1	Exploration of natural red-shifted rhodopsins using a machine learning-based Bayesian
2	experimental design
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#### 31 Abstract

Microbial rhodopsins are photoreceptive membrane proteins utilized as molecular tools in 32 optogenetics. In this paper, a machine learning (ML)-based model was constructed to 33 34 approximate the relationship between amino acid sequences and absorption wavelengths using ~800 rhodopsins with known absorption wavelengths. This ML-based model was specifically 35 designed for screening rhodopsins that are red-shifted from representative rhodopsins in the 36 same subfamily. Among 5,558 candidate rhodopsins suggested by a protein BLAST search of 37 38 several protein databases, 40 were selected by the ML-based model. The wavelengths of these 40 selected candidates were experimentally investigated, and 32 (80%) showed red-shift gains. 39 40 In addition, four showed red-shift gains > 20 nm, and two were found to have desirable ion-41 transporting properties, indicating that they were potentially useful in optogenetics. These findings suggest that an ML-based model can reduce the cost for exploring new functional 42 proteins. 43

#### 45 Introduction

Microbial rhodopsins are photoreceptive membrane proteins widely distributed in bacteria, 46 archaea, unicellular eukaryotes, and giant viruses<sup>1</sup>. They consist of seven transmembrane (TM) 47  $\alpha$  helices, with a retinal chromophore bound to a conserved lysine residue in the seventh helix 48 (Fig. 1a). The first microbial rhodopsin, bacteriorhodopsin (BR), was discovered in the plasma 49 membrane of the halophilic archaea *Halobacterium salinarum* (formerly called *H. halobium*)<sup>2</sup>. 50 BR forms a purple-coloured patch in the plasma membrane called purple membrane, which 51 outwardly transports H<sup>+</sup> using sunlight energy<sup>3</sup>. After the discovery of BR, various types of 52 microbial rhodopsins were reported from diverse microorganisms, and recent progress in 53 genome sequencing techniques has uncovered several thousand microbial rhodopsin genes<sup>1,4–</sup> 54 55 <sup>6</sup>. These microbial rhodopsins show various types of biological functions upon light absorption, leading to all-trans-to-13-cis retinal isomerization. Among these, ion transporters, including 56 light-driven ion pumps and light-gated ion channels, are the most ubiquitous (Fig. 1b). Ion-57 transporting rhodopsins can transport several types of cations and anions, including H<sup>+</sup>, Na<sup>+</sup>, 58 K<sup>+</sup>, halides (Cl<sup>-</sup>, Br<sup>-</sup>, I<sup>-</sup>), NO<sup>-</sup>, and SO<sub>4</sub><sup>2-1,7-9</sup>. The molecular mechanisms of ion-transporting 59 rhodopsins have been detailed in numerous biophysical, structural, and theoretical studies<sup>1</sup>. 60

In recent years, many ion-transporting rhodopsins have been used as molecular tools in 61 optogenetics to control the activity of animal neurons optically in vivo by heterologous 62 expression<sup>10</sup>, and optogenetics has revealed various new insights regarding the neural network 63 relevant to memory, movement, and emotional behaviour<sup>11–14</sup>. However, strong light scattering 64 by biological tissues and the cellular toxicity of shorter wavelength light make precise optical 65 control difficult. To circumvent this difficulty, new molecular optogenetics tools based on red-66 shifted rhodopsins that can be controlled by weak scattering and low toxicity longer-67 wavelength light are urgently needed. Therefore, many approaches to obtain red-shifted 68 rhodopsins, including gene screening, amino acid mutation based on biophysical and structural 69

insights, and the introduction of retinal analogs, have been reported<sup>15-17</sup>. Recently, a new 70 71 method using a chimeric rhodopsin vector and functional assay was reported to screen the 72 absorption maximum wavelengths ( $\lambda_{max}$ ) and proton transport activities of several microbial rhodopsins present in specific environments<sup>18</sup>. This method identified partial sequences of red-73 shifted yellow (560-570 nm)-absorbing proteorhodopsin (PR), the most abundant outward H<sup>+</sup>-74 pumping bacterial rhodopsin subfamily, from the marine environment. Although these works 75 identified several red-shifted rhodopsins<sup>14,15,17,19</sup>, those showing ideally red-shifted absorption 76 and high ion-transport activity sufficient for optical control in vivo have yet to be obtained. 77

As an alternative approach, we recently introduced a data-driven machine learning (ML)-78 based approach<sup>20</sup>. In the previous study, we demonstrated how accurately the absorption 79 80 wavelength of rhodopsins could be predicted based on the amino acid types on each position of the seven TM helices<sup>20</sup>. We constructed a database containing 796 wild-type (WT) 81 rhodopsins and their variants, the  $\lambda_{max}$  of which had been reported in earlier studies. Then, we 82 demonstrated the prediction performance of the ML-based prediction model using a data-83 84 splitting approach, i.e., the data set was randomly divided into a training set and a test set; the former was used to construct the prediction model, and the latter was used to estimate the 85 prediction ability. The results of this "proof-of-concept" study suggested that the absorption 86 87 wavelengths of an unknown family of rhodopsins could be predicted with an average error of  $\pm 7.8$  nm, which is comparable to the mean absolute error of  $\lambda_{max}$  estimated by the hybrid 88 quantum mechanics/molecular mechanics  $(OM/MM)^{21}$ method. 89 Considering the computational cost of both approaches, the ML-based approach is much more efficient than 90 QM/MM approach, while the latter provides insights on the physical origin controlling  $\lambda_{max}$ . 91

Encouraged by this result, in this study, we used an ML-based approach to screen more redshifted rhodopsins from among 3,064 new candidates collected from public databases (nonredundant and metagenomic rhodopsin genes from the National Center for Biotechnology

95 Information [NCBI] and Tara Oceans data sets) for which the absorption wavelengths have not been investigated. The goal of the present study was to identify rhodopsins with a  $\lambda_{max}$  longer 96 than the wavelengths of representative rhodopsins in each subfamily of microbial rhodopsins 97 for which the  $\lambda_{max}$  has already been reported (base wavelengths). Here, we call the red-shift 98 change in the wavelength from the base wavelength the "red-shift gain". We focus on the 99 problem of identifying rhodopsins with large red-shift gains because this would lead to the 100 101 identification of amino acid types and residue positions that play important roles in red-shifting 102 absorption wavelengths. In addition, in optogenetics applications, it is practically important to have a wide variety of ion-pumping rhodopsins from each subfamily to construct a new basis 103 104 for rhodopsin toolboxes with red-shifted absorption and various types of ion species that can be transported. To screen rhodopsins that would have large red-shift gains, it is necessary to 105 consider the uncertainty of prediction in the form of "predictive distributions"<sup>22</sup>. By using 106 predictive distributions, it is possible to consider appropriately the "exploration-exploitation 107 trade-off" in screening processes<sup>23,24</sup>, where exploration indicates an approach that prefers 108 109 candidates with larger predictive variances, and exploitation indicates an approach that prefers 110 candidates with longer predictive mean wavelengths (Fig. 2). In this paper, we employ a 111 Bayesian modeling framework to compute the predictive distributions of candidate rhodopsin 112 red-shift gains. We then consider an exploration-exploitation trade-off by selecting candidate rhodopsins based on a criterion called "expected red-shift gains". 113

