KCl         | 2.7 mM              | 0.2 g  |
| Tween-20    | 0.1%                | 1 mL   |
| ddH₂O       | N/A                 | Add to 1 L |
| Total       | N/A                 | 1 L    |

Mix well, store at room temperature for <12 h.
CRITICAL: Sodium deoxycholate is low toxic, irritant and harmful when swallowed. Appropriate protective clothing, gloves and masks are necessary. 30% Acrylamide-Bisacrylamide has a certain neurotoxicity to the human body, and APS belongs to the fifth category of dangerous goods, which are irritating and corrosive to the skin and mucosa. It should be performed in the fume hood, and lab coats and disposable gloves are necessary to wear. TEMED is volatile, flammable, corrosive and neurotoxic. Please avoid inhaling in large quantities, remember to close the bottle cap after use, keep away from the source of fire and wear appropriate protective clothing, gloves and eye/face protection. In case eyes are contaminated, please rinse immediately with plenty of water and consult a doctor.

**STEP-BY-STEP METHOD DETAILS**

**Susceptible comparison of WT and \( \text{lgp2}^{+/+} \) zebrafish larvae to SVCV infection**

**Timing:** 2–3 days

To determine the biological role of LGP2 in zebrafish survival against virus infection, WT (\( \text{lgp2}^{+/+} \)) and \( \text{lgp2}^{-/-} \) zebrafish larvae are directly immersed with SVCV in two 10-cm dishes, respectively. Dead larvae are easily distinguished from live larvae due to severe body curling, which is beneficial to illustrate by taking pictures (Figure 1).

1. Sterilize the aerated water (system water) by 0.22 µm filter unit.
2. Take \( \text{lgp2}^{+/+} \) or \( \text{lgp2}^{-/-} \) zebrafish larvae into two containers with system water.
   a. Absorb methylene blue-containing water as far as possible.
   b. Wash zebrafish larvae with sterilized system water for 2–3 times to remove residual methylene blue.
3. Transfer 50 \( \text{lgp2}^{+/+} \) larvae and 50 \( \text{lgp2}^{-/-} \) larvae to two 10-cm dishes, respectively. Each dish contains 20 mL of system water supplemented with 200 µL penicillin-streptomycin solution (final concentrations: 100 U/mL of penicillin, and 100 mg/mL of streptomycin sulfate) to prevent the growth of bacteria.
4. Mix SVCV solution to reach a viral titer of \( 2 \times 10^5 \text{TCID}_{50}/\text{mL} \).
5. Place dishes at 28°C for 2 days through checking every few hours.

△ CRITICAL: SVCV infection duration can be adjusted in terms of real-time mortality, which is tightly related to the final viral titers. In our experiments, the work titer of SVCV is between \( 10^5-10^6 \text{TCID}_{50}/\text{mL} \).
6. Add MS-222 (10 g/100 mL) at a ratio of 1/1,000 to anesthetize \( lgp2^{+/+} \) and \( lgp2^{-/-} \) larvae, which will stop fish swimming and keep stationary within 1 min after anesthetization.

7. Take pictures as soon as possible.

△ CRITICAL: SVCV solution in M199 might make system water become red and has an impact on picturing. It is recommended to replace the virus solution with clean aerated water before anesthetizing.

**Death curve of \( lgp2^{+/+} \) and \( lgp2^{-/-} \) zebrafish larvae in response to SVCV infection**

⊙ Timing: 6–7 days

This section describes how to draw a death curve of zebrafish larvae by survival comparison of WT and \( lgp2^{-/-} \) zebrafish larvae immersed with SVCV.

8. Transfer 100 \( lgp2^{+/+} \) and \( lgp2^{-/-} \) larvae to two 10-cm dishes, respectively. Each dish contains 20 mL of system water, supplemented with 200 \( \mu \)L penicillin-streptomycin (final concentrations: 100 U/mL of penicillin, and 100 mg/mL of streptomycin sulfate).

