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Extending Human Perception of Electromagnetic Radiation to the UV Region through Biologically Inspired Photochromic Fuzzy Logic (BIPFUL) Systems.

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Photochromic Fuzzy Logic Systems have been designed that extend human visual perception into the UV region. The systems are founded on a detailed knowledge of the activation wavelengths and quantum yields of a series of thermally reversible photochromic compounds. By appropriate matching of the photochromic behaviour, unique colour signatures are generated in response differing UV activation frequencies.

A deep scientific understanding of how the human brain perceives, thinks, and acts will have a revolutionary impact in science, medicine, economic growth, security and well-being as recently expounded by an interdisciplinary and international team lead by Albus. Perception plays a relevant role within the cognitive architecture of the human nervous system. In fact, it builds and maintains an internal model of the external world and conditions behavior. Among the different sensory systems, we have at our disposal, the visual system is remarkable because it allows us to discern colour, shape and the movement of objects. Human colour perception is founded upon a mosaic of many replicas of three types of photosensitive cells, termed cones. Each type of cone absorbs a particular region of the visible spectrum, although their spectra partly overlap. There is a type of cone that absorbs mainly blue light (with an absorbance peak at 420 nm), another that absorbs green wavelengths (with an absorbance peak at 530 nm), and a third type that absorbs up into the red region (with an absorbance peak at 565 nm). When light, having a particular spectral power distribution, impinges on the retina, it activates each of the three types of cones by differing degrees. The distribution of the degrees of activation of the three types of cones is information that travels as electrochemical signals up to the visual cortex. In the visual cortex, information is encoded as a specific pattern of activity of the cortical neurons in layer 4 of the V1 area. Metameric matches occur when different spectral signals lead to the same activation patterns in the three types of cones and to the same pattern of activity in the visual cortex; thus, the different spectral signals appear to represent the same colour. Of course, the overall information about the colours within our brain is not limited to a simple correspondence between pigment activation and light spectrum. In fact, neurophysiological evidence, such as colour constancy and coloured shadows, reveal the existence of post-receptor mechanisms of colour information processing.

Fuzzy logic is a mathematically rigorous model useful to describe the human computational ability which presently uses words and imprecise reasoning. It is based on the theory of fuzzy logic. Fuzzy logic is a mathematically rigorous model useful to describe the human computational ability which presently uses words and imprecise reasoning. It is based on the theory of fuzzy logic.

Electronic Supplementary Information (ESI) available: [For instrumentation, materials and methods see the Electronic Supporting Information (ESI)]. The synthesis of 2 – 5 are described. The methodology for the determination of the photochemical quantum yields. Quantum mechanical simulations of the absorption spectra of the uncolored forms by the DFT method. The reflectance spectra of the photochromic sheets of white paper]. See DOI: 10.1039/x0xx00000x

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patterns of activity of cortical neurons that are interpreted as the same colour. The Fuzzy Inference Engine is the mechanism of transduction of the electrochemical information stored by the photoreceptor cells in the information encoded as patterns of activity of cortical neurons. This description is useful for the design of biologically inspired chemical systems for UV vision. Humans cannot perceive UV radiation because the lens and cornea of the eye absorb strongly in this wavelength region preventing UV radiation from reaching the retina. However, a wide variety of animal species show sensitivity to UV, ranging from insects to mammals. Most often, the species that see UV are those that have a specific photoreceptor peaked around 350 nm in the UV-A region.\(^{15,16}\) UV sensitivity is useful in activities as diverse as navigation, intra- and inter-species communication, foraging and circadian synchronization. A remarkable case is the mantis shrimp that has at least four types of photoreceptors for UV in addition to eight for the visible region, as befits its habitat of kaleidoscopically colourful tropical coral reefs.\(^{17}\)

In this work, we present the synthesis, study and optimized combinations of five thermally reversible photochromic compounds (1 – 5) that generate Biologically Inspired Fuzzy Logic Systems useful to transform the frequencies of the UV spectrum, invisible to us, into specific colours perceptible to the human eye. The photochromic compounds are 1,3-dihydro-1,3,3-trimethyl-8′-nitro-spiro[2H-indole-2,3′-][3H]naphth[2,1-b][1,4]oxazine (1), 2,2-bis(4-methoxyphenyl)-5,6-dimethyl-2H-naphtho[1,2-b]pyran (2), 6-morpholino-3-(4-morpholinophenyl)-3-phenyl-3H-naphtho[2,1-b]pyran (3), 3,3-diphenyl-6-morpholino-3H-naphtho[2,1-b]pyran (4), 3-(2-fluorophenyl)-3-(4-pyrrolidinophenyl)-3H-naphtho[2,1-b]pyran (5). Their structures and the colours of their solutions containing their ring-opened forms after UV irradiation are shown in Figure 1.

