pKD: re-designing protein $pK_a$ values

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ABSTRACT

The $pK_a$ values in proteins govern the pH-dependence of protein stability and enzymatic activity. A large number of mutagenesis experiments have been carried out in the last three decades to re-engineer the pH-activity and pH-stability profile of enzymes and proteins. We have developed the pKD webserver (http://polymerase.ucd.ie/pKa_Design), which predicts sets of point mutations that will change the $pK_a$ values of a set of target residues in a given direction, thus allowing for targeted re-design of the pH-dependent characteristics of proteins. The server provides the user with an interactive experience for re-designing $pK_a$ values by pre-calculating $\Delta pK_a$ values from all feasible point mutations. Design solutions are found in less than 10 min for a typical design job for a medium-sized protein. Mutant $\Delta pK_a$ values calculated by the pKD web server are in close agreement with those produced by comparing results from full-fledged $pK_a$ calculation methods.

INTRODUCTION

The pH-dependence of enzymatic activity and protein stability is of major importance for the biological function and industrial application of enzymes and proteins. pH-dependent protein characteristics have therefore been the subject of significant research efforts over the last two decades [e.g. (1–8)]. The pH-dependence of protein characteristics is determined by the $pK_a$ values of amino acid residues in the unfolded and folded state of a protein. Consequently a growing number of theoretical methods have been constructed for calculating the $pK_a$ values of protein titratable groups (9–20), and presently the best $pK_a$ calculation methods are accurate to within 0.5 $pK_a$ units when benchmarked against experimentally determined $pK_a$ values. The application of $pK_a$ calculation methods in biology is limited to a relatively small number of studies on catalytic mechanisms and ligand binding [e.g. (2–4,13,21–27)], and there is thus much scope for a wider application of $pK_a$ calculation methodology to biological problems. To our knowledge $pK_a$ calculations have yet to be used for designing proteins with novel pH-dependent properties, although methods have been developed for dissecting the contribution of charged groups to protein $pK_a$ values (25) (J. E. Nielsen, manuscript submitted). These methods can, in principle, be used for targeted re-design of protein $pK_a$ values.

To motivate research into designing proteins with novel pH-dependent properties we have developed the pKD web server (http://polymerase.ucd.ie/pKa_Design), which interfaces with a novel algorithm (B. M. Tynan-Connolly and J. E. Nielsen, manuscript submitted) for re-designing protein $pK_a$ values. The web server allows the user to specify a set of design criteria (target residues, desired $\Delta pK_a$ values, maximum number of mutations and minimum distance between target residue and any mutation) that specifies how the user want to change the $pK_a$ values of the protein. The pKD web server subsequently calculates a number of design solutions consisting of sets of point mutations predicted to change the target $pK_a$ values as specified by the design criteria. In addition to reporting the design solutions, the server produces 3D structural models of the proposed solutions and allows the user to inspect the $pK_a$ values calculated from the wild-type structure. The results and inputs of the pKD server can be analysed with a novel graphical interface ($pK_a$Tool) that facilitates the analysis of protein $pK_a$ calculations and titration curves. $pK_a$Tool is freely available to academic researchers at http://enzyme.ucd.ie/Science/pKa/pKaTool.

MATERIALS AND METHODS

The pKD server combines the functionality of a number of software packages using Python scripts. Figure 1 shows an overview of the functionality of the server and indicates where other software packages are employed. The parameters for the pKD $pK_a$ calculations are described in detail below. The pKD server software is freely available to academic researchers by contacting pka@ucd.ie, or in some cases by download from http://enzyme.ucd.ie/Science/pKa.

Preparation of PDB files

PDB files are prepared for a pKD design by deleting all water molecules and non-protein atoms. Missing protein atoms are rebuilt using the position-specific rotamer libraries (28) in WHAT IF (29).
Calculation of $pK_a$ values of the wild-type structure

The $pK_a$ values of the wild-type structure are calculated with the WHAT IF $pK_a$ calculation package as described elsewhere (19), except that a uniform dielectric constant of eight is used for the protein, and that the neutral–charged, charged–neutral and neutral–neutral interaction energies are calculated only for residue pairs with an interaction energy greater than 10 kT. The latter approximation is employed to speed up the $pK_a$ calculation step for large structures and can introduce errors in the reported $pK_a$ values for tightly coupled titratable groups. Users that find this to be a problem for a particular structure should request a full $pK_a$ calculation to be carried out by emailing pka@ucd.ie.

Modelling point mutations

Point mutations are modelled using position-specific rotamer libraries as implemented in WHAT IF (28). We mutate only residues that are at least 30% solvent exposed and fit well in the wild-type structure as deemed by automatic inspection of the rotamer library population. In addition we allow the mutation of buried charged residues to neutral residues of a similar size.

Finally we allow the user to exclude mutations at sites less than a certain distance (user adjustable but recommended to be at least 5 Å) from the residue whose $pK_a$ value is being redesigned.

Calculation of interaction energies for single point mutations

The interaction energies between a point mutation and all other residues are calculated using the WHAT IF $pK_a$ calculation package as described above.

