Fluorescently Labelled ATP Analogues for Direct Monitoring of Ubiquitin Activation

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Abstract: Simple and robust assays to monitor enzymatic ATP cleavage with high efficiency in real-time are scarce. To address this shortcoming, we developed fluorescently labelled adenosine tri-, tetra- and pentaphosphate analogues of ATP. The novel ATP analogues bear — in contrast to earlier reports — only a single acridone-based dye at the terminal phosphate group. The dye’s fluorescence is quenched by the adenine component of the ATP analogue and is restored upon cleavage of the phosphate chain and dissociation of the dye from the adenosine moiety. Thereby the activity of ATP-cleaving enzymes can be followed in real-time. We demonstrate this proficiency for ubiquitin activation by the ubiquitin-activating enzymes UBA1 and UBA6 which represents the first step in an enzymatic cascade leading to the covalent attachment of ubiquitin to substrate proteins, a process that is highly conserved from yeast to humans. We found that the efficiency to serve as cofactor for UBA1/UBA6 very much depends on the length of the phosphate chain of the ATP analogue: triphosphates are used poorly while penta-phosphates are most efficiently processed. Notably, the novel pentaphosphate-harbouring ATP analogue supersedes the efficiency of recently reported dual-dye labelled analogues and thus, is a promising candidate for broad applications.

Introduction

Ubiquitination — the covalent modification of proteins by the 76 amino acid protein ubiquitin (Ub) — is a ubiquitous protein modification with fundamental roles in numerous cellular processes including protein degradation, DNA damage repair, cell cycle regulation and gene expression.[1,2] Malfunction of the ubiquitination system contributes to a broad variety of human diseases like cancer, diabetes or neurodegenerative disorders.[3,4] For the attachment of Ub to substrate proteins, the consecutive action of at least three classes of enzymes is needed. In the first step, Ub is activated by a ubiquitin-activating enzyme (E1) at the consumption of ATP. Thereby Ub is adenylated and then transferred to the active-site cysteine of the E1 to form a thioester with the C-terminal glycine carboxylate of Ub[5] (Figure 1A). By transthiolation, Ub is transferred to a ubiquitin-conjugating enzyme (E2). Finally, by the aid of ubiquitin-protein ligases (E3), Ub is covalently connected to the target protein by forming an isopeptide bond with the ε-amino group of a lysine residue.

In humans, two E1s (UBA1 and UBA6) for Ub are known.[6,7] Quantifying and following E1 activity in real-time is of great importance to study for example, effectors, but means to do

Figure 1. A) Ubiquitin (Ub) activation by E1 (UBA1) with ATP. Adenylated Ub is loaded on UBA1 by the formation of a thioester. B) Concept of mono-labeled ATP analogues as UBA1 sensors that are investigated in this study: In close proximity to the nucleobase adenine, the fluorescent dye is quenched as a result of photoinduced electron transfer (PET). After enzymatic release of the phosphate chain, the fluorescence is restored.
so are sparse. Available E1 activity assays include SDS-PAGE analysis of E1/E2-Ub thioester conjugates by Western Blot,\textsuperscript{[13,14]} radio-labelling of the involved proteins with \textsuperscript{[15]} \textsuperscript{[16]} or \textsuperscript{[17]} FRET between Ub and E\textsuperscript{1}\textsuperscript{[18,19]} or enzyme-coupled spectrophotometric assays for phosphate determination.\textsuperscript{[12]} Drawbacks of these assays are that they are either laborious, do not allow continuous read-out or are dependent on additional enzymes of the downstream cascade. To fill this gap, we have recently developed a time-resolved ATPase sensor (TRASE) based on a Förster resonance energy transfer (FRET) pair embedded within the ATP scaffold to continuously monitor ATP-dependent enzymes.\textsuperscript{[13,14]} This ATP FRET probe sensor uses two fluorescent dyes, a donor and an acceptor dye, that are attached to the terminal phosphate group and to adenine, respectively. By employing the TRASE assay, we found that γ-modified triphosphates are poorly accepted by UBA1 whereas δ-modified tetraphosphates turned out to be better substrates.\textsuperscript{[15]}

In order to improve the substrate properties of ATP-based E1 sensors, we synthesized and investigated new ATP analogues that contain only one dye and differ in the length of the phosphate chain for their propensity to visualize E1 activity. Indeed, we identified a fluorescent dye with an acridone core structure that is efficiently quenched by the canonical nucleobase adenine.

