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Title	Diurnal fluctuations in oxygen release from roots of <i>Acorus calamus</i> Linn in a modeled constructed wetland
Authors(s)	Dong, C.; Zhu, W.; Gao, M.; Zhao, L.F.; Huang, J.Y.; Zhao, Y.Q.
Publication date	2011-01
Publication information	Journal of Environmental Science and Health, Part A, 46 (3): 224-229
Publisher	Taylor & Francis
Link to online version	http://dx.doi.org/10.1080/10934529.2011.535391
Item record/more information	http://hdl.handle.net/10197/3114
Publisher's statement	This is an electronic version of an article published in Journal of Environmental Science and Health, Part A, 46 (3): 224-229, available online at: http://dx.doi.org/10.1080/10934529.2011.535391
Publisher's version (DOI)	10.1080/10934529.2011.535391

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1 Diurnal fluctuations in oxygen release from roots of *Acorus*
2 *calamus* Linn in a modeled constructed wetland

3

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5

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12

13 **ABSTRACT**

14

15 The detailed mechanisms of oxygen release from the roots of plants in constructed wetlands (CW)
16 remains unclear. This study investigated the variation of root oxygen release rate, and the effect of
17 photosynthesis during day and night periods on the rate of oxygen release from the roots of *Acorus*
18 *calamus* Linn in a model CW. The maximum oxygen release rate was recorded to be in the range of
19 215.2-750.8 $\mu\text{molg}^{-1}\text{h}^{-1}$ and this occurred at 15:00. The maximum value of photosynthetically active
20 radiation

21

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24 (PAR) ranged from 1281.8-1712.0 $\text{mmolm}^{-2}\text{s}^{-1}$, and this occurred at 13:00. It was observed that both
25 the oxygen release rate and PAR approached zero at night. The results indicate that the rate of
26 oxygen release depends largely on the light intensity, which exhibits a diurnal periodic variation.
27 Accordingly, there are two time intervals namely: day time and night time, during the former period
28 oxygen is released by plants. This study on dynamics of plant root oxygen release distribution has
29 shown that the variation of root oxygen release during daytime followed the Gaussian function. The
30 Gaussian function can be used to predict the root oxygen release rate in constructed wetlands.

31

32 **Keywords** Constructed wetlands; diurnal variation; oxygen release rate; plant; root

33

34 INTRODUCTION

35

36 Constructed wetlands (CW), as low-cost wastewater treatment technology, have been gaining
37 increased international interest and application because of their good treatment efficiency and their
38 natural fit into the landscape. Aerobic decomposition of organic matter and nitrification are affected
39 significantly by the oxygen levels in CW during wastewater treatment. It has been reported that
40 subsurface flow wetland systems have not been successful (in some cases) in removing nitrogen
41 mainly due to lack of dissolved oxygen caused by permanent saturation.^[1,2]

42

43 Plants in CW are known to transport and release oxygen into their root-zones, thus enhancing
44 aerobic processes^[3]. Oxygen released by the root systems of wetland plants is one of the important
45 oxygen sources. Brix^[4] stated that reeds transport oxygen into the rhizosphere, thereby creating

46 aerobic microsites adjacent to the roots and rhizomes. The ability of reeds to transport oxygen and
47 thereby to support a population of aerobic microorganisms in the rhizosphere is one of the key
48 mechanisms for efficient BOD and nitrogen removal. The oxygen release rate into the rhizosphere
49 by a plant can be determined under various light intensities ^[5]. Ojeda et al. ^[6] reported high rates of
50 plant oxygen transfer, suggesting a general view that macrophytes played a considerable role in
51 wastewater treatment. Mitsch ^[7] reported that the rate of oxygen released by plant roots varied from
52 0 to 45gO₂m⁻²d⁻¹. Sorrell and Armstrong ^[8] measured the root oxygen release by bathing whole root
53 systems of *Juncus ingens* in titanium (III) citrate buffer. The results show that the rate of root
54 oxygen release ranged from 0 to 121.6 μmolh⁻¹g⁻¹ root dry wt. It was noted that the root oxygen
55 release rate is highly variable. Therefore, further study of the root oxygen release behaviour is
56 desirable.

