

# Mass marking of juvenile *Schizothorax wangchiachii* (Fang) with alizarin red S and evaluation of stock enhancement in the Jinping area of the Yalong River

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*Schizothorax wangchiachii* is a key fish species in the stock enhancement program of the Yalong River hydropower project, China. Alizarin red S (ARS) was used to mark large numbers of juvenile *S. wangchiachii* in the Jinping Hatchery and later used to evaluate stock enhancement in the Jinping area of the Yalong River. In a small-scale pilot study, 7,000 juveniles of the 2014 cohort were successfully marked by immersion in ARS solution, and no mortality was recorded during the marking process. The ARS mark in the fish otoliths remained visible 20 months later. In the large-scale marking study, approximately 600,000 juveniles of the 2015 cohort were successfully marked. Mortalities of both marked and unmarked juveniles were very low and did not differ significantly. Total length, wet mass and condition factor did not differ significantly between unmarked and marked individuals after three months. On 24 July 2015, about 840,000 Jinping Hatchery-produced young *S. wangchiachii*, including 400,000 marked individuals, were released at two sites in the Jinping area. Recapture surveys showed that 1) marked and unmarked *S. wangchiachii* did not differ significantly in total length, wet mass and condition factor; 2) stocked individuals became an important part of recruitment of the 2015 cohort; 3) instantaneous growth rate of marked individuals tended to slightly increase; and 4) most stocked individuals were distributed along a 10–15 km stretch near the release sites. These results suggest that the ARS method is a cost-efficient way to mass mark juvenile *S. wangchiachii* and that releasing juveniles is an effective means of stock recruitment.

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14 **Abstract**

15 *Schizothorax wangchiachii* is a key fish species in the stock enhancement program of the Yalong  
16 River hydropower project, China. Alizarin red S (ARS) was used to mark large numbers of  
17 juvenile *S. wangchiachii* in the Jinping Hatchery and later used to evaluate stock enhancement in  
18 the Jinping area of the Yalong River. In a small-scale pilot study, 7,000 juveniles of the 2014  
19 cohort were successfully marked by immersion in ARS solution, and no mortality was recorded  
20 during the marking process. The ARS mark in the fish otoliths remained visible 20 months later.  
21 In the large-scale marking study, approximately 600,000 juveniles of the 2015 cohort were  
22 successfully marked. Mortalities of both marked and unmarked juveniles were very low and did  
23 not differ significantly. Total length, wet mass and condition factor did not differ significantly  
24 between unmarked and marked individuals after three months. On 24 July 2015, about 840,000  
25 Jinping Hatchery-produced young *S. wangchiachii*, including 400,000 marked individuals, were  
26 released at two sites in the Jinping area. Recapture surveys showed that 1) marked and unmarked  
27 *S. wangchiachii* did not differ significantly in total length, wet mass and condition factor; 2)  
28 stocked individuals became an important part of recruitment of the 2015 cohort; 3) instantaneous  
29 growth rate of marked individuals tended to slightly increase; and 4) most stocked individuals  
30 were distributed along a 10–15 km stretch near the release sites. These results suggest that the  
31 ARS method is a cost-efficient way to mass mark juvenile *S. wangchiachii* and that releasing  
32 juveniles is an effective means of stock recruitment.

### 33 Introduction

34 Habitat degradation and overexploitation contribute to the decline of fisheries around the world.  
35 In order to improve fish stocks, stock enhancement by releasing hatchery-produced fish into wild  
36 habitats has been widely implemented (Brown & Day, 2002; Taylor et al., 2016; Yang et al.,  
37 2013). The effectiveness of stock enhancement can be assessed by mark-release-recapture studies  
38 (Blaxter, 2000), which require effective tagging or marking methods. Various marking  
39 techniques, such as otolith marking (Volk et al., 1999), coded wire tags (Bernard et al., 1998) and  
40 passive integrated transponders (Navarro et al., 2006), have been developed to monitor released  
41 fish. Among these techniques, otolith marking is a feasible method that allows long-term  
42 identification of small fish (Caraguel et al., 2014; Crook et al., 2009). Otolith marks can be  
43 achieved by fluorescent marking (Yang et al., 2016), thermal marking (Volk et al., 1999) and  
44 isotopic marking (Woodcock et al., 2011). The most popular marking protocol is to use  
45 fluorochromes, which can form chelate complexes with calcium ions that are built into skeletal  
46 and otolith structures (Poczyczyński et al., 2011). Fluorescent marks on the otolith are visible  
47 under a specific inducing laser because the calcium-fluorochrome complexes emit fluorescent  
48 light (Bashey, 2004; Taylor et al., 2005; Yang et al., 2016). Fluorochromes commonly used for  
49 otolith marking are alizarin red S (ARS), alizarin complexone, oxytetracycline hydrochloride and  
50 calcein. Compared with other fluorochromes, ARS offers better mark quality and lower cost and  
51 thus is viewed as a promising dye for mass marking fish at early life stages (Taylor et al., 2005;  
52 Yang et al., 2016).

53 In China, releasing hatchery-reared fish to enhance or restore fish stock abundance and fishery  
54 catches has been widely implemented for more than fifty years (Yang et al., 2013). Chinese carps  
55 and several other commonly cultured species that do not breed effectively in still waters were  
56 selected for early artificial rearing-releasing programs (Wu & Zhong, 1964; Liu, 1965). In recent  
57 decades, technical developments and advances in hatchery production have made it possible to  
58 breed considerable numbers of endemic and rare fish annually, including Chinese sturgeon  
59 *Acipenser sinensis* (Chang & Cao, 1999) and Chinese sucker *Myxocyprinus asiaticus* (Zhou et  
60 al., 2002). However, fish stock enhancement programs in China, particularly for freshwater fish  
61 species, have focused mainly on artificial propagation techniques and stocking scale, with little

62 attention paid to monitoring and evaluating the success of fish after release (Cheng & Jiang,  
63 2010; Yang et al., 2013; Zhang et al., 2003). Some marking technologies have been tested in  
64 different fish species in recent years, but rarely has large-scale marking and recapturing been  
65 used in stock enhancement (Zhang et al., 2003; Yang et al., 2013). Therefore, it is necessary to  
66 carry out post-release evaluation based on mass marking and recapturing.

67 *Schizothorax wangchiachii* (Fang, 1936), which belongs to the subfamily Schizothoracinae of  
68 the family Cyprinidae, is distributed mainly in the upper Yangtze River and its tributary, the  
69 Yalong River (Yue, 2000). This species is adapted to torrential mountain rivers in the  
70 southeastern Qinghai-Tibetan Plateau (Yue, 2000). Before the 1990s, *S. wangchiachii* was caught  
71 abundantly in many parts of its distribution range. However, its recruitment has declined  
72 dramatically since the mid-1990s, likely due to habitat degradation, overfishing and hydropower  
73 development (Deng et al., 2000; Duan et al., 1995; Jiang et al., 2007). To improve the health of  
74 the *S. wangchiachii* population in the Yalong River, conservation plans, such as building fish  
75 hatcheries, have been initiated (Wang et al., 2011).