While triphosphates were poorly processed, we found that also in this case acceptance is improved by elongation of the phosphate chain, that is, tetraphosphates are better accepted than triphosphates. In fact, elongation of the phosphate chain to pentaphosphate resulted in even better analogues enabling us to follow the activation of Ub by both enzymes, UBA1 and UBA6, in real-time.

Results and Discussion

Recently, we developed γ-modified ATP analogues that are suitable as model compounds for monitoring enzymatic ATP consumption by fluorescence lifetime readout. Compared to other probes, these compounds contain only one instead of two fluorophores, which promises better enzymatic acceptance. Fluorescence lifetime changes between the intact and the cleaved ATP analogues are caused by the quenching of fluorescence by the nucleobase adenine\textsuperscript{[16]} (Figure 1B). The quenching is caused by photoinduced electron transfer (PET). In PET, the efficiency for electron transfer rates and thus for quenching can be estimated by using the Rehm-Weller equation and/or by comparing the involved highest occupied molecular orbital (HOMO) energy levels.\textsuperscript{[17–19]} The PET process starts with excitation of the acceptor chromophore by irradiation (step 1 in Figure 2). This promotes an electron into the lowest unoccupied molecular orbital (LUMO) of the acceptor.

In case the HOMO of a neighbouring donor molecule in the immediate vicinity is higher in energy than the HOMO of the acceptor molecule, an electron is transferred from donor to acceptor (step 2) and fluorescence is quenched. The cycle is closed by transfer of an electron from the acceptor LUMO to the donor HOMO. This process can only take place for distances between acceptor and donor on a nanometre scale, that is, in the intact ATP analogue. Enzymatic cleavage of the phosphate chain, however, leads to an immediate separation of the dye-adenine pair by diffusion such that fluorescence of the dye is restored.

PET-based quenchers that have previously been employed in biological assays are guanine (−5.33 eV; calculated from its redox potential)\textsuperscript{[20]} and tryptophan (−4.90 eV; calculated from its redox potential)\textsuperscript{[21]} both of which possess high HOMO energies.\textsuperscript{[22–26]} By contrast, due to its lower HOMO energy (−5.78 ± 0.01 eV)\textsuperscript{[16]} adenine has only rarely been used as quencher. To develop new suitable dye-adenine pairs, we used photoelectron spectroscopy in air (PESA) to determine HOMO energy levels for hydrophilic dyes. With this approach, we recently were able to investigate different fluorophores like rhodamines and BODIPYs from dry thin films that suited for fluorescence lifetime readout when attached to ATP\textsuperscript{[15]} While we could demonstrate enzymatic processing and real-time detection of these compounds, their main drawback was the relatively short lifetime of these dyes. This made the detection of lifetime changes especially under biologically relevant conditions difficult.

In order to overcome this shortcoming, we now investigated fluorescent dyes with long fluorescence lifetimes. Acidone and quinacridone as well as their derivatives are interesting candidates for our purpose, especially because they are extremely photostable\textsuperscript{[22,23]} have a long fluorescence lifetime of 14 ns and 22 ns, respectively, and show no spectral pH dependency in the biological relevant range from pH 5–9.\textsuperscript{[24]} However, only few examples are reported where in particular acidone derivatives under the name Puretime 14 (PT14) were used for fluorescence lifetime-based biological assays.\textsuperscript{[29–32]} For functionalization and attachment of acidone or quinacridone to ATP, we followed a known synthesis strategy. Initially, we created water-soluble compounds from the organic pigments by introducing sulfonic acid residues (Scheme 1).\textsuperscript{[28]} We expected that sulfonation would also lower the energy of the frontier orbitals that in turn would result in efficient quenching of the dye when being in close proximity to adenine.