57

58 It is well recognized that the rate of oxygen released by the plant root systems is influenced by
59 many factors. Jespersen et al. ^[9] studied the effect of rhizosphere sediment on oxygen release rate
60 by comparing Typhas grown in two sediments (natural organic sediment and a sediment enriched
61 with acetate), and measuring the root oxygen release using titanium (III) citrate buffer. Oxygen
62 demand in the sediment enriched with acetate was higher compared to that of the natural organic
63 sediment. This phenomenon influenced the growth pattern of plants and root shape, and accordingly
64 influenced oxygen release rate, which was about 120-200 μmolO₂h⁻¹g⁻¹ root dry wt. Sasikala et al.
65 ^[10] investigated the effects of water level fluctuation on plant radial oxygen loss (ROL), root
66 porosity, plant growth performance, and nitrogen removal in vertical subsurface flow CW. Their
67 results showed that the quantity of oxygen released by the root systems of plants could be

68 significantly decreased by water level fluctuation. Stottmeister et al. ^[11] described that gas transport
69 from the sections of the plant above the ground through the rhizome into the fine roots was affected
70 by specific areas of tissue formed in the plant known as the aerenchyma. Other factors include
71 rhizosphere-specific parameters such as the redox state, pH, oxygen concentration, chemical
72 characteristics, and plant-specific parameters such as the mass, the plant species and stage of
73 development of plants, as well as variation of climate and different testing conditions.

74

75 In this study, the daily change in the root oxygen release rates were carefully examined using a
76 titanium (III) citrate buffer. The effect of light intensity on root oxygen release rate was studied. A
77 model (using a Gaussian distribution) to describe the root oxygen release behavior was developed
78 based on the experimental data. The model enables the prediction of the diurnal fluctuation of root
79 oxygen release rate.

80

81 **MATERIALS AND METHODS**

82

83 **Experimental Materials and Procedures**

84

85 The plants of young *Acorus calamus* Linn used in this study were collected from a natural wetland
86 located in Xuanwu Lake, Nanjing, China. After collection, the plants were transplanted to
87 individual plastic pots filled with respective nutrient solutions (self-made nutrient solutions with
88 average concentrations of COD and TN of 50mg/L and 15mg/L, respectively) for three weeks
89 before sowing. The plants were removed from the pots and their roots were gently washed free of

90 debris twelve hours before the commencement of experiments. All plants had 20-40 adventitious
91 roots which varied in length from 12-21cm and were up to 0.087 ± 0.029 cm (n=118) in diameter.
92 Their height above ground was 41-58cm.

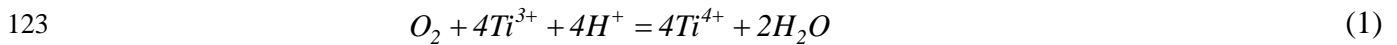
93
94 Oxygen release from the roots was examined using a non-phytotoxic titanium (III) citrate buffer,
95 which allows root oxygen release measurements in a reducing, oxygen-scavenging solution with a
96 low redox potential ^[8,9]. The 1000mL jar was initially filled with 900mL of distilled water, and the
97 water was then sparged with N₂ gas for 60 minutes to remove any oxygen dissolved in the water.
98 Sparging with N₂ gas was continued while titanium (III) citrate stock solution (made by 0.2249g
99 citric acid and 8mL TiCl₃) was added. The basal part of the shoot was wrapped with tinfoil to
100 prevent the oil from infiltrating the aerenchyma. The stirring from the sparging was necessary to
101 ensure complete mixing. The roots of *Acorus calamus Linn* were submerged in the solution. A 5mm
102 thick layer of paraffin oil was placed on top of the solution to prevent re-aeration from the
103 atmosphere. It ensured that the roots were the only possible source of oxygen entry into the chamber.
104 The root chamber was shielded from light using a tight-fitting tinfoil cover. Blank jars without
105 plants were also prepared in a similar way to serve as negative control. Figure 1 shows the
106 experimental device used in the investigation of the root oxygen release rate. The experimental
107 device was exposed to the open air and natural light in a sealed area outside the laboratory building.
108 Light intensity was measured every 1 hour using a luminometer (MODEL ZDS-10F-2D). The unit
109 of light intensity is lux. Details of the experimental set-up are shown in table 1.