9. Mix SVCV solution to reach a viral titer of \( 5 \times 10^5 \) TCID\(_{50}\)/mL.

10. Put dishes at 28°C. Check every few hours and pick out dead larvae immediately.

11. Record deaths in the two groups every 24 h until one group of larvae is completely dead.

12. Draw the death curve by data processing tools such as Prism 8.0.

△ CRITICAL: Infection duration is tightly related to the final titer of SVCV virus. In our experiments, the final titer of SVCV is between \( 10^5 \)–\( 10^6 \) TCID\(_{50}\)/mL. The dead larvae should be picked out immediately to avoid water contamination.

**IFN response comparison of \( lgp2^{+/+} \) and \( lgp2^{-/-} \) zebrafish larvae to SVCV infection**

⊙ Timing: 4–5 days

In response to SVCV infection, IFN response is significantly induced in both \( lgp2^{+/+} \) and \( lgp2^{-/-} \) zebrafish larvae by upregulation of \( ifn \) and ISGs. The subtle differences can be detected by time lapse comparison of mRNA levels of \( ifn \) and ISGs between \( lgp2^{+/+} \) and \( lgp2^{-/-} \) zebrafish larvae (6dpf) following SVCV infection.

13. Take 300 \( lgp2^{+/+} \) or \( lgp2^{-/-} \) zebrafish larvae to two 10-cm dishes, respectively. Each dish contains 20 mL of sterilized system water, supplemented with 200 \( \mu \)L penicillin-streptomycin solution (final concentrations: 100 U/mL of penicillin, and 100 mg/mL of streptomycin sulfate).

14. Add SVCV solution into two dishes up to a viral concentration of \( 5 \times 10^5 \) TCID\(_{50}\)/mL, and place dishes at a 28°C incubator.

15. Collect 30 \( lgp2^{+/+} \) larvae and 30 \( lgp2^{-/-} \) larvae, at 0, 3, 6, 12, 24, 36, 48, 72 h post infection, to two 1.5 mL RNase-free microcentrifuge tubes, respectively.

16. Remove the residual virus solution by washing with PBS twice.

17. Add 200 \( \mu \)L of RNAlater RNA Stabilization Reagent (Qiagen) into each tube, and store at \(-80^\circ\)C.

△ CRITICAL: To obtain replicable and stable expression patterns by follow-up real-time PCR, the sampling time points can be set more densely to ensure that there are enough sampling points in the early and later stages of SVCV infection.

18. Thaw all samples after sampling is finished. For each sample at a time point:
   a. Take all larvae out with dissection forceps.
   b. Absorb residual solution on a filter paper.
c. Put the sample into a fresh 1.5 mL RNase-free microcentrifuge tube.

19. Lyse samples:
   a. Add 1 mL of tissue lysis solution (according to different Kits. e.g., Trizol-based lysis buffer provided in TRizol Plus RNA Purification kit) together with several baked ceramic beads.
   b. Homogenize fully on a homogenizer. Or homogenize samples with other available methods.

20. Centrifuge at 15,000 × g for 10 min at 4°C.

21. Transfer supernatants carefully to fresh 1.5 mL microcentrifuge tubes.

22. Extract total RNA and remove cellular DNA by RNase-free DNase.

23. Synthesize first-strand cDNA by random primers or Oligo(dT) primers.

24. Detect mRNA levels of *ifn* and ISGs by real-time PCR.

25. Analyze data with data processing software such as Prism 8.0.

Death curve of *lgp2*+/+ and *lgp2*−/− zebrafish adults in response to SVCV infection

© Timing: 6–7 days

This section describes the steps to compare mortality rates between WT and *lgp2*−/− zebrafish adults in response to SVCV infection. Unlike *lgp2*+/+ and *lgp2*−/− zebrafish larvae which are easily immersed with SVCV solution in 10-cm dishes, *lgp2*+/+ and *lgp2*−/− zebrafish adults (60 dpf) are housed in two tanks at 28°C and infected with SVCV by intraperitoneal injection.