Both N-alkylated fluorescent dyes were treated with hot sulphuric acid, which yielded Sacridone-COOH (2.1) and S\textsubscript{4}quinacridone-COOH (4), respectively. For disulfonation of the functionalized acidone, we used 65% SO\textsubscript{3} in H\textsubscript{2}SO\textsubscript{4} to yield S\textsubscript{4}acidone-COOH (2.2).
We speculated that extended phosphate groups are introduced. Since the measured HOMO energies are decreasing, the more electron-withdrawing phosphates are even better substrates for nucleic acid polymerases than shorter congeners.

We found that all functionalized acridones (1, 2.1 and 2.2) possess lower HOMO energies than ATP, whereas the sulfonated \( S_2 \text{quinacridone} \) is higher in energy. For the acridone derivatives (1–2.2), we measured HOMO energies of \(-5.91 \pm 0.05 \text{ eV} \) (1), \(-5.93 \pm 0.03 \text{ eV} \) (2.1) and \(-6.03 \pm 0.02 \text{ eV} \) (2.2). These findings nicely confirm our expectation that the HOMO energies are decreasing, the more electron-withdrawing groups are introduced. Since the measured HOMO energies are lower than that of ATP, we moved on to the synthesis of a \( \gamma \)-modified ATP analogue bearing disulfonated acridone. Here, we followed a well-established synthesis strategy. In brief, commercially available ATP was alkylated, followed by azide reduction with tris(2-carboxyethyl)phosphine (TECP) and subsequent NHS ester coupling by activating the carboxylic acid of \( S_2 \text{acridone-COOH} \) with \( N,N,N’,N’ \)-tetramethyl-O-(N-succinimidyl)uronium tetra-fluoroborate (TSTU) in situ which was finally coupled without purification to the free amine to yield \( \text{Ap}_2 \)-Dye (5) (Scheme 2).

As a first proof of concept, we determined the fluorescence lifetimes of \( \text{Ap}_2 \)-Dye (5) in the intact and cleaved state. As a model enzyme for cleavage, we used the ATP hydrolysing phosphodiesterase I from \( \text{Crotalus adamanteus} \) (Snake Venom Phosphodiesterase, SVPD). Before cleavage, we measured a fluorescence lifetime value of 8.71 ± 0.10 ns. By addition of SVPD, the triphosphate is cleaved and the quencher adenine is released as AMP and separated from the fluorophore. Here, we measured a value of 15.64 ± 0.25 ns, corresponding to an absolute lifetime change of 6.93 ± 0.27 ns. With typical experimental errors in fluorescence lifetime determination in the range of 100 ps, this difference is well suited to use lifetimes for quantification of ATP analogue cleavage. Additional absorption and emission spectra recorded before and after cleavage with SVPD did not reveal a spectral shift assuming a low interaction between the sulfonated dye and the nucleobase adenine (data are shown in the Supporting Information).

Encouraged by these results, we next focused on the synthesis of ATP analogues with longer phosphate chains to determine their acceptance by UBA1. From our previous experiments on the enzymatic turnover of ATP analogues by UBA1, we knew that modified \( \delta \)-tetraphosphates are superior substrates compared to modified \( \gamma \)-triphosphates in UBA1-mediated activation of Ub. We speculated that extended phosphate anhydride chains in the present ATP analogue will also increase the activity towards UBA1. Therefore, we synthesized a modified \( \delta \)-tetraphosphate by activating ATP with 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (EDC) to form a triphosphate which was treated with 1-(6-azido)hexyl phosphate according to the literature. Afterwards the product was reduced and coupled to \( S_2 \text{acridone-NHS} \) (Ap\(_2\)-Dye (6), Scheme 2). Interestingly, for \( \text{Ap}_2 \)-Dye (6) we found a fluorescence lifetime of 8.74 ± 0.03 ns, which is similar to that of \( \text{Ap}_2 \)-Dye (5) (8.71 ± 0.10 ns).