110

111 **Sampling and Analytical Methods**

112

113 Since the oxygen released from the roots was oxidized by Ti^{3+} in titanium (III) citrate buffer, rates
114 of root oxygen release could be calculated from the rate of decrease in the concentration of Ti^{3+} in
115 the jars. As the brown titanium (III) citrate solution gradually became clear during oxidation, the
116 samples were taken every 1 hour using a small syringe and the absorbance at 527nm was measured
117 immediately using a spectrophotometer. The absorbances of the samples were compared to those of
118 solutions with a known concentration of Ti^{3+} . At the same time, the light intensity, temperature and
119 humidity were examined. The relation of Ti^{3+} and O_2 is described in the Equation 1. It is seen that
120 1mol O_2 is consumed when 4mol Ti^{3+} were reduced. The oxygen consumption (ΔO_2 , mg) is thus
121 calculated using Equation 2. Thereafter, the root oxygen release rate (V_o , $\mu\text{molg}^{-1}\text{h}^{-1}$) could be
122 calculated using Equation 3.



124
$$\Delta O_2 = \frac{32 \times V \times (C_0 - C_e)}{4 \times 47.73} \quad (2)$$

125 where V is the volume of titanium (III) citrate buffer, 0.9L. C_0 , C_e are the initial and end Ti^{3+}
126 concentration, respectively.

127
$$V_o = \frac{\Delta O_2 \times 1000}{24 \times 32 \times \text{Root dry weighs}} \quad (3)$$

128

129 **RESULTS**

130

131 **Plant Root Oxygen Release Rate**

132

133 Figure 2 illustrates the daily changes of titanium(III) citrate concentration on 22nd April, 2009. The

134 precision of the detecting device is confirmed by comparing the measured Ti^{3+} concentration with
135 that in blank jars. Titanium (III) citrate concentration in the blank jars did not change during the
136 whole experiment. This suggests that the variation of Ti^{3+} concentration was caused by the oxygen
137 released by the plants in the jars.

138

139 According to the experiments conducted on 22nd, 25th and 26th April, 2009, respectively, variation of
140 the root oxygen release rates could be obtained based on Ti^{3+} concentrations in tested jars using the
141 Equations (1)-(3). The dry weights of plant root were measured after drying for 24h at 105 °C. Daily
142 changes of oxygen release rate and PAR are illustrated in Figure 3. The results reveal a significant
143 difference in the root oxygen release rate during day and night. Oxygen release increased gradually
144 with increasing light intensity in the morning. However, a decrease in the oxygen release rate
145 occurred following the decreased light intensity in the afternoon, and approached $0\mu\text{molg}^{-1}\text{h}^{-1}$ at
146 night. These variations indicate a significant time dependency for oxygen release by plants during
147 the day and night. For all three sets of experiments, the start time and end time of oxygen release
148 were closely related to light. The maximum oxygen release rate ($215.2\text{-}750.8\mu\text{molg}^{-1}\text{h}^{-1}$) was
149 observed during the daytime at 15:00 hrs while the maximum light intensity was observed at 13:00
150 hrs. The maximum value of PAR ranged from 1281.8 to 1712.0 $\text{mmolm}^{-2}\text{s}^{-1}$. Clearly, the peaks of
151 root oxygen release occurred after the peak of light intensity.

152

153 **Daily Variations of Root Oxygen Release and Light Intensity—Application of a Gaussian**
154 **Function**

155

156 The variation of the root oxygen release during the day-night is schematically summarized in Figure
 157 4. There are two time intervals during the day-night cycle, i.e. periods of brightness and darkness.
 158 The t_{Ls} and t_{Le} , are the start and end time of the bright period, respectively. They also correspond to
 159 the sunrise (t_{Ls}) and sunset (t_{Le}) time. L_{max} is the maximum light intensity at the corresponding time
 160 t_{Lmax} . The t_{Os} and t_{Oe} , are the start and end time of oxygen release period, respectively. V_{Omax} is the
 161 maximum oxygen release rate at the time of t_{Omax} . Since the time of maximum oxygen release rate
 162 occurred at a later period than the time of maximum light intensity, this time difference is termed as
 163 lag time (Δt). It may be caused by photosynthesis and oxygen transport in the aerenchyma.