76 Jinping Hatchery (28°18'39.09"N, 101°38'50.10"E; Fig. 1) is the first and most important fish  
77 hatchery located in the lower Yalong River. The hatchery is used to domesticate and propagate *S.*  
78 *wangchiachii* and many other fish species that are threatened by hydropower development in the  
79 Yalong River. Since 2011, annual release of *S. wangchiachii* (total length 40–80 mm) from  
80 Jinping Hatchery has been carried out in the Jinping area of the Yalong River (Deng et al., 2016).  
81 The objectives of this study were to assess the feasibility of mass marking *S. wangchiachii* using  
82 ARS and subsequently to evaluate the effectiveness of stock enhancement by recapturing marked  
83 individuals of the 2015 cohort after release.

## 84 **Materials and Methods**

### 85 **Experimental fish and fluorochrome**

86 Juveniles of *S. wangchiachii* used for marking and stocking in this study were produced at  
87 Jinping Hatchery using an artificial propagation technique. The breeding stocks were native  
88 spawners caught from the wild in the Jinping area of the Yalong River in 2011 and 2012.  
89 Juveniles were reared in numerous cylindrical tanks in the juvenile rearing room. These tanks are  
90 made of fiberglass, have a diameter of 2 m and a height of 1 m, and each has a temperature-

91 controlled water supply (water temperature, 15.5–17.5°C; dissolved oxygen concentration, 7.0–  
92 7.4 mg L<sup>-1</sup>; pH, 7.1–7.4) from the recirculating aquaculture system.

93 The fluorochrome ARS (C<sub>14</sub>H<sub>7</sub>NaO<sub>7</sub>S) used for marking was analytically purified powder.  
94 During immersion marking, ARS was dissolved directly in the rearing water according to the  
95 experimental design. To optimize marking quality and minimize juvenile mortality, several  
96 preliminary experiments were conducted in 2013, and results showed that immersing juvenile *S.*  
97 *wangchiachii* in water containing ARS doses ≤ 100 mg L<sup>-1</sup> for 24 h resulted in the lowest death  
98 rate while producing a mark that could be seen clearly in the otolith.

#### 99 **Small-scale marking pilot study**

100 A small-scale marking pilot study was performed in the rearing tank. Approximately 7,000 40-  
101 day-old juvenile *S. wangchiachii* of the 2014 cohort (total length 23.52 ± 1.50 mm, mean ± S.D.,  
102 n = 20) reared in a tank were selected for immersion marking on 5 May 2014. These juveniles  
103 were starved for 24 h prior to the treatment. The inner wall of the tank was carefully cleaned, and  
104 the rearing water was completely replaced with about 500 L of clean water. Thirty-five grams of  
105 ARS were pre-dissolved in about 10 L of water, which was immediately added to the tank. The  
106 juveniles were immersed in the ARS solution (70 mg L<sup>-1</sup>) for 24 h. During immersion, the  
107 solution was aerated continuously and the fish were not fed. After immersion was completed, the  
108 ARS solution was discharged into a sewage pool, and at the same time clean water was pumped  
109 into the tank to thoroughly rinse out the remnant dye. One day after immersion, dead individuals  
110 were counted and removed from the tank. To check the visibility and persistence of the marks, 10  
111 marked fish from the tank were haphazardly sampled and sacrificed with an overdose of MS-222  
112 (100 mg L<sup>-1</sup>) on 29 May 2014, 28 December 2014, 4 May 2015 and 24 January 2016. Sampled  
113 fish were kept in 100% ethanol until otolith examination.

#### 114 **Large-scale marking application**

115 From late April to early May 2015, five batches of juvenile *S. wangchiachii* (total length 20.85 ±  
116 1.41 mm, mean ± S.D., n = 140) of the 2015 cohort were marked at Jinping Hatchery using the  
117 ARS immersion protocol described above. In total, an estimated 600,000 fish were marked. To  
118 ensure the safety of the very small juveniles during immersion marking, the concentrations of  
119 ARS solution were controlled within a range of 30–50 mg L<sup>-1</sup>. One day after immersion was

120 completed, dead fish in each tank were counted and recorded. The mortalities of three batches of  
121 unmarked fish in the rearing room also were recorded. Five days after marking, both marked and  
122 unmarked fish were transferred to four outdoor fishponds in the hatchery. The fish were fed to  
123 satiation three times a day with a commercial artificial compound diet.

124 To assess effects of the marking process on growth, 100 marked and unmarked *S.*  
125 *wangchiachii* at similar daily age were haphazardly taken from two fishponds on 24 July 2015.  
126 These fish were starved for 24 h prior to further treatment. Afterwards, they were anaesthetized  
127 with MS-222 at a concentration of 100 mg L<sup>-1</sup>. Their total length and body length was measured  
128 to the nearest 0.01 mm with a digital caliper, and their wet mass was weighed to the nearest  
129 0.0001g with a precision electronic balance. To assess the mark effectiveness, 400 marked fish  
130 also were selected haphazardly from those in the four fishponds. They were anaesthetized with  
131 MS-222 at a concentration of 100 mg L<sup>-1</sup> and then stored in 100 % ethanol until otolith  
132 examination.

### 133 **Release and recapture**

134 On 24 July 2015, 840,000 young *S. wangchiachii* of the 2015 cohort, of which 400,000  
135 individuals were had been marked by ARS, were released at sites 2 and 3 in the Jinping area of  
136 the Yalong River (Fig. 1). Site 2 (28°18'47.80"N, 101°38'51.19"E) is located in the wide and  
137 deep part of the river (> 5 m maximum depth). Although it was not a suitable habitat for *S.*  
138 *wangchiachii*, site 2 was used because of stairs that provided access to the river. Site 3  
139 (28°19'41.32"N, 101°38'52.18"E) was near a sand quarry, and the substrate was covered with  
140 gravel and small stones: this was an appropriate habitat for young fish. The distance between the  
141 two release sites was about 2 km. The fish were first captured from each fishpond with a nylon  
142 trawl, and 150 marked individuals were haphazardly selected for measuring and weighing.  
143 Afterwards, fish were transferred to the release sites by a pickup truck and released into the river  
144 using buckets. For comparison of growth, about 10,000 unmarked *S. wangchiachii* were raised as  
145 a control group in a fishpond in Jinping Hatchery. They were fed twice daily with commercial  
146 feed at a ratio of about 3% of body weight per day.