In this paper, we updated the ML-based model used in our previous study<sup>20</sup> so that it could properly compute expected red-shift gains and applied this new model to 3,064 ion-pumping rhodopsin candidates derived from archaeal and bacterial origins that can be easily expressed in *Escherichia coli* (Fig. 1b). We then selected 66 candidates for which the expected gains were > 10 nm, and experimentally investigated their wavelengths by introducing the synthesized rhodopsin genes into *E. coli*. Of these 66 selected candidates, 40 showed significant colouring 120 in E. coli cells, 32 showed actual red-shift gains, seven showed blue-shifts, and one showed no change, suggesting that our ML-based model enables more efficient screening of red-shifted 121 rhodopsin genes compared with random choice (i.e., 80.5% [32/40] of the selected candidates 122 showed red-shift gains with  $p < 10^{-3}$  in a binomial test). We then investigated the ion-123 transportation properties of the rhodopsins whose red-shift gains were > 20 nm, and found that 124 some actually had desired ion-transporting properties, suggesting that they (and their variants) 125 could potentially be used as new optogenetics tools. Furthermore, the differences in the amino-126 acid sequences of the newly examined rhodopsins and the representative ones in the same 127 128 subfamily could be used for further investigation of the red-shifting mechanisms.

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#### 130 **Results**

# 131 Construction of an ML-based model for computing expected red-shift gain

To compute the expected red-shift gains of a wide variety of rhodopsins, we updated various 132 aspects of the ML model used in our previous study<sup>19</sup>. Figure 3 shows a schematic of the 133 134 updating procedure. First, we added 97 WT microbial rhodopsins and their variants for which the  $\lambda_{\rm max}$  had recently been reported in the literature or determined by our experiments, to a 135 previously reported data set<sup>20</sup>. In other words, the new training data set consisted of the amino 136 acid sequences and  $\lambda_{max}$  of 893 WT microbial rhodopsins and their variants (Extended Data 137 Table 1). Second, the new ML model used only N = 24 residues located around the retinal 138 chromophore (Extended Data Figure 1) because our previous study<sup>19</sup> indicated that amino acid 139 140 residues at these 24 positions play significant roles in predicting absorption wavelengths (Fig. 141 3a). Third, M = 1818 amino acid physicochemical features (Extended Data Table 2) were used as inputs in the ML model, as opposed to the amino acid types used in the previous ML 142 model. This enabled us to predict the absorption wavelengths of a wide range of target 143 144 rhodopsins that contain unexplored amino acid types in the training data at certain positions.

145 Therefore, an amino acid sequence is transformed into an  $M \times N = 432$  dimensional feature 146 vector  $\mathbf{x} \in \mathbb{R}^{MN}$  by concatenating  $x_{i,j}$ , the *j*-th feature of the *i*-th residue (Fig. 3b). We 147 consider a linear prediction model  $f(\mathbf{x}) = \mu + \sum_{i=1}^{N} \sum_{j=1}^{M} \beta_{i,j} x_{i,j}$ , where  $\beta_{i,j}$  is the 148 parameter for the *j*-th feature of the *i*-th residue, and  $\mu$  is the intercept term.

Finally, to consider the exploration-exploitation trade-off appropriately in the screening 149 process, we introduce a Bayesian modeling framework, which allows us to compute the 150 predictive distributions of red-shift gains. Specifically, we employed Bayesian sparse modeling 151 called BLASSO<sup>25</sup> (see the Methods section for details). This enables us to provide not only the 152 mean, but also the variance of the predicted wavelengths. Unlike classical regression analysis, 153 BLASSO regards the model parameters  $\beta_{i,i}$  and  $\mu$  as random variables generated from 154 underlying distributions, as illustrated in Figure 3c. Therefore, the wavelength prediction f(x)155 is also represented as a distribution. The red-shift gain is defined as gain = max(f(x) - f(x))156  $\lambda_{\text{base}}$ , 0), where  $\lambda_{\text{base}}$  is the wavelength of the representative rhodopsin in the same subfamily 157 whose  $\lambda_{max}$  has been experimentally determined and reported in the literature (Extended Data 158 Table 3). Note that the red-shift gain is positive if f(x) is greater than  $\lambda_{\text{base}}$ ; otherwise, it 159 takes the value of zero. Since f(x) is regarded as a random variable in BLASSO, the red-shift 160 gain is also regarded as a random variable. Therefore, we employ the expected value of the red-161 shift gain, denoted by  $\mathbb{E}[gain]$ , as the screening criterion where  $\mathbb{E}$  represents the expectation 162 of a random variable. Illustrative examples of  $\mathbb{E}[gain]$  are shown in Figure 3d. Unlike the 163 simple expectation of the wavelength prediction  $\mathbb{E}[f(x)]$ ,  $\mathbb{E}[gain]$  depends on the variance 164 165 of the predictive distribution. This encourages the exploration of rhodopsin candidates having large uncertainty (for exploration), as opposed to only those having longer wavelengths with 166 high confidence (for exploitation). 167

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#### 169 Screening potential red-shifted microbial rhodopsins based on expected red-shift gains

170 The target data set to explore red-shifted microbial rhodopsins was constructed by collecting putative microbial rhodopsin genes collected by a protein BLAST (blastp) search<sup>26</sup> of the NCBI 171 non-redundant protein and metagenome databases<sup>27</sup>, as well as the *Tara* Oceans microbiome 172 and virome databases<sup>28</sup>. As a result, we obtained a non-redundant data set of 5,558 microbial 173 rhodopsin genes (Fig. 1b). The sequences were aligned by ClustalW and categorized to 174 175 subfamilies of microbial rhodopsins based on the phylogenic distances, as reported previously<sup>29</sup>. Among these, 3,064 rhodopsin genes from bacterial and archaeal origins were 176 extracted because their  $\lambda_{max}$  can be easily measured by expressing in *E. coli* cells. We calculated 177 178 the  $\mathbb{E}[gain]$  of these 3,064 genes (Extended Data Table 4), and then selected 66 genes of 179 putative light-driven ion pump rhodopsins showing an  $\mathbb{E}[gain] > 10$  nm for further experimental evaluation, as ion pump rhodopsins can be used as new optogenetics tools. 180

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# 182 Experimental measurement of the absorption wavelengths of microbial rhodopsins 183 showing high red-shift gains

We synthesized the selected 66 genes that showed an  $\mathbb{E}[gain] > 10$  nm. These were then 184 185 introduced into E. coli cells, and the proteins expressed in the presence of 10 µM all-trans retinal. As a result, 40 E. coli cells showed significant colouring, indicating significant 186 187 expression of folded protein, and their  $\lambda_{max}$  were determined by observing ultraviolet (UV)visible absorption changes upon bleaching of the expressed rhodopsins through a hydrolysis 188 reaction of their retinal with hydroxylamine, as previously reported<sup>20</sup> (Fig. 4). The observed 189 gains were compared with the E[gain] shown in Table 1. A full list of unexpressed genes is 190 shown in Extended Data Table 5. In total, 32 of 40 genes showed a longer wavelength than 191 192 their base wavelength (that is, positive red-shift gain) (Fig. 5), suggesting that our ML-based model can significantly improve the efficiency of screening to explore new red-shifted 193 microbial rhodopsins compared with random sampling (p < 0.0002). 194