The absorption spectra of the closed uncoloured forms are depicted in Figure 2A. Naphthoxazine 1 has the largest absorption coefficient in the UV-A (320–400 nm) and almost in the entire UV-B region (280–320 nm). In the portion of UV-C included between 250 and 280 nm that will be considered in this work, the absorption of 1 is overwhelmed by the contributions of the other four naphthopyrans. Naphtho[2,1-b]pyrans 3 and 4 have fairly similar absorption spectra in the UV-A; in UV-B and UV-C regions, 3 absorbs more than 4 due to the presence of a morpholino group bound to one of the two phenyl rings (see Figure 1). Naphtho[1,2-b]pyran 2 is characterized by small values of the absorption coefficients in the range 387–294 nm. Compound 5 is the naphtho[2,1-b]pyran that commences absorption at the shortest wavelengths among the other photochromes. Upon UV irradiation (see Figure 2B), 1 gives rise to a merocyanine that has an absorption band with a maximum at 610 nm and its solution becomes blue; 5 generates a band having a maximum at 554 nm and its solution becomes purple; 2 produces a band in the visible region which peaks at 497 nm and its solution becomes pink; 3 gives rise to a band centred at 463 nm and its solution becomes orange; finally, 4 generates a narrow band with a maximum at 413 nm and its solution appears yellow. The band due to 1 has an absorption coefficient at 610 nm that is more than four times larger than the values of the other coloured species; it is generally accepted that photochromic oxazines typically afford ring-opened species, which are more hyperchromic and bathochromic than those derived from diarylnaphthopyrans.\(^{18-21}\)

The photochemical quantum yields (\(\Phi_{\text{UC}}\)) of the five photochromic compounds have been determined by irradiating at different wavelengths in the UV. The experimental methodology followed for their determination is described in the Electronic Supplementary Information (ESI). The results are listed in Table 1. For all compounds, \(\Phi_{\text{UC}}\) in the UV-C region is larger than \(\Phi_{\text{UC}}\) in UV-A and UV-B regions. This is particularly true in the case of naphthopyrans. In fact, naphthopyrans are known to give ultrafast electrocyclic ring opening reactions that kinetically compete with the other unreactive relaxation pathways and their photochemical quantum yields are usually wavelength-dependent.\(^{22}\) Simulations of the absorption spectra by Density Functional Theory computation (DFT, see ESI) reveal that the electronic transitions in UV-A and UV-B involve mainly the naphthopyran rings, whereas those in UV-C have charge transfer character from the naphthopyran rings to the two phenyl groups bound to the sp\(^3\) carbon atom in the pyran ring. The open forms produced by irradiating with UV have lifetimes...
of tens of seconds, which are independent of the frequency of irradiation (see data in Table 1). In particular, the open form of 5 has the shortest lifetime, lasting 19 s, whereas the open form of 4 is approximately three times more persistent.

With knowledge of the spectral and photochemical properties of the five photochromes, some or all of them can be mixed in different ratios to create chemical systems able to transform the frequencies of the UV-A, UV-B and UV-C regions into different colours. The matching criteria are founded upon two considerations. First, the absorption bands of the uncoloured forms must be conceived as input fuzzy sets and the irradiation intensity \( I(\lambda_{irr}) \) at \( \lambda_{irr} \) will belong to each of them with a degree \( \mu_{\nu_{ij}} \) given by:

\[
\mu_{\nu_{ij}} = \Phi_{PCj}(\lambda_{irr}) \cdot I_{abs}(\lambda_{irr})
\]

where \( k_{s,i} \) is the reciprocal of the \( i \)-th open form’s lifetime. The final absorption spectrum recorded at the photo-stationary state will be the sum of as many terms represented by equation (2) as there are photochromic components present in the mixture. Of course, the sum must be extended to all the wavelengths (\( \lambda_{irr} \)) belonging to the visible spectrum.

Table 1. Photochromic quantum yields (\( \Phi_{\nu} \)) determined by irradiating the five photochromic compounds in different regions of the UV spectrum, and lifetimes (\( t \)) of their coloured forms (in MeCN solution).