147 Before the recapture surveys were conducted, appropriate recapture sites along the Jinping area  
148 of the Yalong River were chosen. Criteria included ease of fishing, suitable habitats for young

149 fish and the distances from the release sites. Seven sites, including the two release sites, along a  
150 60 km stretch of the Yalong River in the Jinping area were selected for the recapture surveys (Fig.  
151 1). Site 1 (28°17'46.38"N, 101°38'39.19"E) was a shallow riffle area (< 0.5 m in depth) with a  
152 substrate of gravel and small stones located about 1.5 km upstream of site 2 and 7 km  
153 downstream of Jinping Dam II. Site 1 was an important nursery ground for *Schizothorax* fish at  
154 early life stages. Site 4 (28°20'50.68"N, 101°39'18.04"E), site 5 (28°24'10.28"N,  
155 101°43'24.86"E), site 6 (28°27'41.38"N, 101°44'49.65"E) and site 7 (28°36'58.04"N,  
156 101°55'56.04"E) were located about 3, 15, 20 and 50 km downstream of site 3, respectively, and  
157 had a similar substrate of small stones and occasional boulders.

158 Recapture surveys were carried out at three-month intervals (in October 2015, January 2016  
159 and April 2016) using a similar method and sampling effort each time. At each site, three 9-m  
160 long fishing pots with 6-mm mesh were used for recapturing fish for four successive days, and  
161 catches in each pot were removed once a day. The fishing pot used in these surveys was a trap-  
162 type stationary fishing device that was especially suitable for catching small fish with total length  
163 < 20 cm. Electrofishing permitted by the Sichuan Municipal Bureau of Aquatic Products was  
164 performed only one time at each site, and it involved using a 30-cm-diameter anode and a 6-mm  
165 mesh landing net to sample for 40 min along the river. Specimens of *S. wangchiachii* assumed to  
166 be from the 2015 cohort based on personal experience of age-total length were sacrificed with an  
167 overdose of MS-222 and measured and weighed, whereas other fish were released. Meanwhile,  
168 50 individuals sampled from the hatchery control group also were measured. All sampled fish  
169 were stored in 100 % ethanol for further processing.

#### 170 **Otolith removal and examination**

171 Three pairs of otoliths were removed from all fish sampled, and the left three otoliths were  
172 mounted on glass slides using neutral balsam. To check the ARS mark and to read age, otoliths  
173 were observed under an Olympus BX40 fluorescence microscope fitted with a Q-Imaging  
174 MicroPublisher 5.0 RTV digital camera using the green laser and normal transmitted light (Yang  
175 et al., 2016).

#### 176 **Data analysis**

177 In this study, instantaneous growth rate of mean total length ( $G_t$ ) was calculated following Ricker

178 (1975) as:

$$179 \quad G_l = (\ln l_2 - \ln l_1) / 3$$

180 where  $l_1$  is the mean total length in millimeters of *S. wangchiachii* of at a given time point,  $l_2$  is  
181 the corresponding mean total length of the same batch three months later, and 3 is the  
182 sampling interval of three months.

183 Instantaneous growth rate of mean wet mass ( $G_w$ ) was calculated as:

$$184 \quad G_w = (\ln w_2 - \ln w_1) / 3$$

185 where  $w_1$  is the mean wet mass in grams of *S. wangchiachii* at a given time,  $w_2$  is the  
186 corresponding mean wet mass of the same batch three months later, and 3 is the sampling  
187 interval of three months.

188 The percent ( $P_i$ ) of marked individuals out of all marked *S. wangchiachii* for each recapture  
189 survey was calculated as:

$$190 \quad P_i = n_i / N_t$$

191 where  $n_i$  is the number of marked fish at site  $i$  of the given recapture date and  $N_t$  is the total  
192 number of marked fish of the given recapture.

193 The condition factor of *S. wangchiachii* was calculated following Fulton (1904) as:

$$194 \quad \text{condition factor} = 10^6 \times w / l^3$$

195 where  $w$  is the wet mass in grams of *S. wangchiachii* at a given time point and  $l$  is the total length  
196 in millimeters of *S. wangchiachii* at the same time.

197 ANOVA test for total length, wet mass and condition factor between marked and unmarked  
198 *S. wangchiachii* was performed with SPSS 19.0 software. The Bonferroni Tests were used for  
199 post hoc tests when the variances were equal, and alternatively, the Games-Howell Tests were  
200 used when the variances were unequal. The significance level was set as  $P < 0.05$ .

## 201 **Results**

### 202 **Small-scale marking pilot study**

203 One day after immersion was completed, no marked fish had died. Twenty-three days after  
204 marking, all sampled individuals showed a visible red-orange mark in their otoliths (Fig. 2a).

205 Without polishing, visible marks also were easily identified in the otoliths of marked individuals  
206 sampled on 28 December 2014 (Fig. 2b), 4 May 2015 (Fig. 2c) and 24 January 2016 (Fig. 2d).

207 More than one year after immersion, the ARS mark still remained visible, and there was no  
208 evidence that the mark was significantly decaying.

### 209 **Large-scale marking application**

210 In the large-scale immersion marking of juvenile *S. wangchiachii*, mortalities in each batch were  
211 very low ( $\leq 0.50\%$ ), and no significant difference in the mortality between marked and unmarked  
212 batches was detected (Table 1). Three months after being reared in outdoor fishponds, samples of  
213 marked and unmarked individuals showed no significant difference in total length, wet mass and  
214 condition factor (Table 1). A visible red-orange mark in the otoliths could be identified under  
215 green laser in all marked fish sampled.

### 216 **Recapture and evaluation**

217 Otolith checking confirmed that a total of 852 *S. wangchiachii* of the 2015 cohort were caught  
218 during the three recapture surveys. Of these fish, 262 individuals had a clear ARS mark in their  
219 otoliths, and 590 individuals had no mark (Fig. 3; Table 2). The percent of marked individuals in  
220 each recapture survey were 32.73% in October 2015, 26.36% in January 2016 and 30.53% in  
221 April 2016.

222 In the October 2015 recapture effort, total length, wet mass and condition factor of both  
223 marked and unmarked *S. wangchiachii* were significantly lower than those of the hatchery  
224 control group (Table 2; one-way-ANOVA with *post hoc* Games-Howell Test,  $P < 0.001$ ). The  
225 condition factor of marked individuals when they were recaptured was also significantly lower  
226 than when they were released (Table 2; independent t-test,  $P < 0.001$ ). Between marked and  
227 unmarked individuals, there was no significant difference in wet mass ( $P = 0.134$ ) and condition  
228 factor ( $P = 0.735$ ), whereas a slight difference in total length was detected ( $P = 0.030$ ). In  
229 addition, the total length, wet mass and condition factor differed significantly among samples  
230 caught from different recapture sites (Table 3; two-way-ANOVA test without the data for site 7,  
231  $P < 0.001$ ), but a significant difference was not observed between marked and unmarked  
232 individuals ( $P > 0.05$ ). Recapture site and ARS mark had interaction effects on wet mass ( $P =$   
233 0.006).