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#### 196 Ion-transport function of red-shifted microbial rhodopsins

Overall, four of the 40 rhodopsins showed red-shifted absorption > 20 nm compared with the 197 base wavelengths (Table 1): three were halorhodopsins (HRs) from bacterial species<sup>9,30,31</sup> (to 198 distinguish classical HRs from archaeal species, these are hereafter referred to as bacterial-199 halorhodopsins [BacHRs]), and one was a PR<sup>32</sup>. Their ion-transport activities were then 200 investigated by expressing in *E. coli* cells and observing the pH change in external solvent (Fig. 201 202 6). Upon light illumination, BacHRs from *Rubrivirga marina* and *Myxosarcina* sp. GI1 showed significant alkalization of external solvent, which was enhanced by addition of the 203 protonophore (CCCP), which increases the H<sup>+</sup> permeability of the cell membrane, and the light-204 dependent alkalizations disappeared when anions were exchanged from Cl<sup>-</sup> to  $SO_4^{2-}$ , indicating 205 206 that these were light-driven Cl<sup>-</sup> pumps, similar to other rhodopsins in the same BacHR subfamily<sup>9,30</sup>. By contrast, *Cyanothece* sp. PCC 7425 did not show any significant transport. 207 While no transporting function can be attributed to the heterologous expression in E. coli, it 208 209 would have considerably different molecular properties from other BacHRs. PRs from a metagenome sequence (ECV93033.1) showed acidification of external solvent that was 210 211 abolished by the addition of CCCP and was independent from ionic species in the solvent. Hence, this was a new red-shifted outward H<sup>+</sup> pump compared with typical PRs whose  $\lambda_{max}$ 212 are present at ca. 520 nm<sup>32</sup>. These light-driven ion-pumping rhodopsins with red-shifted  $\lambda_{max}$ 213 214 have the potential to be applied as new optogenetics tools, and thus, warrant further study in the near future. 215

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#### 217 Discussion

218 Microbial rhodopsins show a wide variety of  $\lambda_{max}$  by changing steric and electrostatic 219 interactions between all-*trans* retinal chromophores and surrounding amino acid residues. An 220 understanding of the colo<u>u</u>r-tuning rule enables more efficient screening and the design of new 221 red-shifted rhodopsins that have value as optogenetics tools, and our ML-based data-driven 222 approach therefore provides a new basis to identify colour-regulating factors without 223 assumptions.

224 We previously demonstrated that an ML-based model based on ~800 experimental results could predict the  $\lambda_{max}$  of microbial rhodopsins with an average error of  $\pm 7.8$  nm. Encouraged 225 226 by this result, in the present study, we constructed a new ML-based model to compute expected red-shift gains for a wide range of unknown families of microbial rhodopsins. As a result, 33 227 of 40 microbial rhodopsins were found to have red-shifted absorption compared with the base 228 wavelengths of each subfamily of microbial rhodopsins (Table 1), suggesting that our data-229 driven ML approach can screen red-shifted microbial rhodopsin genes more efficiently than 230 231 random choice.

By considering the exploration-exploitation trade-off, that is, to consider not only the 232 expected value of the prediction, but also the uncertainty, it was possible to construct a red-233 234 shift protein screening process, as shown in Figure 7. Figure 7a shows the relationships 235 between the prediction uncertainty (as measured by the standard deviation) and the observed red-shift gains. It can be seen that rhodopsins with red-shift gain are found in areas of not only 236 237 low (small standard deviation), but also high prediction uncertainty (large standard deviation). Figure 7b shows the two-dimensional projection of the d = 432 dimensional feature space by 238 principal component analysis. It can be seen that red-shift gains (red) are found for target 239 proteins not only close to training proteins (green), but also far from training proteins. Figure 240 8 shows that the observed wavelengths and red-shift gains tend to be smaller than the predicted 241 242 ones. We conjecture that these differences between the observed and predicted wavelengths and red-shift gains are due to modeling errors, possibly caused by a lack of sufficient 243 information (e.g., three-dimensional structures) and modeling flexibility (e.g., nonlinear 244

effects); in other words, rhodopsins having high prediction values partly by modeling errors
have a high chance of being selected. Therefore, it would be valuable to develop a statistical
methodology to eliminate selection bias due to modeling errors.

Four rhodopsins showed red-shifted absorption > 20 nm than the base wavelength, three of 248 which showed light-driven ion-transport function. Interestingly, while one BacHR from 249 250 *Rubrivirga marina* (accession No.: WP 095512583.1) showed a 40-nm longer  $\lambda_{max}$  (577 nm) than the base wavelength, another 11-nm red-shifted BacHR (WP 095509924.1) was also 251 252 identified from the same bacteria (Table 1). These BacHRs are highly similar to each other (55.2% identity and 70.6% similarity), and only four of 24 amino acid residues around the 253 254 retinal chromophore differ. Hence, R. marina evolved two BacHRs with 29-nm different  $\lambda_{max}$ by small amino acid replacement; the amino acid residue(s) responsible for this color-tuning 255 256 should be investigated in the future.

The differences in amino acids in three of 24 retinal-surrounding residues are known to 257 play a color-tuning role in natural rhodopsins without affecting their biological function. These 258 259 correspond to positions 93, 186, and 215 in BR (BR Leu93, Pro186, and Ala215, respectively)<sup>16</sup>. 260 Position 93 is known to be diversified in the PR family (the well-known position 105 in PRs). Green-light-absorbing PRs (GPRs) have leucine as a BR, whereas glutamine is conserved in 261 262 blue-light-absorbing PRs<sup>4,18</sup>. This colour-tuning effect by the difference between leucine and glutamine is known as the "L/Q-switch"<sup>33</sup>. Interestingly, while 29.8% of 3,064 candidate genes 263 have glutamine at this position, all 40 genes whose large red-shift gains were suggested by our 264 ML-based model have amino acids other than glutamine, which suggests that our ML-based 265 model avoided the genes having glutamine at position 93. Especially, 12 (37.5%) of 32 genes 266 267 that actually showed red-shifted absorption compared with the base wavelengths had methionine at this position (Extended Data Figure 2), which is substantially higher than the 268 proportion of methionine-conserving genes in the 3,064 candidates (16.1%). The red-shifting 269

