

Structural basis for the fast maturation of Arthropoda green fluorescent protein

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Since the cloning of *Aequorea victoria* green fluorescent protein (GFP) in 1992, a family of known GFP-like proteins has been growing rapidly. Today, it includes more than a hundred proteins with different spectral characteristics cloned from *Cnidaria* species. For some of these proteins, crystal structures have been solved, showing diversity in chromophore modifications and conformational states. However, we are still far from a complete understanding of the origin, functions and evolution of the GFP family. Novel proteins of the family were recently cloned from evolutionarily distant marine Copepoda species, phylum Arthropoda, demonstrating an extremely rapid generation of fluorescent signal. Here, we have generated a non-aggregating mutant of Copepoda fluorescent protein and solved its high-resolution crystal structure. It was found that the protein β -barrel contains a pore, leading to the chromophore. Using site-directed mutagenesis, we showed that this feature is critical for the fast maturation of the chromophore.

Keywords: GFP; Arthropoda; TurboGFP; structure; chromophore maturation

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INTRODUCTION

Protein-related fluorescence in hydromedusa *Aequorea victoria* was first studied more than 40 years ago (Johnson *et al*, 1962). Cloning (Prasher *et al*, 1992) and characterization of *A. victoria* green fluorescent protein (GFP) provided the scientific community with a protein capable of visible-spectrum fluorescence without the requirement of exogenous cofactors. Subsequently, GFP and its derivatives have been extensively used as non-invasive, *in situ*

and *in vivo* markers (Lippincott-Schwartz & Patterson, 2003; Chudakov *et al*, 2005). In 1996, the crystal structure of GFP was solved (Ormo *et al*, 1996; Yang *et al*, 1996), which spurred the development of scientifically useful fluorescent variants.

Divergent evolution doctrine suggests that other fluorescent organisms are likely to contain GFP homologues. Indeed, more than 120 fluorescent and coloured GFP-like proteins were cloned from nearly 70 *Cnidaria* species (Labas *et al*, 2002; Matz *et al*, 2006). The similarity of these proteins to GFP ranges from 80–90% to less than 25% identity in amino-acid sequence, but all these proteins share the same β -barrel fold. One would expect GFP-like proteins to be found in organisms that are evolutionarily close to *Cnidaria*. However, surprisingly, we have recently reported several fluorescent proteins from evolutionarily distant Copepoda species (Arthropoda: Crustacea: Maxillopoda: Copepoda: Pontellidae; Shagin *et al*, 2004). The biological role for Pontellidae fluorescence is unclear. These organisms are not luminescent, which eliminates the need for the fluorescent protein as a chromatic shift agent. Presumably, visual mate recognition is important, as Pontellidae have elaborate eyes and show sexual dimorphism in eye design (Ohtsuka & Huys, 2001).

Some of the Copepoda GFPs, such as pfluGFP2 from *Pontellina plumata* (GenBank accession number AY268072), showed extremely rapid development of bright green fluorescent signal when expressed in *Escherichia coli* and mammalian cells (HeLa, mouse embryonic fibroblasts 3T3 and human embryonic kidney 293T). Solving the crystal structure of pfluGFP2 would increase understanding of this phenomenon. As Copepoda GFPs are prone to form aggregates, protein purification and crystallization experiments have been hindered. Overexpression of these proteins in mammalian cells also leads to formation of micro-crystals that eventually rupture membranes, which limits their applicability as biological tags (Fig 1A).

To overcome these problems, we performed solubility-enhancing optimization of pfluGFP2. We produced a highly soluble, rapidly maturing variant, named TurboGFP. Having solved its crystal structure, we were able to analyse the chromophore environment and explain the rapid development of the fluorescent signal *in vivo*.

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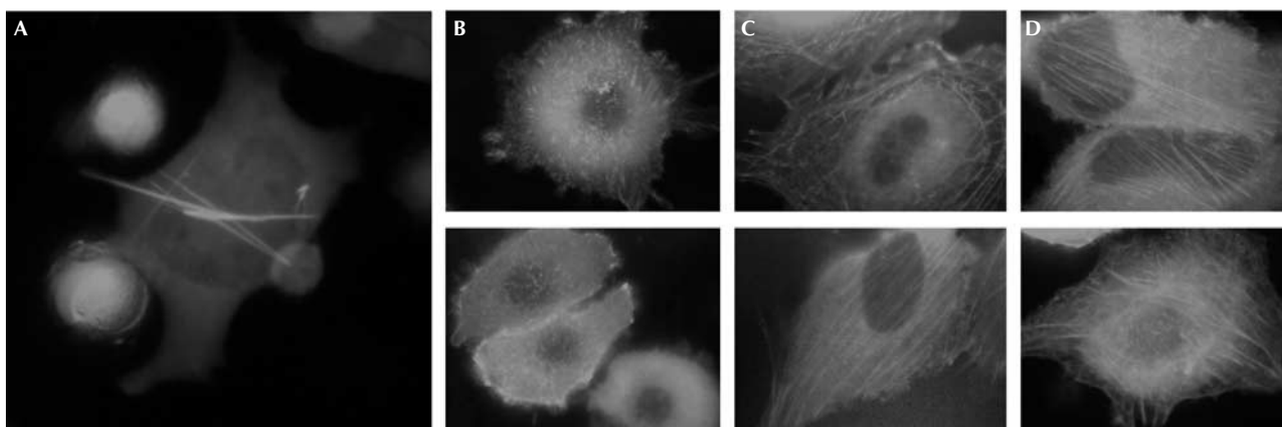


