A comparison between the responses of neutral red and acridine orange: Acridine orange should be preferential and alternative to neutral red as a dye for the monitoring of contaminants by means of biological sensors

Sabrina Manentea, Silvia De Pieria, Alessandra Ierob, Chiarafrancesca Rigoa, Marcantonio Bragadinb

a Department of Environmental Sciences, Ca’ Foscari University of Venice, Dorsoduro, Venice 30123, Italy
b Institute of Applied Ecology, Health Science, and Design, University of Canberra, Bruce, ACT 2617, Australia

ABSTRACT

The acridine orange (AO) and neutral red (NR) dyes, commonly used as probes to measure the internal pH in acidic vesicles, are compared in this article. The comparison between the two dyes (arising from calculations taking into account their analytical constants) illustrated that the use of AO is preferential to that of NR because the AO response is sensitive over the whole pH range between 4.0 and 7.4, whereas the NR response is effective only between pHs 4.0 and 6.0. In addition, it became evident from the mitochondrial respiration response that NR, unlike AO, is a protonophore. When taken into consideration, these two properties suggest that AO is more suitable than NR as an indicator of toxicity measurements in water samples because the environmental toxic compounds induce pH changes in the acidic vesicles of biological structures that are used as environmental biosensors.

Many methods have been proposed for the evaluation of the presence of toxic compounds in water and to assess the risks to both humans and animals. These methods, based on the responses to toxic compounds in fish and invertebrates, are often impractical because they are expensive and time-consuming. These factors have stimulated researchers to develop more rapid and inexpensive methods of evaluation, and a wide range of alternative in vitro tests have been proposed [1–6]. One of these in vitro tests uses the response of mussels to monitor the environmental status of sea water [3,7–9].

When marine molluscs such as mussels are exposed to contaminants, one of the characteristic pathological alterations is a decreased integrity in the lysosomal membrane [10]. This loss of integrity is assayed using the dye neutral red (NR) [1]. The dye enters the lysosomes of the mussels and accumulates inside, with the driving force being the internal acidic pH. The damage to the lysosomes, induced by the toxic compounds, induces a release of the dye from the lysosomes to the resuspending medium. A spectrophotometric measurement of the amount of NR in the resuspending medium gives a measurement of the level of damage induced by the toxic compounds. We believe, and demonstrate in this article, that the procedure has some weak points; the acridine orange (AO) dye, which is widely used for acidic pH measurements in isolated lysosomes, offers some advantages and is preferred to measurements based on the NR response.

Materials and methods

Acid pH measurements using the response of the AO dye

The measurements were performed using a Jenway 6400 spectrophotometer under stirring at room temperature. The lysosomes, having been prepared and purified according to the classical procedure [11], were resuspended in a medium (2.5 ml) (final concentration 0.2 mg/ml). The medium composition was 0.125 M KCl, 10 mM Hepes–Mops (pH 7.4), and 5 mM MgCl2. Following the addition of the AO dye (5 μM), the instrument was adjusted to zero absorbance. The addition of 2 mM adenosine triphosphate (ATP), therefore, induces an absorbance quenching at 492 nm.

Oxygen consumption experiments

The mitochondria from rat liver were prepared following the usual procedure [12]. The mitochondrial protein content was determined using the method of Lowry and coworkers [13]. The mitochondrial oxygen consumption was measured using a Clark oxygen electrode (Yellow Springs Instruments, Yellow Springs, OH, USA) fitted in a closed thermostated chamber (2 ml) at 20 °C equipped with magnetic stirring. The mitochondria (final...
concentration 2.5 mg/ml) were resuspended in 2 ml of the medium. The medium composition was 0.25 M sucrose, 10 mM Hepes–Mops (pH 7.4), and 1 mM MgCl₂.

All of the reagents were of analytical grade. The succinate, AO, NR, and p-trifluoromethoxy carbonyl cyanide phenylhydrazone (FCCP) were purchased from Sigma–Aldrich.