164
 165 In order to describe the daily variation of the root oxygen release in a mathematical way, data
 166 obtained from the experiments were preliminarily fitted using several functions (t_{Os} was 4:00 and
 167 t_{Oe} was 20:00). The results reveal that the best fitting could be achieved by Gaussian function ^[13],
 168 which represents unimodal distribution model as shown in Figure 5(a). The goodness of fit (R^2) is
 169 0.7574, 0.5357, 0.6796 with 95% confidence bounds on 22nd, 25th and 26th April, respectively.
 170 Based on the form of Gaussian function, diurnal variation of root oxygen release rate could be
 171 described as:

$$V_O = ae^{-\frac{(t-t_{Omax})^2}{c^2}} \quad (4)$$

172
 173
 174 where t is time (4:00~20:00); a (in the Gaussian function) is the maximum value of oxygen release
 175 rate in a whole day; t_{Omax} is the location of the symmetry axis in Gaussian function; c expresses the
 176 gradient of the Gaussian function. A decrease in c indicates a steep Gaussian function while an
 177 increase in c indicates a gentle Gaussian function. Figure 5(a) shows the root oxygen release data

178 (on 26th April) with Gaussian function fitting, where a , c and t_{Omax} are $613.1\mu\text{molg}^{-1}\text{h}^{-1}$, 3.884 and
 179 15:00, respectively.

180

181 Light intensity data during daytime (4:00-20:00) also follow a Gaussian function (Fig. 5(b)). It can
 182 be described as:

$$183 \quad PAR = be^{-\frac{(t-t_{Lmax})^2}{d^2}} \quad (5)$$

184 where PAR is the photosynthetically active radiation, $\mu\text{mol}\cdot\text{m}^{-2}\text{ s}^{-1}$; t is time; b is the peak value of
 185 PAR in a whole day; and d is the gradient of the unimodal function. Figure 5(b) shows the light
 186 intensity data (on 26th April) with a Gaussian function fitting, where b , d and t_{Lmax} are
 187 $1702\mu\text{mol}\cdot\text{m}^{-2}\text{ s}^{-1}$, 3.672 and 13:00, respectively.

188

189 The peak value of root oxygen release was observed after the maximum light intensity for 2 hrs.
 190 The correlation between the light intensity and the oxygen release data (collected 2 hrs later) was
 191 analyzed and illustrated in Figure 6. It is clear from Figure 6 that the root oxygen release rate was
 192 influenced by light intensity dramatically. Oxygen release rate increased exponentially with
 193 increased PAR ($R^2 = 0.8689$):

$$194 \quad V_o = 62.22e^{0.00138PAR} \quad (6)$$

195 By combining Equations. (4), (5) and (6), the following equation is obtained:

$$196 \quad ae^{-\frac{(t-t_{Omax})^2}{c^2}} = 62.22e^{0.00138\left(be^{-\frac{(t-t_{Lmax})^2}{d^2}} \right)} \quad (7)$$

197 It should be pointed out that, in Equation 7, $t_{Lmax}=t_{Omax}$ since the oxygen release curve (see Figure 4)
 198 was shifted for 2 hrs when the light intensity and the oxygen release data was correlated, as

199 described in Equation 6.

200

201 In the special case, when the time (t) is equal to the peak time (t_{Omax} or t_{Lmax}), Equation 7 becomes:

$$202 \quad a=62.22e^{0.00138b} \quad (8)$$

203 Equation 8 indicates that the parameters of a in oxygen release behaviour and b in light intensity
204 follow an exponential function.

205

206 The parameters c and d also follow an exponential function. The daily changes of root oxygen
207 release and light were examined. Values of c and d derived from the experiments were fitted using
208 an exponential function (R^2 is 0.9587). The relationship is as follows:

$$209 \quad c=0.66e^{0.4856d} \quad (9)$$

210 Therefore, the Gaussian function can be used to predict the oxygen release rate. The procedure is as
211 follows: (1) obtain light intensity data; (2) fit the data using Gaussian function from which the
212 parameters of b and d could be obtained using Equation 5; (3) determine parameters a and c using
213 Equation 8 and Equation 9, respectively; (4) calculate oxygen release using Equation 4.