234 In the January 2016 recapture effort, the total length and wet mass of both marked and  
235 unmarked *S. wangchiachii* were still significantly lower than those of the hatchery control group

236 (Table 2; one-way-ANOVA with *post hoc* Games-Howell Test,  $P < 0.001$ ), but the difference in  
237 condition factor was not significant ( $P = 0.304$ ). Between marked and unmarked individuals, no  
238 significant differences in total length ( $P = 0.828$ ) and wet mass ( $P = 0.816$ ) were detected. The  
239 total length, wet mass and condition factor of *S. wangchiachii* differed significantly among  
240 samples caught from different recapture sites (Table 3; two-way-ANOVA test without the data for  
241 sites 1 and 7,  $P < 0.001$ ), but ARS mark and the interaction between it and recapture site did not  
242 have significant effects on the three indexes ( $P > 0.05$ ).

243 In the April 2016 recapture effort, the hatchery control group was not sampled because of  
244 fishpond cleaning. There was no significant difference in total length, wet mass or condition  
245 factor between unmarked and marked *S. wangchiachii* (Table 2; independent t-test,  $P > 0.05$ ).  
246 The total length, wet mass and condition factor of *S. wangchiachii* differed significantly among  
247 samples caught from different recapture sites (Table 3; two-way-ANOVA test, including only data  
248 for sites 3, 4 and 6,  $P < 0.001$ ). A significant difference in condition factor (Table 3;  $P = 0.009$ )  
249 was observed between marked and unmarked individuals, but a significant difference was not  
250 observed for total length ( $P = 0.181$ ) or wet mass ( $P = 0.07$ ). The interaction effects of recapture  
251 site and ARS mark were not significant ( $P > 0.05$ ).

252 The  $G_l$  and  $G_w$  of marked individuals tended to slowly increase after release. In the first  
253 trimester after release, the  $G_l$  and  $G_w$  values of marked individuals were 0.0566 and 0.1065,  
254 respectively, which were lower than those of the hatchery control group (0.1430 and 0.4964,  
255 respectively). In the second trimester, both  $G_l$  (0.0683) and  $G_w$  (0.2832) of marked individuals  
256 had increased slightly. At that time, the  $G_w$  of marked individuals was slightly higher than that of  
257 the hatchery control group (0.2569), whereas the  $G_l$  of marked individuals was still lower than  
258 that of the hatchery control group (0.1080). In the third trimester, the  $G_l$  of marked individuals  
259 had increased to 0.0878, whereas the  $G_w$  (0.2815) remained almost the same.

260 After being released, the hatchery-produced *S. wangchiachii* began to move away from the  
261 release area. In October 2015, marked individuals were recaptured at all recapture sites upstream  
262 and downstream of the release sites (Table 2). Site 2 was not suitable as nursery ground for *S.*  
263 *wangchiachii*, but  $P_2$  (18.90%) nevertheless represented a high percent of total recaptures (Fig.  
264 4). In the subsequent recaptures, catches at site 2 were very small because of unsuitable habitat

265 (Table 2). In the three surveys, mean  $P_3$  ( $28.69 \pm 1.03\%$ ) and  $P_4$  ( $31.62 \pm 25.98\%$ ) were much  
266 higher than that of  $P_5$  ( $11.49 \pm 7.80\%$ ),  $P_6$  ( $8.58 \pm 4.38\%$ ) and  $P_7$  ( $4.17 \pm 3.23\%$ ).  $P_i$  significantly  
267 decreased with distances from the release sites.  $P_i$  for the distant sites 6 and 7 in the latter two  
268 recaptures increased slightly compared to that in the first recapture, but the values were still much  
269 lower than those of sites 3–5 (Fig. 4). This implies that stocked fish were mainly distributed over  
270 a 10–15 km long stretch around the release sites.

## 271 Discussion

### 272 Feasibility of ARS mass marking

273 Previous studies of marking different fish species demonstrated that ARS treatment produces an  
274 excellent mark quality and has no significant harmful effects on the fish (Baer & Rösch, 2008;  
275 Bashey, 2004; Caraguel et al., 2014; Liu et al., 2009). However, faced with sustained pressure to  
276 produce enough fish seed to achieve the annual goals of release programs, managers of many  
277 hatcheries continue to worry that mass marking using the ARS method will cause high mortality,  
278 and this concern has a negative impact on the use of marking to evaluate fish stock enhancement.  
279 In this study, marking juvenile *S. wangchiachii* (mean total length  $23.52 \pm 1.50$  mm) by  
280 immersion in  $70 \text{ mg L}^{-1}$  ARS solution for 24 h did not cause death. In the following large-scale  
281 marking application, the mortality of five marked batches and three unmarked ones was  
282 negligible ( $\leq 0.50\%$ ), and no significant difference in mortality between marked and unmarked  
283 fish was detected. After rearing for three months in outdoor fishponds, no significant differences  
284 in total length, body mass, or condition factor between marked and unmarked groups were  
285 detected. Because juvenile *S. wangchiachii* experienced natural mortality, the extremely low  
286 mortalities that occurred during the immersion marking process might not have been due to ARS  
287 solution. In addition, immersion marking was carried out directly in the rearing tanks, which  
288 avoided manipulations of fish, reduced stress and its cost.

289 The ARS mark that develops in the otolith remains highly readable for several years, whether  
290 fish are reared in the laboratory or in the field (Champigneulle & Cachera, 2003; Nagiec et al.,  
291 1995; Partridge et al., 2009; Poczyczyński et al., 2011). Because sunlight and turnover of skeletal  
292 calcium can cause fluorescent marks to fade, external fluorescent marks on scales and fin rays  
293 cannot be readily detected over time (Bashey, 2004; Elle et al., 2010). In contrast, otoliths are

294 protected by the skull and previously deposited otolith materials are not resorbed (Campana &  
295 Neilson, 1985), which prolongs the lifetime of the mark. In the 2014 marking effort, *S.*  
296 *wangchiachii* marked with 70 mg L<sup>-1</sup> ARS retained highly visible marks on otoliths after rearing  
297 for about 20 months in an indoor tank, and they did not present clear signs of significant fading.  
298 In the mass marking of 2015, fish marked with 30–50 mg L<sup>-1</sup> ARS were transferred to outdoor  
299 fishponds and reared for about three months. Jinping Hatchery is located in the arid river valley  
300 region of the western Sichuan Province, where sunshine is very strong all year long (Yuan et al.,  
301 2013). Nevertheless, ARS marks of fish sampled from each marking batch were highly visible.  
302 All of the ARS marks on the otoliths of recaptured fish were as clear as they had been at the time  
303 of release. However, because otoliths continuously grow and thicken, over time the mark can be  
304 covered by otolith materials, and marks can become faint and difficult to detect unless exposed  
305 by grinding and polishing the otoliths (Baer & Rösch 2008; Sánchez-Lamadrid, 2001; Taylor et  
306 al., 2005). In this study, although it was not experimentally tested, the final retention time of the  
307 ARS mark in the otoliths of *S. wangchiachii* was long enough to monitor the released individuals  
308 to evaluate stocking effectiveness.