effect of the L-to-M mutation of this residue in GPRs previously reported<sup>33</sup> and the current 270 271 result imply that many rhodopsins have evolved methionine to absorb light with longer wavelengths. Position 215 in BR is also known to have a colour-tuning role. The mutation from 272 alanine to threenine or serine (A/TS switch) has a blue-shifting effect of  $9-20 \text{ nm}^{16,34-36}$ . Five 273 274 of seven genes that showed blue-shifted  $\lambda_{max}$  compared with the base wavelengths have threonine or serine at this position, suggesting that these types of genes should be avoided to 275 explore red-shifted rhodopsins. By contrast, asparagine was conserved in more than half 276 277 (58.4%) of the 3,064 candidate genes, especially in those belonging to the PR subfamily. A significant portion (37.5%) of the genes with red-shifted absorption compared with the base 278 279 wavelengths also had asparagine at this position (Extended Data Figure 2). The A-to-N mutation at this position had a smaller effect  $(4-7 \text{ nm})^{20,35}$  than that of the A-to-S/T mutation; 280 thus, the difference between alanine and asparagine is not so critical to explore red-shifted 281 rhodopsins. Position 186 in BR is proline in most microbial rhodopsins (in 98.7% of the 3.064 282 candidate genes), and the mutation to non-proline amino acids induces red-shift of absorption<sup>16</sup>. 283 We identified sodium pump rhodopsin (NaR) from Parvularcula oceani, which also has a 284 285 threonine at this position, and showed 10-nm longer absorption than the base wavelength. Although genes having non-proline amino acids are rare in nature, it would be beneficial to 286 identify new red-shifted rhodopsins. These results indicate that ML-based modelling can 287 288 provide insights for identifying new functional tuning rules for proteins based on specific amino acid residues. 289

The number of reported microbial rhodopsin genes is rapidly increasing because of the development of next-generation sequencing techniques and microbe culturing methods. New microbial rhodopsins with molecular characteristics suitable for optogenetics applications are expected to be included in upcoming genomic data. Our ML-based model could be expected to reduce the costs associated with identifying red-shifted rhodopsins from these data.

Especially, we expect that our ML-based model could be applied to ion channel and enzymatic rhodopsins, which were not a focus of this study because of their eukaryotic origins; however, their use in optogenetics research could help identify more useful optogenetics tools with redshifted absorption in the future.

- 299
- 300 Methods

#### 301 Construction of training and target data sets

In this study, we constructed a new training data set (Extended Data Table 1) by adding 97 genes for which the  $\lambda_{max}$  had recently been reported in the literature or determined by our experiments, to a previously reported data set<sup>20</sup>. The sequences were aligned using ClustalW<sup>37</sup> and the results were manually checked to avoid improper gaps and/or shifts in the TM parts. The aligned sequences were then used for ML-based modeling.

To collect microbial rhodopsin genes for the training data set, BR<sup>38</sup> and heliorhodopsin 307 48C12<sup>39</sup> sequences were used as queries for searching homologous amino acid sequences in 308 NCBI non-redundant protein sequences and metagenomic proteins<sup>27</sup> and the *Tara* Oceans 309 microbiome and virome database<sup>28</sup>. Protein BLAST (blastp)<sup>26</sup> was used for the homology 310 search, with the threshold E-value set at < 10 by default, and sequences with > 180 amino acid 311 residues were collected. All sequences were aligned using ClustalW<sup>37</sup>. The highly diversified 312 313 C-terminal 15-residue region behind the retinal binding Lys (BR K216) and long loop of HeR 314 between helices A and B were removed from the sequences to avoid unnecessary gaps in the alignment. The successful alignment of the TM helical regions, especially the 3rd and 7th 315 helices, was checked manually. The phylogenic tree was drawn using the neighbor-joining 316 method<sup>40</sup>, and the microbial rhodopsin subfamilies were categorized based on the phylogenetic 317 distances, as reported previously<sup>29</sup>. Based on the phylogenetic tree, 3,064 putative ion-pumping 318 319 rhodopsin genes from bacterial and archaeal origins were extracted, and their aligned sequences

320 were used as the training data set for the prediction of  $\lambda_{\text{max}}$ .

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#### 322 ML modeling

Suppose that we have *K* pairs of an amino acid sequence and an absorption wavelength  $\left\{\left(\boldsymbol{x}^{(k)}, \lambda_{\max}^{(k)}\right)\right\}_{k=1}^{K}$ , where  $\boldsymbol{x}^{(k)} \in \mathbb{R}^{MN}$  is the feature vector of the *k*-th amino-acid sequence and  $\lambda_{\max}^{(k)} \in \mathbb{R}$  is the absorption wavelength of the *k*-th rhodopsin protein. The least-absolute shrinkage selection operator (LASSO) is a standard regression model in which important regression coefficients can be automatically selected by the penalty on the absolute value of the coefficient, as follows:

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$$\min_{\mu,\beta} \sum_{k=1}^{K} \left( \lambda_{\max}^{(k)} - \mu - \sum_{i=1}^{M} \sum_{j=1}^{N} \beta_{i,j} x_{i,j}^{(k)} \right)^{2} + \gamma \sum_{i=1}^{M} \sum_{j=1}^{N} |\beta_{i,j}|,$$

where  $\boldsymbol{\beta} \in \mathbb{R}^{MN}$  is a vector of  $\beta_{i,j}$  and  $\gamma > 0$  is the regularization parameter. BLASSO is a Bayesian extension of LASSO for which the model is defined through the following random variables:

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$$\lambda_{\max}^{(k)} \sim N(\mu + \boldsymbol{\beta}^{\mathsf{T}} \boldsymbol{x}^{(k)}, \sigma^2),$$

$$\boldsymbol{\beta} \sim \pi(\boldsymbol{\beta} \mid \sigma^2),$$

where  $N(\mu, s^2)$  is a Gaussian distribution with mean  $\mu$  and variance  $s^2$ , and  $\pi(\beta | \sigma^2) =$   $\Pi_{i=1}^{M} \prod_{j=1}^{N} \frac{\gamma}{2\sqrt{\sigma^2}} e^{-\gamma |\beta_{i,j}|/\sqrt{\sigma^2}}$  is the conditional Laplace prior. In this model, the maximum of the conditional distribution of the parameter  $\beta | \{(\mathbf{x}^{(k)}, \lambda_{\max}^{(k)})\}_{k=1}^{K}, \lambda, \sigma$  is equivalent to the LASSO<sup>41</sup> estimator. For the computational details, see the original paper<sup>25</sup>. Since the resulting predictive distribution of  $f(\mathbf{x})$  is not analytically tractable, the parameters  $\beta$  and  $\mu$  are sampled from the estimated distribution T = 10,000 times. For each candidate  $\mathbf{x}$ , we approximately obtain  $\mathbb{E}[\text{gain}]$  by

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$$\mathbb{E}[\text{gain}] \approx \frac{1}{T} \sum_{t=1}^{T} \max(\mu^{(t)} + \boldsymbol{\beta}^{(t)\top} \boldsymbol{x} - \lambda_{\text{base}}, \boldsymbol{0}),$$

342 where  $\mu^{(t)}$  and  $\beta^{(t)}$  are the *t*-th sampled parameters.

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#### 345 **Protein expression**

The synthesized genes of microbial rhodopsins codon-optimized for *E. coli* (Genscript, NJ) were incorporated into the multi-cloning site in the pET21a(+) vector (Novagen, Merck KGaA, Germany). The plasmids carrying the microbial rhodopsin genes were transformed into the *E. coli* C43(DE3) strain (Lucigen, WI). Protein expression was induced by 1 mM isopropyl  $\beta$ -D-

1-thiogalactopyranoside (IPTG) in the presence of 10  $\mu$ M all-*trans* retinal for 4 h.