214

215 **Validation**

216

217 The experiments were conducted with the same testing device in October 2009 to validate the
218 model of the proposed Gaussian function. PAR was tested every 1 hour during the daytime
219 (4:00~20:00), and the parameters obtained are tabulated in table 2.

220

221 Measured oxygen release rates and the corresponding predicted values using a Gaussian function
222 are jointly illustrated in Figure 7. From the results of the stimulation, it can be seen that the
223 Gaussian function can be satisfactorily used to predict the daily changes of oxygen release by roots
224 of wetland plants. The model data closely correlate with the experimental values.

225

226 **DISCUSSION**

227

228 It is well recognized that the oxygen release rate of wetland plants is associated with light intensity.
229 In addition, the oxygen release rate varies with time during natural changes in light intensity.
230 However, no detailed information on the variation of oxygen release rate is found in the literature.
231 In this study, the oxygen release rate of a wetland plant, *Acorus calamus Linn*, was examined in detail.
232 It has been shown that the oxygen release rate for *Acorus calamus Linn* (Fig. 3) appears to be much
233 higher than those reported in the literature although different wetland plants were tested ^[14,15]. Mei
234 et al., ^[15] reported oxygen release rate of 7.40-13.24 $\mu\text{molO}_2 \text{ h}^{-1}\text{g}^{-1}\text{root dw}$ for *Shengtail* and
235 *Suyunuo*, while oxygen release rate of 1.6 $\mu\text{molO}_2\text{h}^{-1}\text{g}^{-1}\text{dw}$ for *Cladium* was reported by Chabbi et al.
236 ^[14]. It is noted in the literature that even for the same wetland plant, the reported oxygen release rate
237 is significantly different. For example, Sorrel and Armstrong ^[8] reported oxygen release rate of 126
238 $\mu\text{molO}_2\text{h}^{-1}\text{g}^{-1}\text{dw}$ for *Juncus ingens*, while a value of 1.5 $\mu\text{molO}_2\text{h}^{-1}\text{g}^{-1}\text{dw}$ for *Juncus bulbosus* was
239 reported by Chabbi ^[16]. The reason for the variation in reported values may be partially attributed to
240 the light intensity.

241

242 Oxygen is produced during photosynthesis ^[17] and is transferred from the leaves to the roots

243 through the gas-filled tissues of plant by the process of diffusion and convection ^[18]. Oxygen is then
244 released to the rhizosphere by gas exchange. The photosynthetic rate of plants was highly correlated
245 with light intensity. The photosynthetic characteristics can affect their ability to provide oxygen ^[19].
246 The light-dark switch generates a large and rapid fluctuation in the internal oxygen levels of plants
247 ^[18]. Thus, plants also experience great released oxygen fluctuations. In this paper, the rate of oxygen
248 release was shown to depend largely on the light intensity, which exhibits a diurnal periodic
249 variation. The variation of oxygen release and light intensity followed unimodal distribution and,
250 furthermore, followed the Gaussian function during the daytime. In particular, the maximum root
251 oxygen release rate was shown to occur 2 hours after the occurrence of maximum light intensity
252 (Fig. 3), from which the relationship between the root oxygen release rate and the light intensity
253 was established. Although a recent study has shown that the maximum root oxygen release with up
254 to 35% oxygen saturation at the root surface occurred under light conditions while a decrease of
255 about 30% was observed under dark conditions ^[5], the present study has given a detailed profile
256 showing the daily changes of root oxygen release rate with natural light.

257

258 More significantly, this study presented a methodology of root oxygen release prediction using a
259 Gaussian function. This allows us to use the light intensity data to calculate the quantity of oxygen
260 likely to be released. However, further studies are still needed to demonstrate the application of the
261 Gaussian function when other wetland plants are tested. It should also be noted that the method of
262 the Gaussian function was established based on the experimental data collected at Nanjing with a
263 unique climate. Thus, validation studies in other places with different natural light should be
264 considered before the methodology established in current study is applied more generally.