Findings from other groups showed that modified pentaphosphates are even better substrates for nucleic acid polymerases than shorter congeners which motivated us to additionally synthesize an \( \varepsilon \)-modified alkylated pentaphosphate. Described synthesis routes of alkylated pentaphosphates that follow P\(^\varepsilon\)N activation and subsequent substitution often lead to unwanted side-products. Therefore, we decided to explore a known iterative polyphosphorylation approach strategy. The reaction is conducted in several steps without purification in one pot. Only modified monophos-

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**Scheme 1.** Synthesis of disulfonated acridone (upper panel) and quinacridone (lower panel) derivatives: a) conc. \( \text{H}_2\text{SO}_4 \), 120 °C, 20 h. b) 65% \( \text{SO}_3 \) in conc. \( \text{H}_2\text{SO}_4 \), rt, 24 h. c) conc. \( \text{H}_2\text{SO}_4 \), 110 °C, 20 h.

**Scheme 2.** Synthesis of \( S_2 \text{acridone} \) functionalized tri- and tetraphosphates.

**Figure 3.** Relative HOMO energies in air. The dashed line represents the value for ATP.
phosphates are needed as starting materials, which can be easily obtained in high yields or are even commercially available. Scheme 3 shows an overview of the synthesis of Ap₅⁻Dye. Disisopropylamino dichlorophosphine (7) is reacted first with pyrophosphate to form a cyclic pyrophosphoryl P-amidite (8) which is coupled to 6-azidohexyl phosphate by subsequent oxidation to form 1-(6-azido)hexyl phosphorocyclotriphosphate (9). The cyclic trimetaphosphate is then opened by adding adenosine monophosphate as nucleophile and MgCl₂ to yield 6-O-6-azidohexyl)-adenosine-OS⁻pentaphosphate (10) in a one-pot reaction with an overall yield of 11%. Reduction of the azide followed by coupling of the activated S-acridone using its NHS ester yielded the desired compound (Ap₅⁻Dye (11), 28% over two steps, Scheme 3). For the Ap₅⁻Dye 11, the fluorescence lifetime of 9.86 ± 0.19 ns is slightly higher than that observed for the tri- and tetraphosphates 5 and 6 (8.71 ± 0.10 ns and 8.74 ± 0.03 ns, respectively). The same fluorescence lifetime of the free dye was measured when 10 was treated with SVPD as for (5 ± 6) (15.64 ± 0.25 ns). The reduced quenching for longer distances between dye and quencher pairs is in accordance to the literature where efficient PET-quenching takes place on a sub-nanometre scale.[42]