351

# 352 Measurement of the absorption spectra and $\lambda_{max}$ of rhodopsins by bleaching with 353 hydroxylamine

E. coli cells expressing rhodopsins were washed three times with a solution containing 100 354 mM NaCl and 50 mM Na<sub>2</sub>HPO<sub>4</sub> (pH 7). The washed cells were treated with 1 mM lysozyme 355 356 for 1 h and then disrupted by sonication for 5 min (VP-300N; TAITEC, Japan). To solubilize the rhodopsins, 3% n-dodecyl-D-maltoside (DDM, Anatrace, OH) was added and the samples 357 were stirred for overnight at 4 °C. The rhodopsins were bleached with 500 mM hydroxylamine 358 and subjected to yellow light illumination ( $\lambda > 500$  nm) from the output of a 1-kW 359 360 tungsten-halogen projector lamp (Master HILUX-HR; Rikagaku) through coloured glass (Y-52; AGC Techno Glass, Japan) and heat-absorbing filters (HAF-50S-15H; SIGMA KOKI, 361 Japan). The absorption change upon bleaching was measured by a UV-visible spectrometer (V-362 363 730; JASCO, Japan).

364

#### 365 Ion-transport assay of rhodopsins in E. coli cells

366	To assay the ion-transport activity in E. coli cells, the cells carrying expressed rhodopsin were
367	washed three times and resuspended in unbuffered 100 mM NaCl. A cell suspension of 7.5 mL
368	at $OD_{660} = 2$ was placed in the dark in a glass cell at 20 °C and illuminated at $\lambda > 500$ nm from
369	the output of a 1 kW tungsten-halogen projector lamp (Rikagaku, Japan) through a long-pass
370	filter (Y-52; AGC Techno Glass, Japan) and a heat-absorbing filter (HAF-50S-50H; SIGMA
371	KOKI, Japan). The light-induced pH changes were measured using a pH electrode (9618S-
372	10D; HORIBA, Japan). All measurements were repeated under the same conditions after the
373	addition of 10 µM CCCP.
374	
375	Reporting Summary
376	Further information on experimental design is available in the Nature Research Reporting
377	Summary linked to this article.
378	
379	Data Availability
380	Data supporting the findings of this manuscript are available from the corresponding author

381 upon reasonable request.

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472		

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481

#### 482 Author contributions

K.I., R.G., O.B., and H.K. contributed to the study design; K.I., D.Y., K.Y., and O.B. collected 483 sequences of non-redundant and metagenomic rhodopsin genes from the GenBank and Tara 484 Oceans metagenomic data sets and conducted multiple amino-acid alignments of rhodopsins; 485 M.Karasuyama, Y.I., and I.T. constructed the machine learning method to estimate the 486 absorption wavelengths of microbial rhodopsins; R.N. constructed DNA plasmids of microbial 487 rhodopsins and introduced them into E. coli cells; R.N., K.M., and T.N. conducted expressions 488 489 of microbial rhodopsins in *E. coli* cells and determined their  $\lambda_{max}$  by hydroxylamine bleaching; 490 M.Konno carried out the ion-transport assay of rhodopsins in E. coli cells; K.I., M.Karasuyama., H.K., and I.T. wrote the paper; All authors discussed and commented on the manuscript. 491

# 493 **Competing interests**

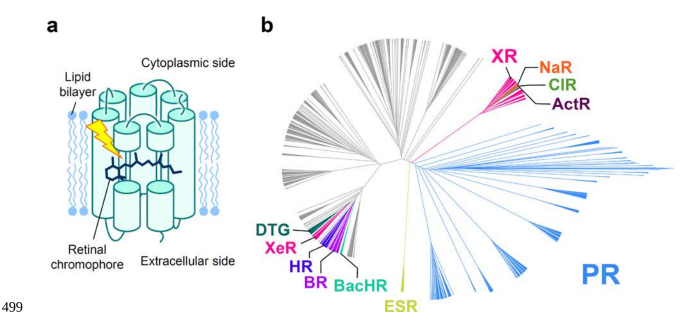
494 The authors declare no competing interests.

# 496 **Table**

# 497 Table 1. Predicted and observed gains of 40 microbial rhodopsins expressed in *E. coli*.

Origin	Accession	Subfamily	Motif	Base wavelength / nm	E [gain]	Observed wavelength / nm	(Observed wavelength) – (base wavelength) / nm
Rubricoccus marinus	WP 094550238.1	BacHR	TSA	537	40.7	541	4
Rubrivirga marina	WP 095509924.1	BacHR	TSA	537	39.8	548	11
Rubrivirga marina	WP 095512583.1	BacHR	TTD	537	35.5	577	40
Bacillus sp. CHD6a	WP 082380780.1	XeR	DTA	565	35.3	566	1
Bacillus horikoshii	WP 063559373.1	XeR	DTA	565	35.3	565	0
Cyanothece sp. PCC 7425	WP 012628826.1	BacHR	TSV	537	32.9	566	29
Cyanobacterium TDX16	OWY65757.1	BacHR	TSD	537	32.9	546	9
Myxosarcina sp. GI1	WP 052056058.1	BacHR	TTV	537	31.2	557	20
Nanohaloarchaea archaeon SW 7 43 1	PSG98511.1	XeR	DSA	565	29.2	572	7
Metagenome sequence	SAMEA2621839 1737175 2	CIR	NTQ	530	25.7	520	-10
Metagenome sequence	SAMEA2620666 5055 4	ClR	NTQ	530	25.1	525	-5
Nonlabens sp. YIK11	AIG86802.2	PR	DTE	520	21.5	531	11
Metagenome sequence	SAMEA2622673 750013 58	CIR	NTQ	530	21.4	534	4
Metagenome sequence	EBN24473.1	PR	DTE	520	20.0	525	5
Metagenome sequence	SAMEA2620404 88891 6	PR	DTE	520	20.0	527	7
Parvularcula oceani	WP_051881578.1	NaR	NDQ	525	19.7	534	9
Rubrobacter aplysinae	WP 084709429.1	DTG	DTG	535	19.5	541	6
Metagenome sequence	SAMEA2619531 1917517 3	PR	DTE	520	18.0	537	17
Metagenome sequence	SAMEA2622766 213679 12	XeR	DSA	565	17.8	572	7
Reinekea forsetii	WP 100255947.1	PR	DTE	520	17.1	524	4
Bacteroidetes bacterium	PSR14004.1	PR	DTE	520	15.4	537	17
Metagenome sequence	SAMEA2620980 19116 14	PR	DTE	520	15.4	536	16
Hassallia byssoidea VB512170	KIF37192.1	BacHR	TSD	537	15.1	535	-2
Erythrobacter gangjinensis	WP 047006274.1	NaR	NDQ	525	13.7	531	6
Pontimonas salivibrio	WP 104913209.1	PR	DTE	520	12.2	538	18
Cyanobacteria bacterium QH 1 48 107	PSO50292.1	CyanDTE	DTD	545	12.0	548	3
Kineococcus radiotolerans	WP 011981580.1	ActR	DTE	540	11.2	536	-4
Sphingopyxis baekryungensis Sphingobacteriales	WP 022671827.1	CIR	NTQ	530	11.0	518	-12
bacterium BACL12 MAG120802bin5	KRP08428.1	PR	DTE	520	10.9	531	11
Metagenome sequence	SAMEA2621401 1198262 5	PR	DTE	520	10.9	534	14
Spirosoma oryzae	WP 106137740.1	NaR	NDQ	525	10.8	533	8
Aliterella atlantica	WP 045053084.1	BacHR	TSD	537	10.8	533	-4
Rosenbergiella nectarea	WP 092678153.1	DTG	DTG	535	10.8	533	-2
Metagenome sequence	SAMEA2620980 1827033 1	PR	DTE	520	10.4	537	17
Fluviicola sp. XM24bin1	PWL28924.1	PR	DTE	520	10.4	538	18
Metagenome sequence	SAMEA2622173 654706 7	PR	DTE	520	10.4	530	10
Metagenome sequence	SAMEA2619399 1397592 7	PR	DTE	520	10.4	529	9
Sphingomonas sp. Leaf34	WP 055875688.1	DTG	DTG	535	10.3	540	5
Sphingomonas sp. Leaf38	WP 056475157.1	DTG	DTG	535	10.3	540	5
Metagenome sequence	ECV93033.1	PR	DTE	520	10.3	542	22