26. Take 50 *lgp2*+/+ zebrafish adults and 50 *lgp2*−/− zebrafish adults in two tanks at 28°C, respectively.

27. Infect fish by intraperitoneal injection with SVCV (10⁸ TCID₅₀/mL), 20 μL each.

△ CRITICAL: How to perform intraperitoneal injection is described previously. It is recommended to use equal amounts of males and females. All fish should be laid on the same day and housed at the same density to avoid differences in body size. It is best to keep zebrafish fasting for 12–24 h before injection. Virus titers can be adjusted appropriately relative to the weight of zebrafish.

28. Place tanks at 28°C.
   a. Check fish every few hours.
   b. Pick out dead fish immediately.

29. Record deaths every 24 h until one group is dead.

30. Draw the death curve by data processing software such as Prism 8.0.

△ CRITICAL: Infection duration is tightly related to the infected amounts of SVCV. Dead fish should be picked out immediately to avoid water contamination.

IFN response comparison of *lgp2*+/+ and *lgp2*−/− zebrafish adults to SVCV infection

© Timing: 4–5 days

IFN response is significantly induced by SVCV infection in zebrafish tissues by upregulation of *ifn* and ISGs. The subtle difference can be detected by time lapse comparison of mRNA levels of *ifn* and ISGs in different tissues of *lgp2*+/+ and *lgp2*−/− zebrafish adults (Figure 2).

31. Repeat steps 26–28.

32. Sample at 0, 3, 6, 12, 24, 36, 48, 72 h post infection, with 4 *lgp2*+/+ adults or 4 *lgp2*−/− zebrafish adults (male: female = 1:1) at a given time point.

33. Add MS-222 (10 g/100 mL) at a ratio of 1/1,000 to anesthetize fish, and snip the tail for blood-letting.
34. For each time point, collect gill, liver, spleen, head kidney, trunk kidney and other tissues of WT or lgp2/C0/C0 zebrafish adults (n=4) to a 200 μL RNase-free microcentrifuge tube, respectively.

35. Add 200 μL of RNAlater RNA Stabilization Reagent (Qiagen), and store at −80°C.

36. Repeat steps 18–25.

△ CRITICAL: To obtain replicable and stable expression patterns by real-time PCR, the sampling time points can be set more densely to ensure that there are enough sampling points in the early and later stages of SVCV infection.

Luciferase assays to determine IFN response by titration of poly(I:C)

△ Timing: 4–5 days

Transfection of fish cells with poly(I:C), a synthetic analogue of virus dsRNA, is believed to mimic virus infection. The concentrations of poly(I:C) correspond to cellular concentrations of virus dsRNA produced during virus infection: low concentrations of transfected poly(I:C) might be analogue to the intracellular concentrations of viral dsRNA at the early stage of virus infection and high concentrations correspond to the late stage of virus infection (Figure 3).

37. Seed EPC cells in 24-wells plates. Each well contain 0.5 mL M199 supplemented with 10% FBS, to reach monolayers after overnight (8–14 h) culture at 28°C.

38. Prepare transfection mixture.
   a. Mix three kinds of plasmids at a ratio of 10:10:1, DrIFN1pro-luc (200 ng), DrLGP2 (200 ng) and pRL-TK (20 ng).
   b. Dilute the indicated plasmids in 50 μL Opti-mem, which contain Polyethylenimine (PEI, MW25000, 1 μg/μL of storage concentration) with a ratio of 1:3 plasmid/PEI (μg/μL).

39. Add the prepared transfection mixtures into EPC cells, each well with 50 μL. Each concentration of poly(I:C) is performed in triplicate.

40. 24 h later, transfert cells again by PEI with poly(I:C) at increasing concentrations (0, 0.25, 0.5, 1, 1.5, 2, 4, 8, 10, 20, 40, 80 ng/mL).

41. Another 24 h later, collect cells for luciferase assays according to the Dual-Luciferase Reporter Assay System (Promega, USA).