### 309 **Effectiveness of stocking enhancement**

310 After release, trimonthly recapture surveys confirmed that some of the stocked *S. wangchiachii*  
311 had survived. Assuming that the percent of marked (47.62%) and unmarked (52.38%) *S.*  
312 *wangchiachii* remained unchanged in the stocked cohort, the percent of catches that originated  
313 from stock enhancement were estimated to be 68.73 % in October 2015, 55.35% in January 2016  
314 and 64.11% in April 2016, respectively. This demonstrated that stocked *S. wangchiachii*  
315 constituted an important part of the young fish with a mean level of 62.73% recruitment of the  
316 2015 cohort. In previous successful stock enhancements, such as those for vendace *Coregonus*  
317 *albula* (Poczyczyński et al., 2011), brown trout *Salmo trutta* (Caudron & Champigneulle, 2009),  
318 Japanese Spanish mackerel *Scomberomorus niphonius* (Obata et al., 2008), and red sea bream  
319 *Pagrus major* and Japanese flounder *Paralichthys olivaceus* (Kitada & Kishino, 2006), hatchery-  
320 produced fish contributed considerably to population recruitment, and stocking successes were  
321 often attributable to appropriate release sizes and environmental conditions at release sites. In this  
322 study, the relatively high percent of released *S. wangchiachii* might be explained by the fact that

323 young fish for release were fully covered with scales, which would protect the skin against  
324 mechanical injury and bacteria and parasites (Yan et al., 2014). In addition, the negligible fishing  
325 pressure on young *S. wangchiachii* under age 2 and few predators, such as *Percocypris pingi* and  
326 *Silurus asotus*, might have had positive effects.

327 However, the comparative analysis of recapture data showed that total length, wet mass, and  
328 condition factor of recaptured *S. wangchiachii* differed significantly among different recapture  
329 sites (Table 3). The body sizes of fish recaptured at release sites were often smaller than those at  
330 other sites for both marked and unmarked fish (Table 2). This finding suggests that fish at release  
331 sites did not grow as well as fish at other sites. Pebbly nursery grounds in shallow waters are  
332 essential for *Schizothorax* fish at early life stages. In the Jinping area of the Yalong River, water  
333 flow sharply decreases (at a maximum percent of about 95%) due to the upstream dam of Jinping  
334 Dam II, which leads to marked physical habitat degradation (Wang et al., 2007). Excavation of  
335 sand in the river, which take place frequently at five sites along the 60 km long survey area,  
336 further destroyed the habitats. This reduction of essential habitats could have a significant  
337 negative effect on the river's carrying capacity for *Schizothorax* fishes. It is likely that the  
338 released fish moved very slowly so that nine months after release most of them still were  
339 distributed over a 10–15 km long stretch near the release sites, although a few marked fish were  
340 caught about 50 km downstream three months after release. The relatively slow migration speed  
341 would maintain a high density of fish in the release area, which would result in both released and  
342 wild fish having to compete intensely with each other for resources.

343 Wild fish can be replaced with hatchery-reared fish when the latter are released in numbers that  
344 exceed the carrying capacity, but it is difficult to verify the extent to which they replace the wild  
345 ones (Kitada & Kishino, 2006). The surveys conducted in this study showed that, there are some  
346 spawning grounds for *Schizothorax* fishes in the Jinping area of the Yalong River, where some  
347 naturally born juveniles were caught in April 2014. This study showed that natural recruitment  
348 still accounted for a sizeable percent (approximate 40%) of *S. wangchiachii* recruitment of the  
349 2015 cohort. Additionally, the non-significant difference in total length, wet mass and condition  
350 factor between the marked and unmarked fish indicated that ARS marking did not have  
351 significant harmful effects. This finding suggests that the stocked *S. wangchiachii* should grow as

352 the naturally born fish in the Jinping area of the Yalong River. Releases at the scale used in this  
353 study might have not exceeded the carrying capacity of release sites or replaced wild fish, but  
354 repeated large annual releases might do so. To maintain the fish population at sustainable levels,  
355 new release sites around site 7 should be added (Fig. 1). Moreover, sand excavation in the river  
356 should be stopped immediately, and nursery habitats must be restored to expand the carrying  
357 capacity of the river.

358 Wild *S. wangchiachii* live in rapid flowing river water and consume adherent alga  
359 (Bacillariophyta) using the sharp outer horny sheath on their lower jaw to scrape it off the  
360 substrate. In contrast, the hatchery-reared fish are reared in still water ponds and fed on  
361 commercial diets. It is well known that environmental differences between hatchery-reared and  
362 wild fish can influence their behavior, especially foraging behavior and avoidance of predators,  
363 which may subsequently affect post-release success (Hervas et al., 2010; Le Vay et al., 2007;  
364 Johnsson et al., 2014). On first release into the wild, hatchery-reared fish must not only avoid  
365 predators but also adapt to a new food supply (Blaxter, 2000). In this study, condition factors of  
366 the stocked *S. wangchiachii* recaptured three months after release were significantly lower than  
367 those of the hatchery control group as well as their own cohort at the time of release (Table 2).  
368 This suggests that these hatchery-produced fish might need to be acclimatized to the wild habitat,  
369 as their growth was negatively affected by the change of environmental conditions. Perhaps,  
370 hatchery-reared fish suffered a high level of short-term post-release mortality during the first  
371 trimester after release. However, it was difficult to precisely estimate this mortality due to lack of  
372 historical data on fisheries and prior investigation in the area. In the subsequent recaptures, the  
373 condition factor of stocked fish had returned to a level as good as that of the hatchery control  
374 group. In addition, the  $G_l$  and  $G_w$  of marked fish displayed a slowly increasing trend (Tables 2  
375 and 4). Therefore, survival of hatchery-produced *S. wangchiachii* suggests that they gradually  
376 adapted to the wild habitat, and they exhibited favorable growth six months after release.