# 498 Figures

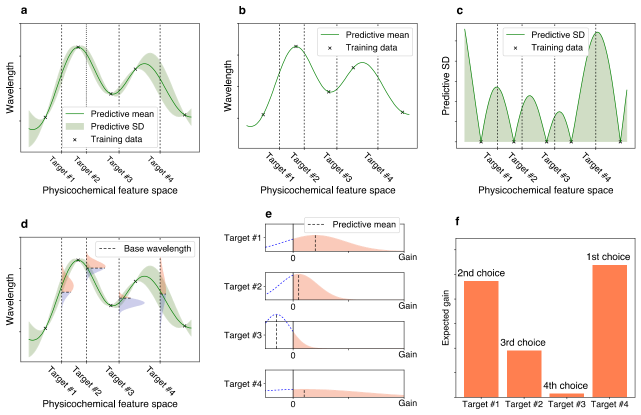


500

## 501 Fig. 1. Structure and phylogenetic tree of microbial rhodopsins.

502 **a** Schematic structure of microbial rhodopsins. **b** Phylogenic tree of microbial rhodopsins. The

- subfamilies of light-driven ion-pump rhodopsins targeted in this study are differently coloured;
- 504 non-ion-pump microbial rhodopsins are shown in grey.



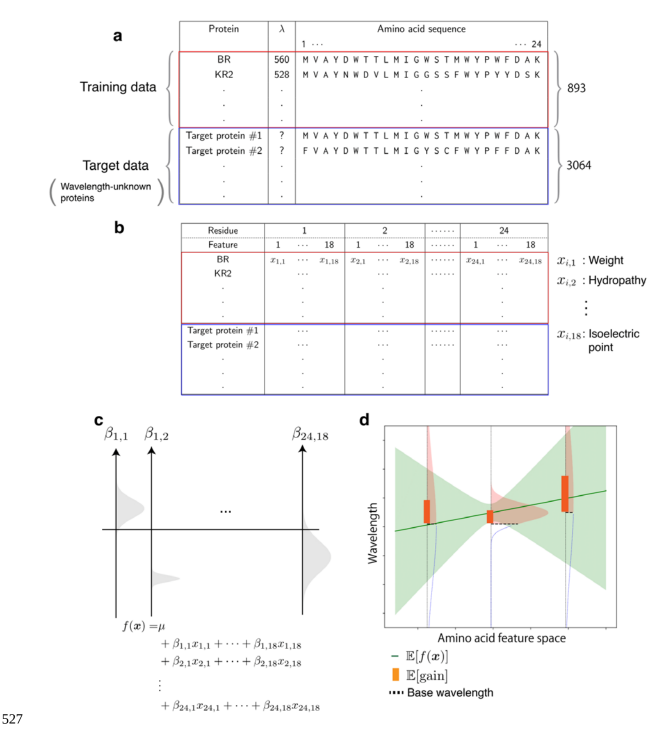
506 Fig. 2. Illustrations of exploration-exploitation for screening rhodopsins with red-shift

507 gain.

505

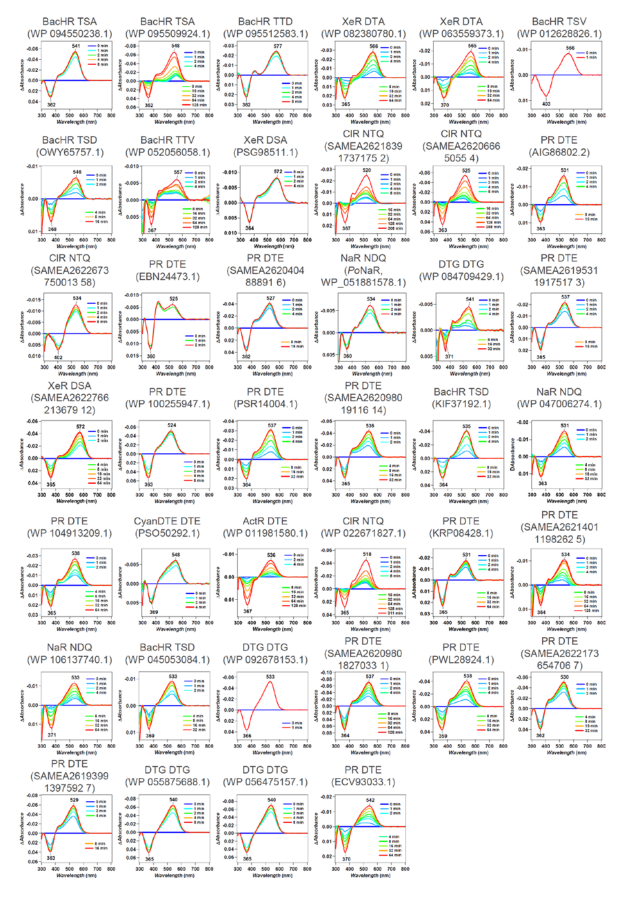
**a** Bayesian prediction model constructed using the current training data (black crosses). The 508 prediction model is represented by the predictive mean and predictive standard deviation (SD). 509 The horizontal axis schematically illustrates the space of proteins defined through 510 physicochemical features. The four vertical dotted lines indicate target proteins (candidates to 511 512 synthesize). **b** Predictive mean. This function is defined as the expected value of the probabilistic prediction by the Bayesian model. c Predictive SD. Since the predictive SD 513 represents the uncertainty of the prediction, it has a larger value when the training data points 514 do not exist nearby. **d** The distributions on the vertical dotted lines represent the predictive 515 516 distributions, and the horizontal dashed lines are the base wavelengths of the target points. The base wavelength is different for each target point because it depends on the subfamily of the 517 518 protein. e The density of the predictive distribution of each target protein on its red-shift gain value. The gain is defined as the predicted wavelength subtracted by the base wavelength, and 519