Calculation of $\Delta pK_a$ values resulting from point mutations

$\Delta pK_a$ values are calculated using a Monte Carlo sampling method (30) implemented as a C++ class, which is imported into a python script. $\Delta pK_a$ values arising from single mutations are calculated by modifying the site–site interaction energy matrix using energies derived from an explicit model of the mutant protein structure. $\Delta pK_a$ values arising from multiple point mutations are calculated by modifying the site–site interaction energy matrix using energies calculated from single point mutation models. Thus no explicit 3D modelling of multiple point mutations takes place until the final solutions are found. Nevertheless we have shown (B. M. Tynan-Connolly and J. E. Nielsen, manuscript submitted) that $\Delta pK_a$ values calculated in this way are in excellent agreement with the $\Delta pK_a$ values found by comparing the results of full-fledged $pK_a$ calculations on wild-type and mutant protein structures, provided that only a small fraction of protein residues are mutated.

Finding the optimal set of point mutations

The search for sets of point mutations that fulfil the design criteria is initiated by selecting 20 sets of combinations of single point mutations, whose cumulative $\Delta pK_a$ values are closest to the design criteria. Since $\Delta pK_a$ values only sometimes can be combined linearly (B. M. Tynan-Connolly and J. E. Nielsen, manuscript submitted), we calculate a more realistic set of $\Delta pK_a$ values for each solution as described above. Finally we perform a short Monte Carlo sampling to find solutions that do cannot be found from a linear combination of individual $\Delta pK_a$ values.

Scoring of design solutions

We apply a scoring function of the form

$$\text{score} = \sum (pK_a_{\text{desired}} - pK_a_{\text{achieved}})^2$$

1
to identify the design solutions in agreement with the design criteria. The sum in equation 1 is calculated over all $pK_a$ criteria specified in the design setup phase. The function is optimized to identify a specific $pK_a$ change rather than identifying the largest possible $pK_a$ change. Therefore, if one is

![Figure 1. The workflow of the pKD server. The server incorporates the functionality of the WHAT IF $pK_a$ calculation package (WIpKa), construction of point mutations (WHAT IF) and the pK$_a$ Design algorithm.](image)
Table 1. Example design solutions and predicted \( \Delta pK_a \) values from pKD, the WHAT IF \( pK_a \) calculation package (19) and the H++ webserver (12) for a re-design of the \( pK_a \) of Glu 35 in HEWL (PDBID 2lzt)

| Design criteria | Minimum distance to target | Mutations | pKD \( \Delta pK_a \) | WI \( \Delta pK_a \) | H++ \( \Delta pK_a \) |
|-----------------|-----------------------------|-----------|---------------------|-----------------|-----------------|
| Glu35 +2.0      | 5.0 \( \text{Å} \)          | R112E+R114Q+N37E+W62D | 1.2               | 0.9             | 1.0             |
| Glu35 −2.0      | 10.0 \( \text{Å} \)         | N103H+D48R+T43K       | −0.6              | −0.6            | −0.7            |

The maximum number of mutations allowed was set to 5. WI \( pK_a \) calculations were carried out as described in Materials and Methods, and H++ calculations were carried out with the default settings on the website. The results illustrate that it often is not possible to achieve the desired \( pK_a \) change with the parameters and the structure in question.

RESULTS AND CONCLUSIONS

We have constructed a the pKD web server, which allows for the re-design of protein \( pK_a \) values by site-directed mutagenesis. Proteins with redesigned \( pK_a \) values will display a change in their pH-dependent characteristics, such as ligand binding, stability and, for enzymes, catalytic rate, and the pKD server is thus aimed at researchers who aim to understand or change the pH-dependent properties of proteins.

The pKD webserver asks the user to select one or more titratable groups for \( pK_a \) value re-design. It must specify how many mutations one is willing to construct and how close these mutations can be to the titratable groups of interest. Subsequently the pKD server predicts a set of point mutations that will change the \( pK_a \) values of the selected titratable groups in the given direction, by the desired amount. Furthermore the user is presented with a set of modelled 3D structures containing the proposed mutations, which can be analysed using pKD Tool (J. E. Nielsen, manuscript submitted).

It is thus possible for the user to achieve an interactive, in-depth understanding of the titrational behaviour of any given protein using these two freely available software tools. Future work will focus on integrating pKD Tool and the pKD server directly to allow for a desktop-based, convenient, interactive analysis facility of the titrational behaviour of wild-type and mutant proteins.

Table 1 and Figure 2 show a typical design solution from the pKD server, and illustrates that good agreement is obtained between the \( \Delta pK_a \) values reported by the pKD server and those predicted by a full-fledged pKD calculation packages. The pKD server uses a full physical model of the pH-dependent behaviour of titratable groups, which ensures that effects arising from differences in intrinsic \( pK_a \) values and complicated pair-wise electrostatic interaction networks are calculated as accurately as possible. Therefore, the solutions calculated with the pKD server are more physically realistic than solutions obtained using the classic relation \( \Delta pK_a = (\Delta \Phi / \ln 10) \) (8), which is known to break down when multiple strong electrostatic interactions are present and when dealing with titratable groups with perturbed intrinsic \( pK_a \) values.

We have calculated design solutions for a large number of protein structures and examined the dependence of the results on the algorithm parameters, and we have found the algorithm to reliably identify mutations that will change the \( pK_a \) values of target residues as judged by theoretical methods. We are currently verifying these \( pK_a \) shifts in the lab, and simultaneously continuing our theoretical studies on factors that influence the pH-dependent properties of proteins.

We believe that the pKD server will be of great benefit to researchers that are interested in re-designing and understanding the pH-dependent characteristics of proteins, and we furthermore hope that the server will encourage more experiments aimed at understanding the complex links between protein structures and their pH-dependent characteristics.

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