With all three compounds in hand, we tested them towards their performance in an E6AP auto-ubiquitination assay that was previously shown to be well-suited to qualitatively evaluate the acceptance of ATP analogues by UBA1 (Figure 4).[15,43] All reaction mixtures were pre-treated with recombinant shrimp alkaline phosphatase (rSAP) which dephosphorylates all terminally bound phosphate groups, for example, of ATP to its nucleoside,[44] while it leaves terminally modified nucleotides unaffected (like for Ap₅⁻Dye). This ensures that the observed activity is due to the Ap₅⁻Dye analogue and does not originate from potential contaminations of natural ATP. After preincubation, rSAP was inactivated by heating the mixture to 65°C for 5 minutes. In the first lane in Figure 4A and B, reactions are de-
Finally, we measured the activation of Ub by UBA1 and UBA6 under conditions, where both thioesters (i.e. UBA1 → Ub and UBA6 → Ub) are unloaded. To do so, we added UbcH5b and E6AP in the same concentration as for the SDS-PAGE experiment (Figure 4). As shown in Figure 6, again neither UBA1 nor UBA6 are able to activate Ub with Ap4-Dye (5) as cofactor, while elongation of the phosphate chain rescues activity as already seen in the SDS-PAGE analysis. Ap2-Dye (6) and Ap4-Dye (11) are both linearly processed which shows once more the suitability of our setup. Moreover, the extension of the tetraphosphate chain to Ap2-Dye (11) increases acceptance of both UBA1 and UBA6 significantly. When all cognate enzymes are present, UBA1 activates Ub approximately 2.9 times faster with Ap2-Dye (11) as ATP source compared to Ap4-Dye (6). This tendency is also seen with UBA6 where the activation with Ap2-Dye (11) is even 4.9 times faster than with Ap4-Dye (6).

Conclusions

In conclusion, we developed and explored novel fluorescently labelled adenosine tri-, tetra-, and pentaphosphates. The ATP analogues bear a single acridone-based dye at the terminus of the phosphate chain. Most importantly, the dye’s fluorescence is quenched by the adenine residue of the ATP analogue. Fluorescence is restored upon cleavage of the phosphate chain and dissociation of the dye from the adenosine moiety. Therefore, the activity of ATP cleaving enzymes can be followed.

In comparison to our earlier approaches, in this approach only one dye modification is appended to the ATP analogue. This has several advantages. Obviously, the synthesis towards the probes is simplified making these sensors more readily available. Another advantage is the absence of any modification at the nucleobase thereby rendering the analogues to be superiorly processed by the enzymes investigated here. This might be due to the fact that upon usage by the adenylate-forming enzymes investigated here, the formed reactive adenylated ubiquitin species is identical to that with natural ATP.

By elongation of the phosphate chain, we were able to increase acceptance and reaction velocity significantly and demonstrated this for UBA1 and UBA6 that accept Ap2-Dye (11)

(Cy5-Ap4-Cy3) (for structure see Supporting Information). Interestingly, at high Ub concentration we observed for all three compounds a reaction inhibition (see Supporting Information Figure S2.3-S2.5), while at low substrate concentration, Michaelis–Menten kinetics were observed. A possible explanation for this observation is that the UBA1-SH binding site is non-covalently occupied by a second Ub upon increasing Ub concentration, thus, inhibiting the transfer of adenylated Ub to the active-site cysteine of UBA1 by forming a thioester with the C-terminal glycine carboxylate of Ub. Therefore, we fitted our data (Figure 5) to a kinetic model describing such a mode of inhibition[46] [Eq. 1]:

\[
V = \frac{V_{\text{max}}}{1 + \frac{K_m}{[S]}}
\]

The high affinity binding site is described by \(K_{hi}\), whereas the inhibitory site, which is in general markedly lower in affinity, is described by \(K\). We found for all analogues a \(K\) of approximately 50 µM and a \(K_{hi}\) of (4.1 ± 0.5) µM for Ap2-Dye (11) which is more than 7 times lower than the \(K_{hi}\) for Ap4-Dye (6) (29.1 ± 3.0) µM and almost 20 times lower than the doubly labelled Cy5-Ap4-Cy3 (76.1 ± 6.7) µM. However, the \(K_m\) for Ub with natural ATP is even lower (0.2 µM) suggesting that our analogues may somewhat interfere with Ub binding.[47]

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best. We could also show that the herein presented analogues can be readily used in real-time assays to follow Ub activation by UBA1 and UBA6. Notably, using the developed disulfonated acridone, the read-out can be both fluorescence intensity and lifetime. These characteristics make the herein developed ATP analogues versatilely applicable for future uses, for example, in the high-throughput screening for effectors of E1 enzymes.

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Conflict of interest

The authors declare no conflict of interest.

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