### 377 **Conclusions**

378 This study offers fishery administrators a cost-efficient method of mass marking juvenile *S.*  
379 *wangchiachii* with ARS. The marking process did not cause significant mortality or affect fish  
380 growth in this study. Release-recapture surveys indicated that the present stock enhancement

381 might make a considerable contribution to the recruitment of young *S. wangchiachii* in the  
382 Jinping area of the Yalong River. Results of this study will be instrumental in promoting  
383 application of mass marking techniques and applying responsible approaches to the development  
384 of stock enhancement in China. However, much information about stock enhancement remains  
385 unknown, including the post-release mortalities of stocked fish, their contribution to the  
386 spawning population, and their genetic impact on the wild population. Therefore, in order to  
387 improve stocking strategy and better protect *S. wangchiachii* and other fish species in the Yalong  
388 River, long-term monitoring and further studies of the released fish should be conducted.

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**Table 1** (on next page)

The mortalities of juvenile *S. wangchiachii* unmarked and marked by immersion in late April to early May 2015, and total length, wet mass and condition factor of these fish three months later.

Mortality (mean and S.D.) of three batches of unmarked juvenile *S. wangchiachii* and five batches marked by immersion in 30–50 mg L<sup>-1</sup> ARS solution for 24 h in late April to early May 2015, was very low and did not differ significantly (independent t-test,  $P = 0.836$ ) between treatments. After being reared for three months in outdoor fishponds, total length ( $P = 0.936$ ), wet mass ( $P = 0.629$ ) and condition factor ( $P = 0.244$ ) (mean and S.D.) of marked and unmarked *S. wangchiachii* did not differ significantly ( independent t-test).

Samples	<i>N</i>	Acute mortality (%)	Total length (mm)	Wet mass (g)	Condition factor (%)
Marked batches	5	0.16(0.20)			
Unmarked batches	3	0.19(0.20)			
Marked individuals	100		49.17(9.76)	1.1460(0.6563)	1.715(0.175)
Unmarked individuals	100		49.07(7.16)	1.1052(0.5288)	1.748(0.212)

*N*, number of samples

**Table 2** (on next page)

Mean total length, wet mass and condition factor of marked and unmarked *S. wangchiachii* of the 2015 cohort.

These data were got at the time of initial release and from recapture surveys carried out in October 2015 and January and April 2016 at seven sites in the Jinping area of the Yalong River. The numbers in parentheses are the standard deviations.

Source	Marked fish				Unmarked fish			
	<i>N</i>	Total length (mm)	Wet mass (g)	Condition factor	<i>N</i>	Total length (mm)	Wet mass (g)	Condition factor
<b>Jul 2015</b>								
Initial release fish	150	44.94(6.67)	0.8225(0.3332)	1.762(0.373)				
Hatchery control fish					50	41.54(4.91)	0.6740(0.2786)	1.893(0.762)
<b>Oct 2015</b>								
Hatchery control fish					50	63.79(11.26)	2.9886(1.8327)	1.995(0.431)
Site 1	45	52.21(7.34)	1.0341(0.4257)	1.430(0.170)	69	54.61(6.72)	1.1551(0.4236)	1.447(0.193)
Site 2	31	53.02(5.61)	1.0889(0.4352)	1.472(0.115)	51	57.27(9.87)	1.5174(1.0875)	1.518(0.133)
Site 3	48	54.03(7.11)	1.1733(0.5013)	1.405(0.187)	118	55.88(6.96)	1.2401(0.4663)	1.349(0.151)
Site 4	6	59.30(6.51)	1.5886(0.6685)	1.400(0.075)	20	58.85(6.36)	1.4423(0.4303)	1.392(0.179)
Site 5	27	51.04(6.32)	0.9882(0.3255)	1.482(0.184)	68	51.54(5.61)	1.0309(0.3654)	1.481(0.195)
Site 6	6	59.88(13.19)	1.8959(1.3197)	1.511(0.158)	9	52.46(5.83)	1.1436(0.3253)	1.570(0.153)
Site 7	1	55.36	1.4478	1.835	2	53.58(3.76)	1.3499(0.3667)	1.618(0.047)
Total	164	53.26(7.23)	1.1320(0.5263)	1.443(0.169)	337	55.03(7.38)	1.2325(0.5939)	1.431(0.182)
<b>Jan 2016</b>								
Hatchery control fish					50	88.20(7.77)	6.4593(1.5860)	1.770(0.261)
Site 1	—				—			
Site 2	0				2	67.13(17.12)	2.2926(1.7039)	1.604(0.265)
Site 3	17	53.57(8.82)	1.3193(0.7693)	1.648(0.186)	36	56.87(7.22)	1.5691(0.7476)	1.676(0.157)
Site 4	21	67.51(15.57)	2.9274(2.1312)	1.622(0.212)	77	67.70(14.15)	2.9843(2.1267)	1.705(0.150)
Site 5	9	70.02(8.99)	3.1357(1.2962)	1.867(0.145)	17	74.55(9.04)	4.1380(1.6331)	1.989(0.245)
Site 6	7	75.37(9.24)	3.5619(1.4448)	1.749(0.118)	18	69.63(8.20)	2.9381(0.9454)	1.823(0.199)
Site 7	4	76.28(3.93)	4.1186(0.7443)	1.996(0.209)	12	72.64(9.62)	3.4644(1.6936)	1.814(0.162)
Total	58	65.37(13.98)	2.6471(1.7583)	1.709(0.214)	162	66.59(12.70)	2.8128(1.8472)	1.748(0.194)
<b>Apr 2016</b>								
Site 1	—				—			

Site 2	0				4	65.62(12.48)	2.7153(1.7478)	1.688(0.104)
Site 3	11	75.28(10.18)	3.8435(1.4643)	1.553(0.094)	20	77.58(14.38)	4.1291(3.2624)	1.450(0.173)
Site 4	22	85.22(17.52)	6.3145(4.1856)	1.653(0.125)	45	79.16(9.76)	4.5388(2.0997)	1.628(0.116)
Site 5	1	96.71	9.1327	1.934	11	99.50(15.42)	10.1536(4.5667)	1.805(0.143)
Site 6	4	104.02(5.54)	10.2389(1.4235)	1.764(0.126)	11	94.99(13.61)	7.7649(3.4934)	1.627(0.164)
Site 7	2	93.32(13.11)	7.5525(3.2873)	1.671(0.076)	0			
Total	40	85.06(16.42)	6.1598(3.7384)	1.644 ±0.134)	91	82.59(14.84)	5.4373(3.5597)	1.613(0.171)

–, no fish were captured because we were unable to get to the site

*N*, number of fish

**Table 3**(on next page)

The results of Two-way ANOVAs on the effects of recapture site (R) and ARS mark (A) and their interaction (R×A) on total length, wet mass and condition factor of *S. wangchiachii* of the 2015 cohort.

Original data were obtained from recapture surveys carried out in October 2015 and January and April 2016 at seven sites in the Jinping area of the Yalong River.