<table>
<thead>
<tr>
<th>Compound</th>
<th>( \Phi_{\nu} )(UV-A)</th>
<th>( \Phi_{\nu} )(UV-B)</th>
<th>( \Phi_{\nu} )(UV-C)</th>
<th>( t = 1/k_s ) (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.12</td>
<td>0.07</td>
<td>0.18</td>
<td>25</td>
</tr>
<tr>
<td>2</td>
<td>0.17</td>
<td>0.26</td>
<td>0.45</td>
<td>37</td>
</tr>
<tr>
<td>3</td>
<td>0.24</td>
<td>0.25</td>
<td>0.92</td>
<td>37</td>
</tr>
<tr>
<td>4</td>
<td>0.27</td>
<td>0.42</td>
<td>0.53</td>
<td>61</td>
</tr>
<tr>
<td>5</td>
<td>0.28</td>
<td>0.23</td>
<td>0.62</td>
<td>19</td>
</tr>
</tbody>
</table>

Many combinations of photochromes 1 - 5, containing from three to five compounds and selected by applying equations (1) and (2), have been found effective in distinguishing the three principal UV regions: UV-A (400 - 320 nm) from UV-B (320 - 280 nm) and from UV-C with \( \lambda \geq 250 \) nm. One of the best systems was a quaternary mixture involving 1, 4, 5, and 2 in concentrations of \( 5.2 \times 10^{-5} \) M, \( 7.38 \times 10^{-5} \) M, \( 1.4 \times 10^{-4} \) M, and \( 1.4 \times 10^{-4} \) M, respectively. Its discriminative power is shown in Figure 3. When the system is irradiated by frequencies belonging to UV-A, the solution becomes green; under UV-B, the solution turns to grey, and under UV-C irradiation with wavelengths longer than 250 nm, it turns to orange. The spectra recorded at the photo-stationary states, shown as the grey dashed traces in the bottom panels of Figure 3, are accurately reproducible summing the spectral contribution of each species, expressed by equation 2 (see the red dashed traces in the same panels). Equation (2) provides a powerful means to predict the observed colour when the mixture is irradiated by many UV frequencies, simultaneously. For example, in Figure 4, the experimental (grey dashed traces) and predicted (red dashed traces) spectra obtained under different polychromatic irradiation frequencies are compared.
Under direct sunlight (see top-left panel), the solution becomes orange, and the experimental spectrum can be readily reproduced if we consider that the closed forms of 1 and 4 are completely transformed into their open forms by UV-A, whereas 2 and 5 are completely converted into their coloured forms by UV-B. When we add the contribution of the radiation at 254 nm, emitted by a Hg lamp (see plot B1 in Figure 4), the solution turns red because of the higher $\Phi_{\text{tr}}$ of 2 and 5 in the UV-C region. When the solution is irradiated by skylight but not direct sunlight, it assumes a pale green colour because the spectrum of skylight is poor in UV-B frequencies (see plot C1 in Figure 4). When we add the 254 nm wavelength emitted by the Hg lamp to the spectrum of skylight, the colour of the solution turns red because the very intense UV-C radiation quantitatively transforms the closed forms of 2 and 5 to their respective open structures. In fact, 2 and 5 have higher probabilities of absorbing the 254 nm radiation because they are at higher concentration into the mixture.

The performance of the Biologically Inspired Photochromic Fuzzy Logic (BIPFUL) systems investigated in acetonitrile solutions can be extended to a solid cellulose support as in white paper. Thereafter, the BIPFUL systems described herein mimic the computing power of vertebrates and invertebrates that have different photoreceptors to distinguish between frequencies of the electromagnetic spectrum. Recently, de Silva et al., demonstrated that the parallel processing by the combination of a pH sensor and a photo-acid generator detects the edges of objects, which is a rather complex computational task normally requiring a highly organized biomolecular system. This paper constitutes a further demonstration of how simple molecular logic systems, such as the photochromic compounds designed as “optical transistors”,28-31 can give rise to high-level computing performances. We have shown that our BIPFUL systems can work not only in solution but also on a solid cellulose support such as a sheet of inexpensive filter paper. Therefore, these systems of photochromic compounds are promising for designing new devices distinguishing UV frequencies in a photochemical manner instead of photo-electrically as with the use of semiconductors.32 Such photochemical UV detectors can be directly linked to human vision through the colours they produce and can supplement the performance of our visual system, enabling a further degree of detection and discrimination.

Notes and references
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