Fig 1 | Confocal microscopy of fluorescent protein expression in HeLa cells. (A) ppluGFP2 (green fluorescent protein from *Pontellina plumata*) forms needles 3 days after transfection. (B) ppluGFP2- β -actin fusion protein. (C) TurboGFP- β -actin fusion protein. (D) Enhanced green fluorescent protein- β -actin fusion protein.

RESULTS AND DISCUSSION

Development of non-aggregating ppluGFP2

We have threaded the sequence of ppluGFP2 onto the *A. victoria* GFP structure (Protein Data Bank (PDB) ID: 1GFL) using Swiss-PdbViewer and HyperChem 5.01 (Hypercube Inc., Gainesville, FL, USA). The resulting model was used to generate a plausible ppluGFP2 molecular surface. Analysis of the computed electrostatic potential suggested that the protein surface is mostly negatively charged with the exception of a positively charged patch of residues near the amino and carboxyl termini. We supposed that aggregation might be caused by electrostatic interactions between the positively and the negatively charged surfaces (Himanen *et al*, 1997). With the aim of shielding the positively charged patch, we added polar, negatively charged amino acids to the protein termini. The resulting ppluGFP2 variant aggregated less *in vitro*. However, it still formed microcrystals 7 days after transient transfection of HeLa cells (see supplementary information 1 online for details). Importantly, it was shown to be tetrameric, according to the gel-filtration data. This suggests that wild-type ppluGFP2 is also tetrameric (ppluGFP2 was originally characterized tentatively as a monomer; Shagin *et al*, 2004).

We followed up with another cycle of mutagenesis, changing exposed hydrophobic residues to polar and charged ones. The initial ppluGFP2 model suggested oligomer interfaces, by comparison with structures of oligomeric fluorescent proteins. To loosen the ppluGFP2 tetramer and make the protein more suitable as a fusion tag, we proceeded to make non-conservative substitutions of several key residues in the hypothetical interfaces. Ultimately, we obtained a non-aggregating dimeric ppluGFP2 variant, designated TurboGFP (see supplementary Fig 1 online for sequence).

Characterization of TurboGFP

TurboGFP is a bright green fluorescent protein (see supplementary Fig 2 online for fluorescence spectrum) with a quantum yield of 0.53 and molar extinction coefficient of $70,000 \text{ M}^{-1} \text{ cm}^{-1}$. Its fluorescence has rather high pH stability (pK_a 5.2). The protein is a dimer in solution, at least at concentrations up to 5 mg/ml

(0.2 mM), according to gel-filtration (supplementary Fig 3 online) and light scattering (unimodal distribution of apparent molecular weight centred at 54 kDa) data. It is soluble under physiological conditions (pH 7.4, 20 mM phosphate-buffered 0.14 M NaCl), at least at concentrations up to 60 mg/ml.

We have succeeded in expressing high levels of TurboGFP in bacteria, yeast (see supplementary information 2 online) and mammalian cells. Unlike with the native ppluGFP2, stable cell lines expressing TurboGFP can be established, as was shown for HeLa, 3T3, mouse melanoma M3 and PC12 cell lines (Dr C. Petzelt, personal communication).

We have also generated a destabilized TurboGFP version with a rapid turnover, by fusing the protein with the degradation domain of mouse ornithine decarboxylase (Li *et al*, 1998). For evaluation of the construct, mammalian cells, transiently transfected with destabilized TurboGFP, were treated with 100 $\mu\text{g/ml}$ cycloheximide (an inhibitor of protein synthesis). According to time-lapse measurements, the destabilized TurboGFP had a half-life of 2 h (supplementary Fig 4 online). Remarkably, the wild-type ppluGFP2 was insensitive to fusion of the degradation domain, presumably owing to aggregates being inaccessible to the proteolytic machinery. Rapidly maturing fluorescent proteins with a characteristically short half-life are particularly useful for studies of gene expression timelines.