520	if it is negative, the value is truncated as 0. This can be seen as a "benefit" that can be obtained
521	by observing the target protein. <b>f</b> Expected value of the red-shift gain. This provides a ranking
522	list from which the next candidates to be experimentally investigated can be determined. Target
523	#4 has the largest expected gain, although target #1 has the largest increase in the predictive
524	mean compared with base wavelength in e. Because of its larger SD (as shown in a, c, d, and
525	e), target #4 is probabilistically expected to have a larger gain than the other targets.
506	



528 Fig. 3. Overview of the ML-based exploration of natural red-shifted rhodopsins.

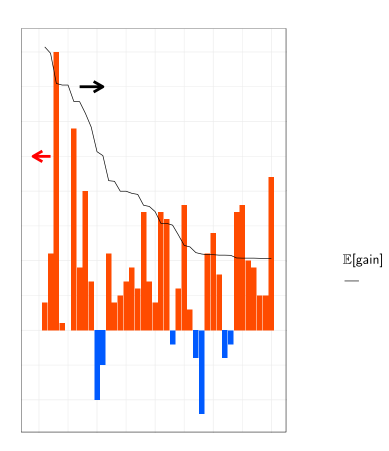
**a** Using existing experimental data, a training data set consisting of pairs of a wavelength  $\lambda_{max}$ and an amino acid sequence was constructed. A particular focus was placed on the 24 amino acid residues around the retinal chromophore to build an ML-based prediction model. A set of protein sequences with no known wavelength was also collected as target proteins. **b** All amino acid sequences were transformed into physicochemical features, leading to  $24 \times 18 = 432$  534 dimensional numerical representations of each protein. c A linear regression model was constructed using the Bayesian approach. Each regression coefficient  $\beta_{i,j}$  was estimated as a 535 distribution (shown as a gray region). The broadness of these distributions represent the 536 uncertainty of the current estimation. d The expected red-shift gain values were evaluated for 537 the target proteins. The green region is the standard deviation of the prediction. The red shaded 538 region in the vertical distribution corresponds to the probability that the wavelength is larger 539 than the base wavelength (dashed line), which is determined by the subfamily of the microbial 540 541 rhodopsin. The bar represents the expected red-shift gain, defined by the expected value of the increase from the base wavelength. 542





## 547 hydroxylamine bleach reaction.

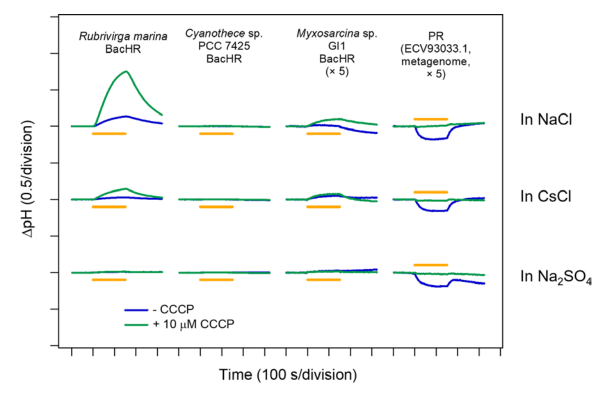
- 548 The difference absorption spectra between before and after hydroxylamine bleaching reaction
- of microbial rhodopsins in solubilized *E. coli* membrane. The  $\lambda_{max}$  of each rhodopsin was
- 550 determined by the peak positions of the absorption spectra of the original proteins, and the
- absorption of retinal oxime produced by the reaction of retinal Schiff base and hydroxylamine
- 552 was observed as a negative peak at around 360–370 nm.



#### 554

#### 555 Fig. 5. Observed wavelengths and expected red-shift gains.

The predicted and observed red-shift (and blue-shift) gains for the 40 candidate rhodopsins that showed significant coloring in *E. coli* cells. Differences between observed and base wavelengths are shown by the bars. The red bars indicate red-shift from the base wavelength, while the blue bars indicate observed wavelengths that were shorter than the base wavelengths. Proteins are sorted in the descending order by  $\mathbb{E}[\text{gain}]$ , as shown by the black line. Among the 40 candidates, 33 (82.5%) showed red-shift gains, suggesting that the proposed ML-based model can screen red-shifted rhodopsins more efficiently than random choice.

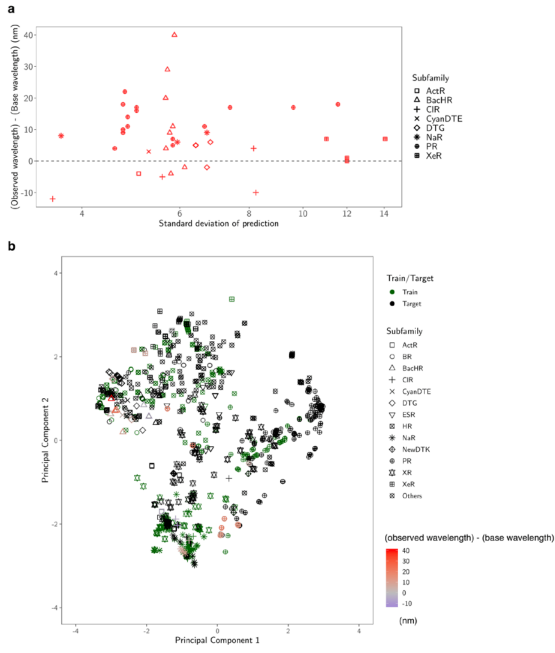


564

Fig. 6. Light-driven ion-transport activities of microbial rhodopsins showed longer  $\lambda_{max}$ . The light-induced pH change in the external solvent of *E. coli* cells expressing four microbial rhodopsins that showed a  $\lambda_{max} > 20$  nm longer than the base wavelength of the subfamily. The data obtained without and with 10  $\mu$ M CCCP are indicated by the blue and green lines, respectively, in 100 mM NaCl (top), CsCl (middle), and Na<sub>2</sub>SO<sub>4</sub> (bottom). Light was

illuminated for 150 s (yellow solid lines).

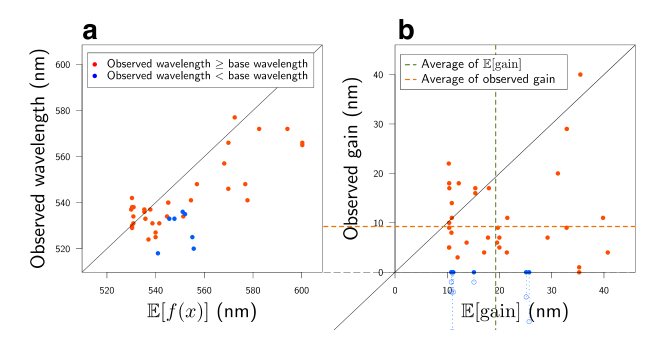
571



573 Fig. 7. Diversity of the selected proteins.

a Predicted standard deviation (horizontal axis) vs. observed gain (vertical axis). The marker shape represents the subfamily of each protein. **b** Two-dimensional projection created by principal component analysis. The original d = 432 dimensional feature space is projected onto the first two principal component directions. The first component (horizontal axis) explains 33% of the total variance of the original space, and the second (vertical axis) explains 17%. The green markers are the training data, and the black markers are the target data. For the synthesized proteins, differences in the observed and base wavelengths are shown by the color

- 581 map. The results indicate that, by considering the exploration–exploitation trade-off, it was
- 582 possible to make a red-shift protein screening process that considered not only the expected
- 583 value of the prediction, but also the uncertainty.