△ CRITICAL: Cell culture must be undertaken in microbiological safety cabinet using aseptic technique to ensure sterility. Cells can be transfected by other available methods, such as Lipofectamine 2000. To obtain the replicable and stable change in stimulatory trends, the
concentration points of poly(I:C) need to be set more densely, particularly within the concentrations of <2 ng/mL in fish cells. Other fish IFN promoters can be used instead of DrIFN⁴₁pro-luc (zebrafish IFN⁴₁ promoter-driven luciferase plasmid).

Luciferase assays to determine IFN response by titration of LGP2 itself

@ Timing: 4–5 days

Overexpression of zebrafish LGP2 in fish cells provokes a strong IFN response at low doses but a weak one at high doses, indicating that the function switch of zebrafish LGP2 can be determined by luciferase assays through titration of zebrafish LGP2 itself (Figure 4). Similar dual regulation of human LGP2 can be detected in mammalian cells only when low concentration of poly(I:C) (such as 4 ng/mL) is present, based on below protocol with a minor modification by transfection again with low concentration of poly(I:C).

42. Seed EPC cells (each well containing 0.5 mL M199 supplemented with 10% FBS) in 24-wells plates to reach monolayers after overnight (8–14 h) culture at 28°C.
43. Prepare transfection mixture similar to step 38:
   a. Mix three kinds of plasmids: DrIFN⁴₁pro-luc (200 ng), pRL-TK (20 ng), together with DrLGP2 at increasing doses (0, 0.02, 0.1, 0.2, 0.5, 1, 2, 5, 10, 40, 100, 200 ng).
   b. Dilute these plasmids with 50 μL Opti-mem, which contains PEI with a ratio of 1:3 plasmid (μg)/PEI (μL).
44. Add the prepared transfection mixture into EPC cells, each well with 50 μL. Each dose of LGP2 is performed in triplicate.
45. 24–48 h later, collect cells for luciferase assays according to the Dual-Luciferase Reporter Assay System (Promega, USA).

△ CRITICAL: Cells can be transfected by other available methods, such as Lipofectamine 2000. To obtain replicable and stable inducible trends, the dose points of LGP2 can be set more densely, particularly within the doses of <10 ng in fish cells. Adequate empty vector pcDNA3.1(+) is added to guarantee equivalent amounts of total plasmid DNA transfected in all wells. Consistent results can be obtained when total doses of transfected plasmids are used at less than a threshold, such as <600 ng in 0.5 mL/well in 24-well plates.
Biochemical assays to determine LGP2’s dual regulation on IFN response

**Timing:** 4–5 days (for step 46)

SVCV infection triggers IRF3 dimerization in fish cells (Figure 5). In SVCV-infected EPC cells that are overexpressed with zebrafish LGP2, the dual regulation of zebrafish LGP2 on cellular IFN response can be determined by time course analyses of the change in IRF3 dimerization by native PAGE, and the change in cellular IFN and ISG expression by real-time PCR.

46. Seed EPC cells in 3.5-cm dishes to reach monolayers after overnight (8–14 h) culture at 28°C, each dish with 2 mL M199 supplemented with 10% FBS.

47. Prepare transfection mixture similar to step 38:
   a. Set DrLGP2 plasmids at increasing doses (0, 40, 1,000 ng).
   b. Dilute plasmids of each dose with 200 µL Opti-mem, which contains PEI with a ratio of 1:3 plasmid (µg)/PEI (µL).

48. Add the prepared transfection mixture into dishes, each dish with 200 µL.

49. Infect cells with SVCV at 24 h post transfection.
   a. Add SVCV solution to dishes to a final concentration at 5 × 10^5 TCID50/mL and mix gently.
   b. Put dishes at a 28°C incubator.

50. Sample at 0, 6, 12, 24, 36, 72 h post infection. Remove culture medium, and wash cells gently with cold PBS 2–3 times.

51. For real-time PCR analyses.
   a. Add cellular lysis buffer to dishes, 500 µL each.
   b. Transfer cell lysates to a 1.5 mL RNase-free microcentrifuge tube, and store at −80°C.
   c. Thaw all samples for RNA extraction according to SV Total RNA Isolation System Protocol (https://www.promega.com.cn/resources/protocols/technical-manuals/0/sv-total-rna-isolation-system-protocol/), after sampling is finished.
   d. Remove cellular DNA by treatment with RNase-free DNase.
   e. Synthesize first-strand cDNA by reverse transcription with random primers or Oligo(dT) primers.
   f. Detect mRNA levels of ifn and ISGs by real-time PCR.
   g. Analyze data with data processing software such as Prism 8.0.