Results and discussion

The measurement of the internal pH in acidic vesicles is based on the responses given by dyes such as AO and NR [14–17]. Both compounds are weak bases (pKₐ = 6.5 for NR, pKₐ = 10.5 for AO [15,18]). These weak bases, being permeant, cross the biological phospholipidic membranes as neutral compounds and, following the classical transport mechanism of weak permeant bases [19], accumulate in the matrix, the driving force being the internal acidic pH (pHi 4.0 in Fig. 1). The accumulation in the matrix gives rise to an absorption change that is probably due to a metachromatic effect [16]. Therefore, in the case of AO, by operating at a wavelength of 492 nm, which is the absorption maximum in water, the accumulation in an acidic matrix gives rise to a spectral change and the acidification is followed by the signal quenching that accompanies the entry of the dye. Fig. 2 shows an example of the use of AO to follow the acidification process in isolated lysosomes, where the pH in the matrix is approximately 4.5 [11,17,19]. Analogous responses are obtained if NR, instead of AO, is used [16]. The AO and NR dyes accumulate in the acidic compartments of the cell by means of a similar mechanism [19,11,20].

This capacity of measuring the internal pH in lysosomes is used in environmental tests to monitor the quality of water samples [7,8] because it is assumed that the status of the lysosomes (i.e., the pH in the lysosomal matrix) in the cells of the biosensors (mussels) is a measurement of the environmental contamination [7–9]. To this end, the lysosomal efficiency in mussels is measured on the basis of the accumulation (and release) of NR. The standard test [21] forecasts an incubation period of 3 h following the addition of the dye, and that the same type of measurements could give some advantages if they were carried out using AO instead of NR.

The accumulation ratio R (i.e., the ratio between the internal and external concentration of the dye),

$$R_{AO} = \frac{[AO]_i + [AOH^+]_i}{[AO]_o + [AOH^+]_o}$$

for AO and

$$R_{NR} = \frac{[NR]_i + [NRH^+]_i}{[NR]_o + [NRH^+]_o}$$

for NR.

Where

$$[AO]_i$$ and $$[NR]_i$$ are the concentrations of the AO and NR dyes in the matrix of the vesicle in undissociated form;

$$[AO]_o$$ and $$[NR]_o$$ are the concentrations of the dyes in the external medium in undissociated form;

$$[AOH^+]_i$$ and $$[NRH^+]_i$$ are the concentrations of AO and NR in protonated form in the matrix of the vesicles;

$$[AOH^+]_o$$ and $$[NRH^+]_o$$ are the concentrations of AO and NR as protonated forms in the external medium (see the scheme in Fig. 1),

Fig. 1. The classical mechanism responsible for the entry and accumulation of weak permeant bases in acidic vesicles. The dye (in this case AO, although this type of behavior is common to all weak permeant bases such as NR) is soluble in the membrane (phospholipidic bilayer) in its undissociated form (AO). For symmetry reasons, the same situation occurs in the internal aqueous phase, where the pH is 4.0. If the internal matrix is acidic, this induces an accumulation of the weak base in the matrix, and the accumulation increases as the internal pH (pHi) becomes more acidic.

Fig. 2. An example of the use of AO as an indicator of the internal pH in lysosomes. The acidic pH in lysosomes induces an accumulation of AO, with a consequent signal quenching at 492 nm (metachromasy). The release of the dye induced by a classical protonophore such as FCCP (100 nM) shows an example of release of the dye induced by membrane permeability enhancement and indicates that the transport of AO is very fast.
in acidic vesicles can easily be calculated when the pKₐ values for AO and RN are taken into account. Another factor is that, as a consequence of the existence of a partition coefficient, 
\[
K = \frac{[\text{AO}]_{\text{membrane}}}{[\text{AO}]_{\text{water}}},
\]
where 
- \([\text{AO}]_{\text{membrane}}\) is the concentration of the soluble AO in the membrane;
- \([\text{AO}]_{\text{water}}\) is the concentration of the AO soluble in water,
and then \([\text{AO}]_{\text{o}} = [\text{AO}]_{\text{i}}\) (see Fig. 1). Analogous calculations regarding NR give the same conclusion (i.e., \([\text{NR}]_{\text{o}} = [\text{NR}]_{\text{i}}\)). The results of the calculations are summarized in Table 1.

Because the internal pH in the lysosomes (pHi) of the cells is modified by the presence of toxic compounds, with a consequent release of either AO or NR from the lysosomes to the resuspending medium, Table 1 reports the accumulation ratio and the concentration of the AO and NR dyes in the supernatant, calculated at different pH values. With regard to these calculations, the pHi in the cytoplasm is 7.5.