265 **CONCLUSIONS**

266

267 The oxygen release rate of wetland plants exhibited diurnal periodic variation. Light intensity is a
268 major factor influencing oxygen release. In the morning, the oxygen release rate increased with
269 increasing light intensity. Both the values of the oxygen release rate and light intensity decreased
270 gradually in the afternoon, and approached $0\mu\text{molg}^{-1}\text{h}^{-1}$ at night due to the absence of the light.
271 More significantly, the variation of the root oxygen release rate and light intensity followed a
272 unimodal distribution. The Gaussian function has been demonstrated to well describe the day time
273 variation of the root oxygen release rate. It can also be used for prediction purposes of the root
274 oxygen release rate in constructed wetlands.

275

276 **ACKNOWLEDGEMENTS**

277

278 This study was financially supported by the National Natural Science Foundation of China
279 (50979028) and the Ministry of Water Resources of China (200801065).

280

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- 331 **Figure 1.** Schematic sketch of the root oxygen release rate detecting device
- 332 **Figure 2** Daily changes of titanium (III) citrate concentration
- 333 **Figure 3** Diurnal fluctuation of root oxygen release rate: (a) 22nd April; (b) 25th April; (c) 26th April
- 334 **Figure 4** Schematic indication of the diurnal fluctuation of the root oxygen release
- 335 **Figure 5** Fitting root oxygen release rate (a) and light intensity (b) using Gaussian function
- 336 **Figure 6** The effect of light intensity on root oxygen release rate
- 337 **Figure 7** Comparison of observed and predicted oxygen release rate in October 2009
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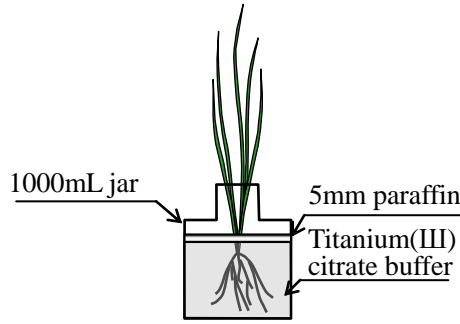


Fig.1

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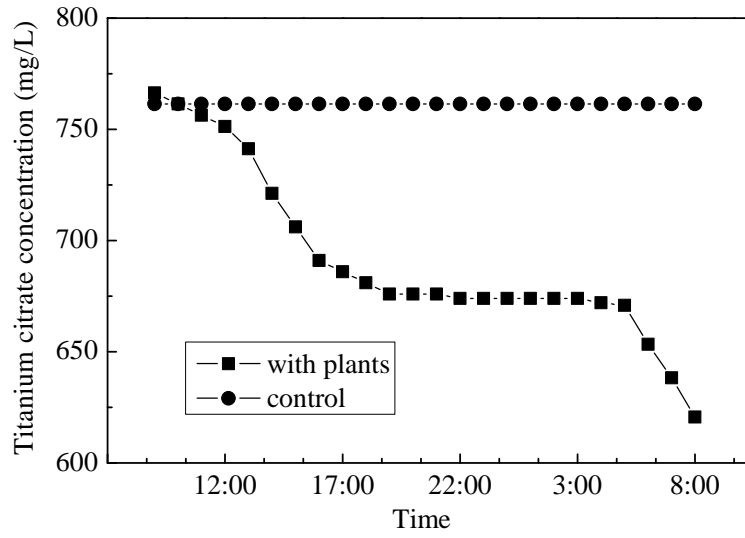
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Fig. 2