52. For Native-PAGE analysis.
53. Detect IRF3 dimerization by native-PAGE electrophoresis.
   a. Prepare 8% native-PAGE gel according to the recipe provided above.
   b. Add 1 × Tris-Gly working buffer (inside and outside inner modular), and run for 30 min at 40 V before loading samples.
   c. Add 12.5 μL 5× non-denaturing PAGE loading buffer to each sample (50 μL lysates) on ice, and mix well.
   d. Load samples and run for 3 h at 80 V on ice.

   **CRITICAL:** Running electrophoresis at low temperature: place the electrophoresis tank in ice, and put an ice box inside the electrophoresis tank at the same time. If needed, replace with fresh ice boxes to sustain low temperature environment during electrophoresis.

   e. Using Bio-Rad transfer system, arrange sponge > filter paper > gel > 0.45 μm PVDF membrane > filter paper > sponge in transfer buffer orderly and thoroughly, and run for 50–80 min at 95 V on ice, to transfer proteins from gel to membrane.

   **Alternatives:** Besides wet transfer above, other transfer methods, such as dry transfer and semi-dry transfer, might be option.

   **CRITICAL:** Before transfer, put PVDF membrane into 100% methanol solution for 15 s, wash PVDF membrane with transfer buffer 1–2 times, and keep membrane wet in transfer buffer.

   f. Wash the membrane 3 × 10 min with TBST after transfer.
g. Block the membrane in TBST containing 5% milk for 30–60 min at RT (20°C–25°C) or 12 h at 4°C.

h. Wash the membrane 3 × 10 min with TBST.

i. Incubate the membrane with IRF3 antibody solution (1:5000 dilution in TBST supplemented with 1% milk, for 12 h at 4°C or 2 h at RT.

j. Remove antibody solution, wash the membrane 3 × 10 min using TBST.

k. Incubate the membrane in secondary antibody solution (1:5000 in TBST supplemented with 1% milk) at RT for 1–2 h.

l. Remove antibody solution, wash the membrane 3 × 10 min using TBST.

m. Do chemiluminescent visualization of protein bands by ImageQuant LAS 4000 mini system
   i. Make a solution by mixing equal volumes of enhancer solution and peroxide solution
      (Clarity Western ECL Substrate, BIO-RAD).
   ii. Drip the solution frequently on the top of membrane for 2 min at RT.
   iii. Capture image with a chemiluminescent imaging system, or other methods.

**CRITICAL:** Antibody specific to crucian carp IRF3 is described previously. Antibody dilution is dependent on the storage titer of a given antibody.

**EXPECTED OUTCOMES**

This protocol is designed to characterize the dual regulatory roles of zebrafish LGP2 in IFN response. Comparison of WT and knockout zebrafish in response to SVCV infection demonstrates that LGP2 is essential for zebrafish survival against viral infection (Figure 1). Real-time PCR analyses of *ifn* and ISG expression indicate that LGP2 acts as an activator of IFN response in the early phase of SVCV infection (<24 h post infection) and an inhibitor in the late phase of SVCV infection (>24 h post infection) (Figure 2). The function switch of zebrafish LGP2 can be observed in fish cells transfected with poly(I:C) at increasing concentrations by luciferase assays (Figure 3).

Given that overexpression of zebrafish LGP2 alone can stimulate IFN promoter activation, titration of LGP2 expression levels shows that low doses of LGP2 function as an activator, and high doses of LGP2 as an inhibitor (Figure 4). As a typical ISG, the change of LGP2 expression is tightly related to the different stages of virus infection; therefore, the assays regarding titration of LGP2 expression levels still indicate the dual regulation of LGP2 on fish IFN response. Finally, the dual regulation of zebrafish LGP2 can be confirmed by biochemical assays, such as time course analyses of the change in IRF3 dimerization by native PAGE, and the change in cellular IFN and ISG expression by real-time PCR.