Source	Dependent variable	SS	df	F	P
<b><i>Oct 2015</i></b>					
Recapture site	Total length	1362.827	5	272.565	< 0.001
	Wet mass	8.040	5	1.608	< 0.001
	Condition factor	1.091	5	0.218	< 0.001
ARS mark	Total length	1.948	1	1.948	0.844
	Wet mass	0.086	1	0.086	0.596
	Condition factor	0.005	1	0.005	0.683
R × A	Total length	487.183	5	97.437	0.088
	Wet mass	5.087	5	1.017	0.006
	Condition factor	0.168	5	0.034	0.316
Error	Total length	24574.918	486		
	Wet mass	149.255	486		
	Condition factor	13.776	486		
<b><i>Jan 2016</i></b>					
Recapture site	Total length	8178.913	4	15.296	< 0.001
	Wet mass	118.057	4	10.572	< 0.001
	Condition factor	1.860	4	15.275	< 0.001
ARS Mark	Total length	2.128	1	0.016	0.900
	Wet mass	0.001	1	< 0.001	0.984
	Condition factor	0.019	1	0.612	0.435
R × A	Total length	433.993	4	0.812	0.519
	Wet mass	9.423	4	0.844	0.499
	Condition factor	0.218	4	1.792	0.132
Error	Total length	27805.420	208		
	Wet mass	580.665	208		
	Condition factor	6.331	208		
<b><i>Apr 2016</i></b>					
Recapture site	Total length	4442.714	2	13.549	< 0.001
	Wet mass	208.859	2	12.230	< 0.001
	Condition factor	0.476	2	13.454	< 0.001
ARS Mark	Total length	297.492	1	1.815	0.181
	Wet mass	28.595	1	3.349	0.070
	Condition factor	0.126	1	7.134	0.009
R × A	Total length	418.750	2	1.277	0.283
	Wet mass	25.205	2	1.476	0.233
	Condition factor	0.049	2	1.388	0.254

Error	Total length	17542.681	107
	Wet mass	913.667	107
	Condition factor	1.895	107

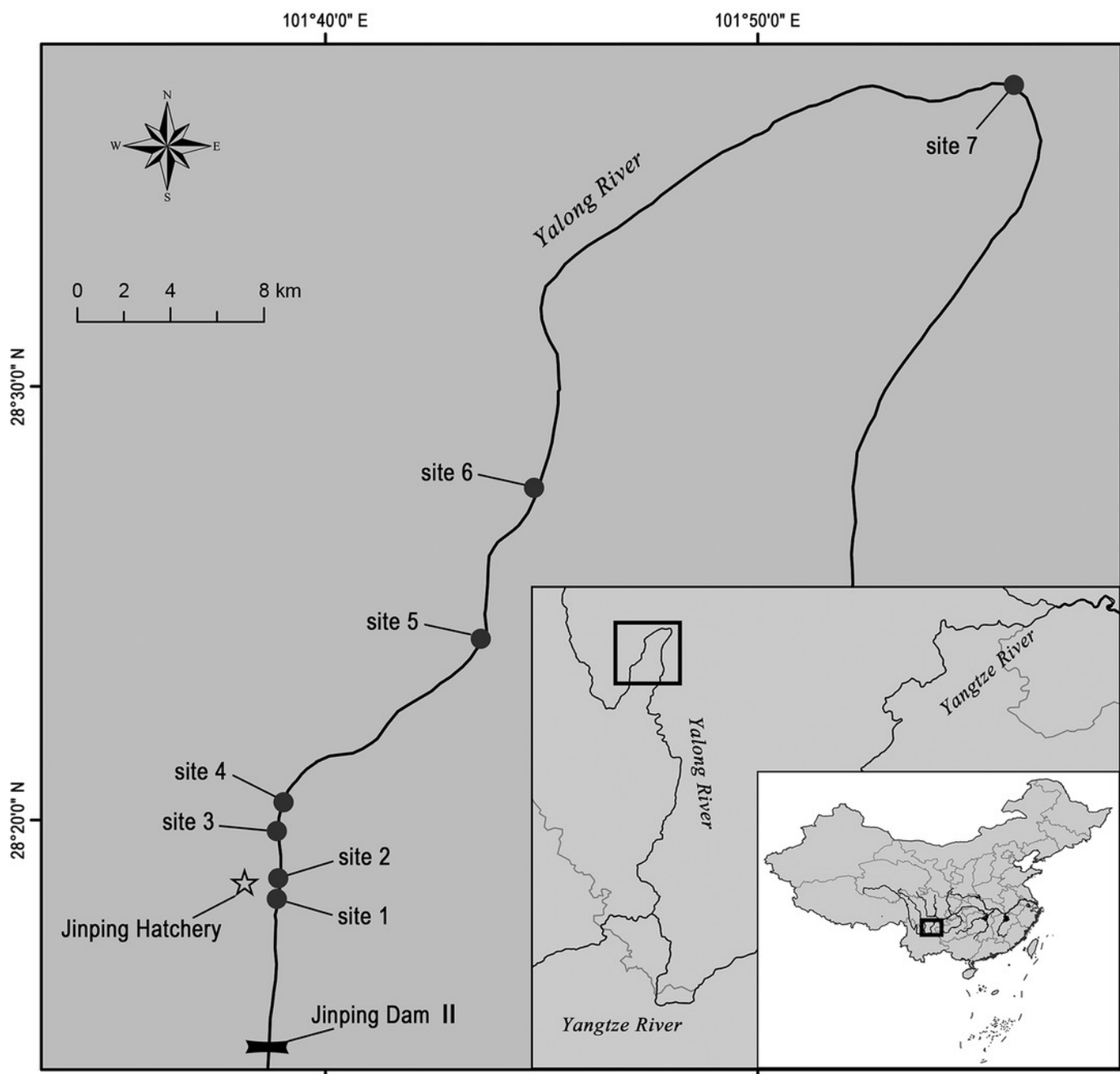
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# Figure 1

Map of the Jinping area of the Yalong River showing the locations of Jinping Hatchery and the sites where the stocked *S. wangchiachii* were released and recaptured.

Fish were released at sites 2 and 3 in July 2015, recaptures were conducted at sites 1–7 from October 2015 to April 2016.