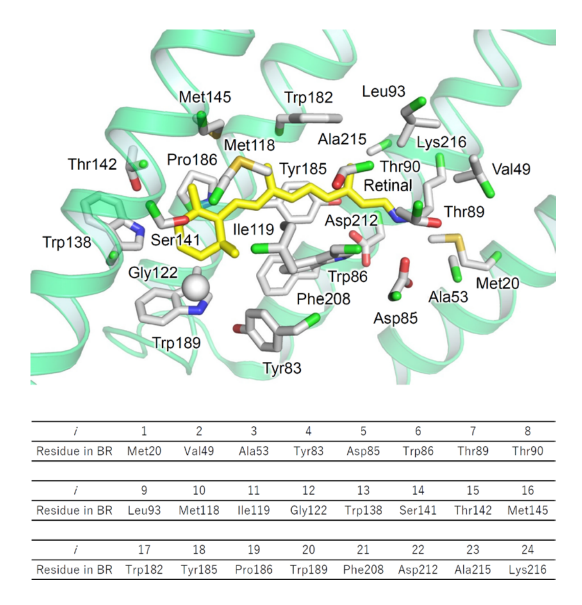




#### 586 Fig. 8. Comparisons of experimental observations and ML predictions.

In these two plots, the red points have longer observed wavelengths than the base wavelength 587  $\lambda_{\text{base}}$ , while the blue points have shorter observed wavelengths than  $\lambda_{\text{base}}$ . a ML-based 588 prediction of  $\lambda_{max}$  (horizontal axis) vs. experimentally observed  $\lambda_{max}$  (vertical axis). **b** 589 Expected red-shift gain (horizontal axis) vs. observed gain (vertical axis). Since we selected 590 591 rhodopsins having expected red-shift gains of > 10 nm, all the points on the horizontal axis are > 10 nm. The observed gain, defined by  $\max(\lambda_{\max} - \lambda_{\text{base}}, 0)$ , is nonnegative by definition. 592 The green and orange dashed lines are the averages of the horizontal and vertical axes (19.3 593 nm and 9.3 nm), respectively. The results indicate that the observed wavelengths and red-shift 594 595 gains tended to be smaller than the predicted ones. We conjecture that these differences between the observed and predicted wavelengths are due to modelling errors (see the Discussion for 596 details). 597

599	Supporting Information:
600	Exploration of natural red-shifted rhodopsins using a machine learning-
601	based Bayesian experimental design
602	Inoue and Karasuyama et al.
603	



604 605

# 606 Extended Data Figure 1. Amino acid residues around the retinal chromophore.

The structure of the 24 amino acid residues around the retinal used in the current ML model in the X-ray crystallographic structure of BR (PDB ID: 1IW6 (Matsui et al. *J. Mol. Biol.* (2002) **324**, pp. 469–481)). The C $\alpha$  atom of Gly122 is shown as a white sphere. For clarity, the ribbon models of helices B, C, and E were omitted. The table lists the residue numbers and names of each residue in BR.

Subfamily	Motif	Origin	₿ [gain]	(observed wavelength) – (base wavelength) / nm	Residue at BR Leu93	Residue at BR Pro186	Residue a BR A215
BacHR	TTD	Rubrivirga marina	35.5	40	L	P	Α
BacHR	TSV	Cyanothece sp. PCC 7425	32.9	29	1	P	А
PR	DTE	Metagenome sequence	10.3	22	М	Р	N
BacHR	TTV	Myxosarcina sp. Gl1	31.2	20	L	Р	А
PR	DTE	Pontimonas salivibrio	12.2	18	L	Р	N
PR	DTE	Fluviicola sp. XM24bin1	10.4	18	М	Р	N
PR	DTE	Bacteroidetes bacterium	15.4	17	м	Р	N
PR	DTE	Metagenome sequence	18.0	17	L	Р	N
PR	DTE	Metagenome sequence	10.4	17	М	Р	N
PR	DTE	Metagenome sequence	15.4	16	м	Р	N
PR	DTE	Metagenome sequence	10.9	14	м	Р	N
PR	DTE	Sphingobacteriales bacterium BACL12 MAG120802bin5	10.9	11	м	Р	N
PR	DTE	Nonlabens sp. YIK11	21.5	11	м	Р	N
BacHR	TSA	Rubrivirga marina	39.8	11	L	Р	А
PR	DTE	Metagenome sequence	10.4	10	М	Р	N
NaR	NDQ	Parvularcula oceani	19.7	9	L	т	S
BacHR	TSD	Cyanobacterium TDX16	32.9	9	L	Р	А
PR	DTE	Metagenome sequence	10.4	9	М	Р	N
NaR	NDQ	Spirosoma oryzae	10.8	8	L	Р	S
XeR	DSA	Nanohaloarchaea archaeon SW 7 43 1	29.2	7	1	Р	С
XeR	DSA	Metagenome sequence	17.8	7	1	P	с
PR	DTE	Metagenome sequence	20.0	7	М	Р	N
DTG	DTG	Rubrobacter aplysinae	19.5	6	L	Р	А
NaR	NDQ	Erythrobacter gangjinensis	13.7	6	L	P	S
DTG	DTG	Sphingomonas sp. Leaf34	10.3	5	L	Р	А
DTG	DTG	Sphingomonas sp. Leaf38	10.3	5	L	Р	А
PR	DTE	Metagenome sequence	20.0	5	М	Р	N
PR	DTE	Reinekea forsetii	17.1	4	L	Р	N
BacHR	TSA	Rubricoccus marinus	40.7	4	L	Р	А
CIR	NTQ	Metagenome sequence	21.4	4	L	Р	S
CyanDTE	DTD	Cyanobacteria bacterium QH 1 48 107	12.0	3	L	Р	А
XeR	DTA	Bacillus sp. CHD6a	35.3	1	L	P	S
XeR	DTA	Bacillus horikoshii	35.3	0	L	Р	s
DTG	DTG	Rosenbergiella nectarea	10.8	-2	L	P	А
BacHR	TSD	Hassallia byssoidea VB512170	15.1	-2	L	P	S
ActR	DTE	Kineococcus radiotolerans	11.2	-4	L	P	Α
BacHR	TSD	Aliterella atlantica	10.8	-4	L	P	S
CIR	NTQ	Metagenome sequence	25.1	-5	L	P	т
CIR	NTQ	Metagenome sequence	25.7	-10	L	P	т
CIR	NTQ	Sphingopyxis baekryungensis	11.0	-12	L	P	Ť

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# 614 Extended Data Figure 2. Amino acid residues at the color-tuning positions.

615 The amino acid residues at the color-tuning positions corresponding to BR Leu93, Pro189, and

616 Ala215.