It can be seen in Table 1 and from the corresponding graph (Fig. 3) that NR is sensitive to the pH only in the pH values ranging from 4.0 (4.5–5.0 is the pH value in the matrix of lysosomes in noncontaminated cells) to 6.0. In the pH values from 6.0 to 7.5 (which is the physiological pH value in the cytoplasm), the response and the release of NR in the supernatant is the same. This means that any damage to the lysosomes that induces an enhancement of the pHi from 6.0 to 7.5 gives the same response, corresponding to that given by the total destruction of the lysosomal membrane (i.e., a pHi of 7.5). Therefore, NR cannot be used as a probe in measurements of the pHi values that range between 6.0 and 7.5. This problem does not occur when using AO (see the graph in Fig. 3) and is due to the different pKₐ values of the two dyes.

There are additional advantages in choosing AO instead of NR. Both dyes are protonophores (i.e., substances that enhance the proton permeability) [1,9,14,22], but AO is less protonophoric than NR. The mechanism that induces an enhancement of the proton conductivity is due to the presence of a potential in the biological membranes. With regard to lysosomes, the matrix pH is approximately 4.5 to 5.0 [19] due to the presence of an ATP-driven proton pump [11,19]. This pump drives the protons in the matrix. The latter gives rise to a positive inside potential that is only partially neutralized by the entry of Cl⁻ ions through a selective channel [19]. In steady-state conditions, the potential is approximately +40 mV [23]. This value is not particularly high, but it could induce an extrusion of the accumulated dye as a charged cation, thereby giving rise to a cyclic mechanism. The mechanism is called a protonophore effect because, at any cycle, a proton is extruded with a consequent pH enhancement. Obviously, such an effect should be as minimal as possible because it is equivalent to that produced by environmental contaminants that damage the membrane, as already mentioned.

Therefore, the crucial question is as follows: what is the extent of this effect with regard to AO and NR? Because the main inducer of the protonophore effect is the potential (the transport of the undissociated compound is rapid, as Fig. 2 shows in the case of AO), a measurement of the protonophore effect (or the uncoupling effect [22]) in mitochondria is useful at this point, where the internal potential is high at approximately ~200 mV.

There are many procedures used to assess the presence of a protonophore in mitochondria. Once it has been verified that the protonophore does not inhibit the stimulated respiratory rate (not shown), the simplest procedure is to measure the stimulation induced by a protonophore in the basal mitochondrial respiration (state 4 mitochondrial respiration [14,22]). Fig. 4A shows the mitochondrial state 4 stimulation induced by FCCP, which is a classical protonophore. Fig. 4B shows that a similar effect is induced by means of 20 μM NR. Fig. 4C shows that no stimulation is obtained with 80 μM AO. The behavior of the AO dye in this case confirms the data already reported [14]. This leads us to the conclusion that NR, as in literature [24], is a protonophore, whereas AO is not in analogous conditions. Consequently, AO is “less contaminating” or invasive than NR; that is, the presence of AO in the biological membrane is less toxic and destructive than that of NR.

### Conclusions

The above-discussed results lead to the following conclusions. First, the analytical calculations indicate that the response of AO covers the whole pH range between 4.0 and 7.5, whereas NR is insensitive in the pH range between 6.0 and 7.5. Second, the experimental measurements indicate that, in similar conditions, AO is not a protonophore, whereas NR has a protonophore effect that is similar to that induced by toxic compounds. These two crucial points, together with the facts that the absorption coefficient (ε) is 18,000 for AO and 18,500 for NR (allowing the use of lower concentrations of AO to obtain the same responses as with NR) and that the accumulation of AO is instantaneous (Fig. 2) and no incubation times are necessary (whereas the standard test implies an incubation period of 3 h in the case of NR [14]), lead us to the conclusion that AO is preferable to NR for toxicity measurements in...
aqueous samples using biological structures such as the mussels used in the standard test procedure.

References


Fig. 4. A typical experiment regarding the mitochondrial respiratory rate stimulation induced by a classical uncoupler (or protonophore), FCCP (100 nM) (A), or by increasing amounts (up to 20 μM) of NR (B). Each arrow in panel B corresponds to the addition of 5 μM NR. The addition of AO up to 80 μM does not induce any respiratory stimulation (C). In panel C, each arrow indicates the addition of 20 μM AO. In panel A, the mean value of the respiratory rate (μL oxygen/mg protein/min) after the addition of succinate (2 mM), as results from three experiments, is 16 ± 2. After the addition of FCCP, the mean value of the rate is 209 ± 10. In panel B, after the addition of succinate, the respiratory rate is 16 ± 2. After the addition of NR (5 μM), the rate is 33 ± 2. After two successive additions of 5 μM NR, the rates are 69 ± 3 and 108 ± 6, respectively. In panel C, the basal respiration is 16 ± 2 and the addition of AO does not induce respiratory rate modifications.