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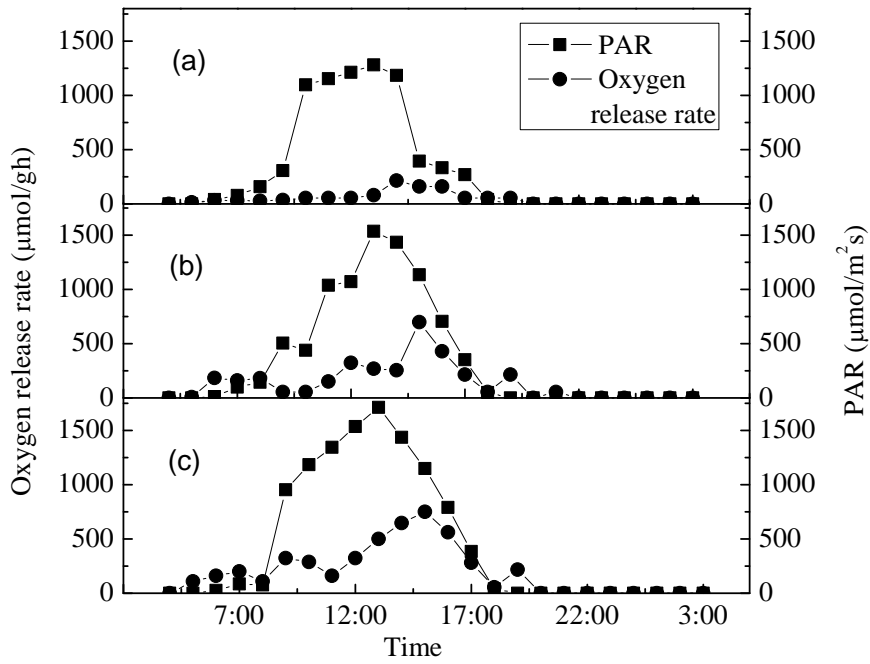
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Fig. 3

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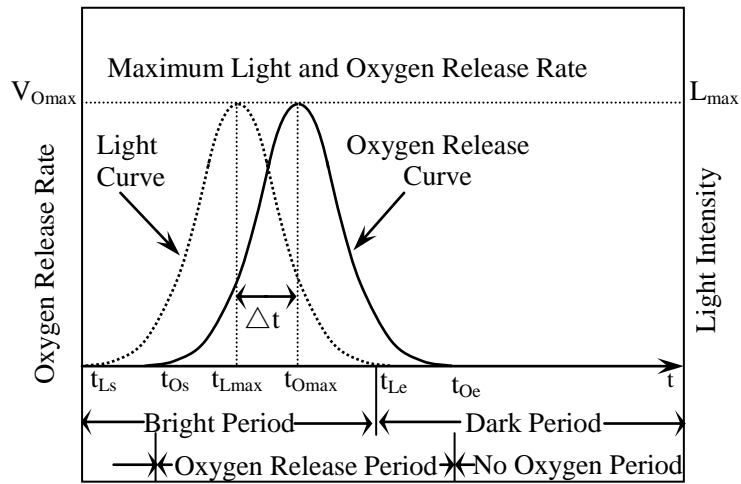


Fig. 4

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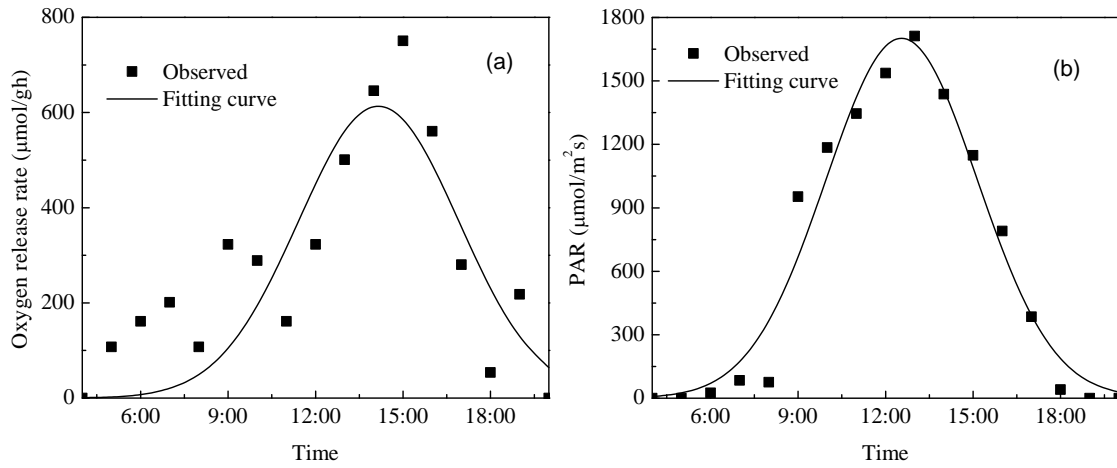


