## roGFP Derivations & Sensitivity Analysis

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## 1 Converting 410/470 intensity ratio to the proportion of oxidized sensor molecules

The  $roGFP_{12}$  sensor is genetically encoded into a tissue. The intensity values emitted from the  $roGFP_{12}$ -containing tissue are recorded after being stimulated 410nm and 470nm light. The ratios between these values describe relative levels of tissue oxidation. For example, a 410/470 ratio of 2.0 indicates that a tissue is more oxidized than a ratio of 1.0. In this brief summary, we algebraically convert the 410/470 intensity ratio into the more biologically-meaningful measure of redox potential (E) via the description of ratio of of  $roGFP_{12}$  molecules in an oxidized (as opposed to reduced) state (OxD).

Assume a fully reduced state. Then, the intensities observed at a wavelength  $\lambda$  are equal to the product of  $N_T$ , the total number of roGFP molecules, and  $I_{\lambda,R}$ , the intensity of each roGFP molecule at a given wavelength in the reduced state.

$$I_{\lambda,R} = N_T * I_{\lambda,R} \tag{1}$$

The same is true for the fully oxidized state:

$$I_{\lambda,Ox} = N_T * I_{\lambda,Ox} \tag{2}$$

At a redox state between maximally reduced and maximally oxidized, the intensity at a given wavelength is a weighted sum of the molecules found at either discretely oxidized or reduced state. We therefore can rewrite any state in terms of equations (1) and (2)

$$I_{\lambda} = \frac{N_{Ox}}{N_{T}} * I_{\lambda,Ox} + \frac{N_{Red}}{N_{T}} * I_{\lambda,Red}$$
(3)

Because all sensor molecules must be in either an oxidized or reduced state,  $N_{Red} = N_T - N_{Ox}$ . So we can rewrite equation (3):

$$I_{\lambda} = \frac{N_{Ox}}{N_T} * I_{\lambda,Ox} + \left(1 - \frac{N_{Ox}}{N_T}\right) * I_{\lambda,Red} \tag{4}$$

Using equation (4), consider the intensity ratio of 410 nm / 470 nm:

$$\frac{I_{410}}{I_{470}} = \frac{\frac{N_{Ox}}{N_T} * I_{410,Ox} + (1 - \frac{N_{Ox}}{N_T}) * I_{410,Red}}{\frac{N_{Ox}}{N_T} * I_{470,Ox} + (1 - \frac{N_{Ox}}{N_T}) * I_{470,Red}} =$$

For brevity, let  $OxD = \frac{N_{Ox}}{N_T}$ . Then cross-multiply:

$$I_{410} * OxD * (I_{470,Ox} + (1 - OxD) * I_{470,Red}) =$$

$$I_{470} * OxD * (I_{410,Ox} + (1 - OxD) * I_{410,Red})$$

Simplify and express OxD in terms of known quantities:

$$OxD = \frac{I_{470}I_{410,R} - I_{410}I_{470,R}}{I_{410}I_{470,Ox} - I_{410}I_{470,R} - I_{470}I_{410,Ox} + I_{470}I_{410,R}}$$
(5)

To simplify, let:

$$R_{Red} = \frac{I_{410,R}}{I_{470,R}} \tag{6}$$

$$R_{Ox} = \frac{I_{410,Ox}}{I_{470,Ox}} \tag{7}$$

$$\frac{I_{410}}{I_{470}} = \frac{I_{410}}{I_{470}} \tag{8}$$

$$\delta_{470} = \frac{I_{470_{Ox}}}{I_{470_{Red}}} \tag{9}$$

We can now re-derive the definition of OxD in terms of ratio values.

Step: Re-arrange terms, multiply by  $\frac{-1}{-1}$ :

$$OxD = \frac{I_{410}I_{470,R} - I_{470}I_{410,R}}{I_{410}I_{470,R} - I_{470}I_{410,R} + I_{470}I_{410,Ox} - I_{470,Ox}I_{410}}$$

Step: Work to factor out  $I_{470,R}i_{470}$  from the numerator and denominator write some in terms of ratio values:

$$OxD = \frac{I_{470,R}I_{470}(\frac{I_{410}}{I_{470}} - R_{Red})}{I_{470,R}I_{470}(\frac{I_{410}}{I_{470}} - R_{Red} + \delta_{470}(R_{Ox} - \frac{I_{410}}{I_{470}}))}$$

And simplify:

$$OxD = \frac{\frac{I_{410}}{I_{470}} - R_{Red}}{\frac{I_{410}}{I_{470}} - R_{Red} + \delta_{470}(R_{Ox} - \frac{I_{410}}{I_{470}})}$$
(10)

Where  $\delta_{470}$  describes the sensor-specific dynamic range 470nm.

## 2 Analyzing the conversion between intensity ratio and ratio of oxidized molecules

There are 4 parameters that fit into the determination of the fraction of oxidized molecules at any given point:

- 1.  $R_{410/470}$  is the ratio of intensity at 410/470, which is usually the observed value of interest.
- 2.  $R_{Red}$  is the minimal intensity value at 410/470 (when the sensor is maximally reduced).
- 3.  $R_{Ox}$  is the maximal intensity value at 410/470 (when the sensor is maximally oxidized).
- 4.  $\delta_{470}$  is the dynamic range of the sensor at 470 (ratio of 470 at maximally reduced and maximally oxidized, respectively).

Experimentally, our function's true input is  $R_{410/470}$ , so we can plot the fraction oxidized as a function of  $R_{410/470}$ .

Note that varying  $\delta_{470}$  varies the linearity of the relationship between intensity ratio and fraction oxidized (Figure 1), whereas varying  $R_{Red}$  or  $R_{Ox}$  simply translates the function by a constant.

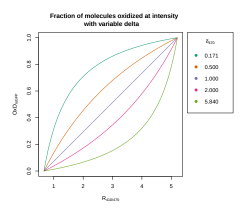


Figure 1: The relationship between observed intensity and OxD with different  $\delta_{470}$  constants

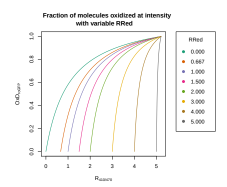


Figure 2: The relationship between observed intensity and OxD with different  $R_{Red}$  constants

- 3 Converting fraction of oxidized sensors into an electric potential
- 4 Reparameterizing the intensity ratio