**LIMITATIONS**

Zebrafish *ifn* and several ISGs are modulated in comparison of IFN response between WT and knockout zebrafish in this protocol. Transcriptome and proteome might be further performed at the same time points after SVCV infection to assist the understanding of the changes in *ifn* and ISG expression. In vitro assays are performed by overexpression of zebrafish LGP2 in fish cells. To avoid the interference from endogenous LGP2, these overexpression assays might be carried out in LGP2-deficient fish cells to further clarify LGP2 function.

**TROUBLESHOOTING**

**Problem 1**

*lgp2*+/− and *lgp2*−/− zebrafish adults do not lay eggs on the same day, or zebrafish embryos are not enough for follow-up infection assays (Refer to: Breed WT or *lgp2*−/− zebrafish).

**Potential solution**

Before feeding fish more frequently up to 3–4 times daily for 2–3 days, facilitating females to empty dead eggs by setting up breeding tanks and pre-breeding is helpful to improve breeding efficiency.
To avoid female dystocia due to over-feeding, feeding frequency and feeding duration can be adjusted in time. Breeding females are characterized by abdominal bulge, with yellow and translucent abdomen near the fecal orifice.

Problem 2
Infection assays fail to observe the differential mortality between \( lgp2^{+/+} \) and \( lgp2^{-/-} \) zebrafish due to dying too quickly (all dead <24 h post infection) or too slowly (starting to die >72 h post infection) (Refer to step-by-step method details: 8–12).

Potential solution
A pretest is recommended to obtain the optimized viral titer for infection assays under same conditions.

Problem 3
During viral infection assays, system water in dishes or tanks is contaminated by other microbes (Refer to step-by-step method details: 3–5, 8–10, 13–14).

Potential solution
System water is sterilized through 0.22 \( \mu \)m filter units before use. In addition, a supplement with penicillin and streptomycin is helpful to inhibit bacteria growth. Importantly, check frequently and pick up dead larvae immediately.

Problem 4
Real-time PCR or luciferase assays does not show a positive regulation of LGP2 on IFN response at the early infection and a negative regulation at the later infection (Refer to step-by-step method details: 23–24, 36, 41, 45).

Potential solution
The key point of this protocol is to utilize titration methods, including dense titration of viral titer and infection duration, the doses of diverse stimuli, such as poly(I:C) and even LGP2 itself. It is recommended to set viral titers, infection duration and stimuli doses more densely to optimize experiment parameters.

Problem 5
Overloading during plasmid transfection causes imperfect results (Refer to step-by-step method details: 43 and 47).

Potential solution
During transfection, adequate empty vector pcDNA3.1(+) is added to guarantee equivalent amounts of total plasmid DNA transfected in all wells. Consistent results can be obtained when total doses of transfected plasmids are used at less than a threshold, such as <600 ng in 0.5 mL/well in 24-well plates. The total doses of transfected plasmids are proportional to the number of fish cells seeded in different plates or dishes. For example, 4-fold doses of the plasmids are transfected in 6-well plates compared to that in 24-well plates, and a 3.5-cm dish corresponds to a well of 6-well plates.

Problem 6
Fish IRF3 dimerization is not shown by Native-PAGE assays (Refer to step-by-step method details: 52–53).

Potential solution
Collecting cells, lysing cells, loading samples and running electrophoresis are performed at 4°C. Avoid freezing and thawing of cellular lysates frequently, and use antibodies specific to fish IRF3.
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Yi-Bing Zhang (ybzhang@ihb.ac.cn).

Materials availability
The expression plasmids generated in this protocol are available from the lead contact.

Data and code availability
This paper does not report original code. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

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AUTHOR CONTRIBUTIONS
Y.-B.Z. conceived the project, and Y.-B.Z. and X.-Y.G. designed the experiments. X.-Y.G. performed the experiments. Y.-B.Z. and X.-Y.G. wrote the manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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