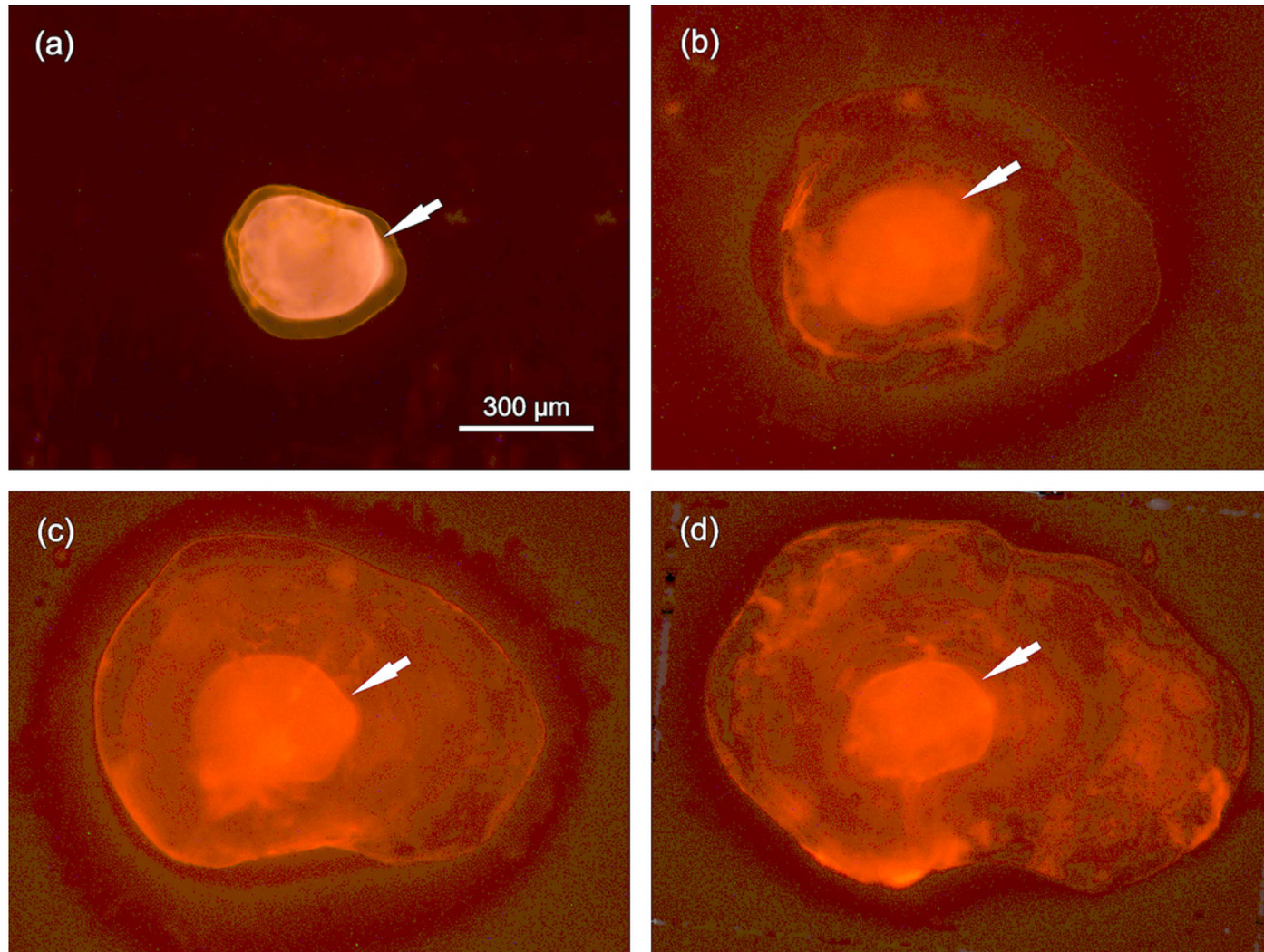


## Figure 2

Photographs of lapillus otoliths of ARS marked *S. wangchiachii* of the 2014 cohort sampled on 29 May 2014 (a), 28 December 2014 (b), 4 May 2015 (c) and 24 January 2016 (d).

These fish were marked by immersing in  $70 \text{ mg L}^{-1}$  ARS for 24 h on 5 May 2014. Photographs were taken under green laser and  $\times 40$  magnification. White arrows show the ARS marks.

*\*Note: Auto Gamma Correction was used for the image. This only affects the reviewing manuscript. See original source image if needed for review.*

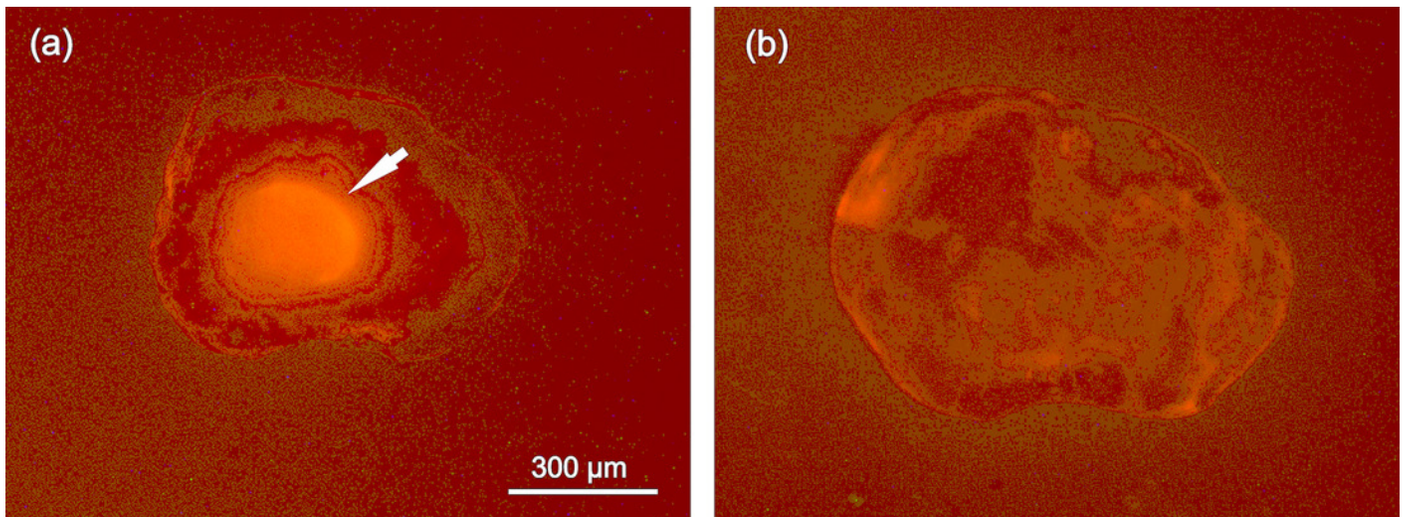


## Figure 3

Photographs of ARS marked (a) and unmarked (b) lapillus otolith of *S. wangchiachii* of the 2015 cohort recaptured in the Jinping area of the Yalong River.

Photographs were taken under green laser and  $\times 40$  magnification. White arrow shows the ARS marks.

*\*Note: Auto Gamma Correction was used for the image. This only affects the reviewing manuscript. See original source image if needed for review.*



## Figure 4

Percent of all marked *S. wangchiachii* captured at each site in three recapture surveys in October 2015 (n = 164), January 2016 (n = 58), and April 2016 (n = 40).

(x, no recapture survey was conducted; \*, the percent was 0%).