Fig. 5

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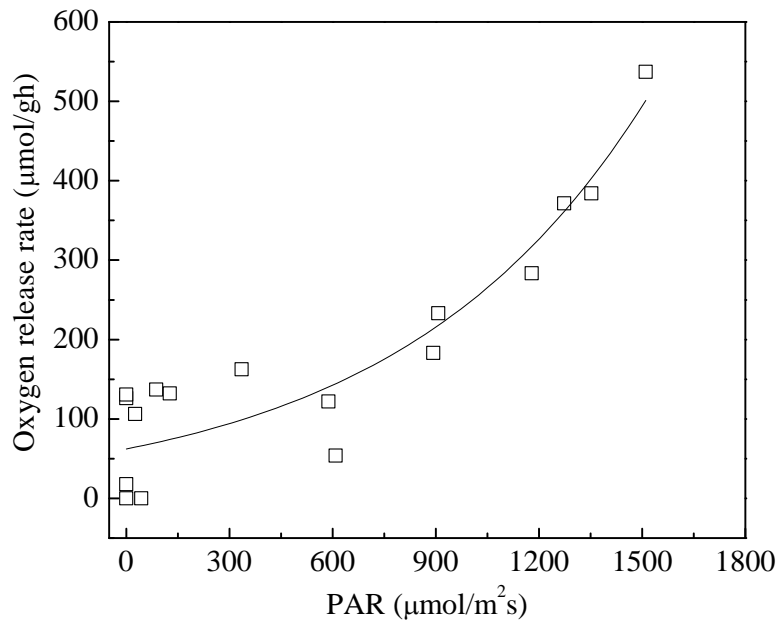


Fig. 6

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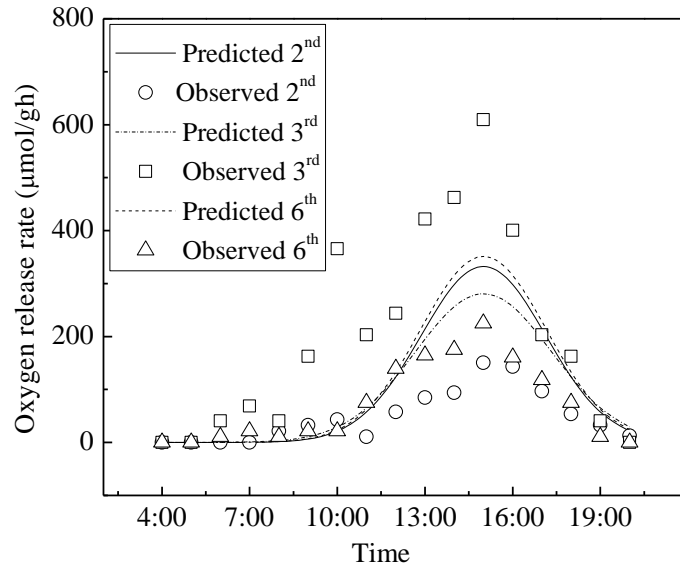


Fig. 7

Table 1. Light intensity, temperature and humidity during experiments

Experimental date	PAR ($\mu\text{mol}\cdot\text{m}^{-2}\text{ s}^{-1}$)		Temperature ($^{\circ}\text{C}$)		Humidity (%)	
	Average	range	Average	range	Average	range
22 April 2009	444.2	0-1281.8	24	19-28	38	15-61
25 April 2009	501.1	0-1536.4	22	18-25	27	10-52
26 April 2009	630.5	0-1712.0	22	18-25	27	9-55

* Sample number is 24. PAR is photosynthetically active radiation. One lux is $0.019 \mu\text{mol}\cdot\text{m}^{-2}\text{ s}^{-1}$ [12].

Table 2. The parameters of modeling

Date	PAR		Oxygen release	
	b ($\mu\text{mol}\cdot\text{m}^{-2}\text{ s}^{-1}$)	d	a ($\mu\text{molg}^{-1}\text{h}^{-1}$)	c
2 nd October	1214	3.137	332.31	3.020
3 rd October	1092	3.33	280.82	3.317
6 th October	1254	3.155	351.16	3.047