To evaluate the suitability of TurboGFP as a fusion partner for *in vivo* fluorescent tagging, we compared the performance of ppluGFP2, TurboGFP and the commonly used enhanced green fluorescent protein (EGFP; Clontech, Mountain View, CA, USA) fusions with β -actin in HeLa cells. Fig 1C,D shows that the fluorescence patterns of mammalian cells expressing TurboGFP- β -actin are not visually distinguishable from those expressing EGFP- β -actin. Both show formation of typical elongated actin filaments and an apparently normal cytoskeleton. By contrast, the ppluGFP2- β -actin filaments are much shorter, leading to distortion of the cytoskeleton morphology (Fig 1B).

Similar to the wild-type ppluGFP2, TurboGFP is characterized with a rapid generation of fluorescent signal in living cells. In transient transfections of HeLa, 3T3 and Phoenix Eco cell lines,

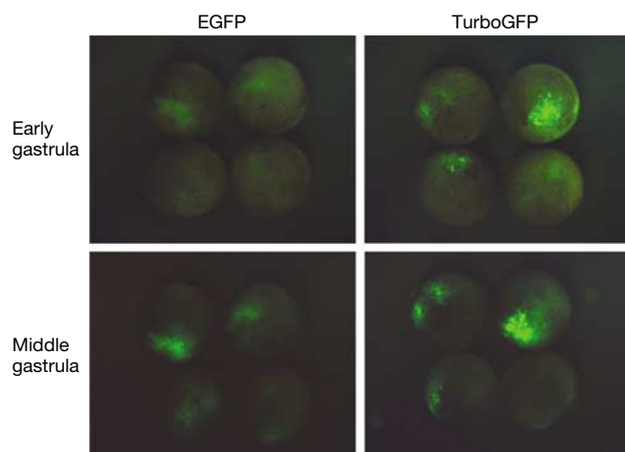


Fig 2 | Comparison of TurboGFP and enhanced green fluorescent protein maturation speed in developing *Xenopus laevis* embryos. At the stage of two blastomeres, embryos were microinjected with TurboGFP-C1 and pEGFP-C1 vectors. Living embryos were photographed from the animal pole side at the early and mid-gastrula stages. EGFP, enhanced green fluorescent protein.

TurboGFP fluorescence was detectable 3–5 h earlier than that of EGFP. Furthermore, we compared TurboGFP and EGFP in developing *Xenopus laevis* embryos (Fig 2). TurboGFP fluorescence was about threefold brighter at the early (10 h after fertilization) and twofold brighter at the middle (12 h after fertilization) gastrula stages, demonstrating the speed advantage offered by this tool to visualize gene expression.

TurboGFP crystal structures

TurboGFP expression in *E. coli* easily yielded enough material to pursue structural studies. The details of cloning, protein production, crystallization and structure solution are provided in Methods and supplementary information 3 online. As expected, despite the modest degree of identity between TurboGFP and other members of the GFP family, the structure of this protein is a β -barrel fold (PDB IDs 2G6X and 2G6Y; Fig 3A). TurboGFP is a dimer in solution at concentrations at least up to 5 mg/ml, whereas it forms tetramers in the crystal form. It is likely that TurboGFP dimers undergo reversible association into tetramers at higher concentrations. There are virtually no differences in the structures of the two crystal forms ($P2_1$ and $C222_1$; see supplementary Table 1 online for details)—they superimpose with inter-monomer and inter-tetramer $C\alpha$ root-mean-square deviations (r.m.s.d.) of 0.35 and 0.5 Å, respectively. All four monomers in the TurboGFP asymmetric unit are also virtually identical (pairwise $C\alpha$ r.m.s.d. of 0.3 Å), with the exception of residues 185–187 of chain B in the $P2_1$ crystal form, which adopt a slightly different conformation, probably because of a crystal contact. The quaternary structure of TurboGFP tetramer is similar to that of DsRed (Wall *et al*, 2000; Yarbrough *et al*, 2001). In fact, the two tetramers superimpose with a least-squares minimized $C\alpha$ r.m.s.d. of 1.45 Å.

The first dimer interface (monomers A|B and C|D) occludes about 1,093 Å² of molecular surface per monomer. The interface is composed of residues A137, V139, H141, H143, M145, Y165,

P190, F192, F194, F215 and P218 contributed by two-fold non-crystallographic symmetry-related monomers (Fig 3F). We believe this junction to be tight, as the interface residues and the interactions between them are similar to those found in protein hydrophobic cores. The atomic temperature factors of these residues are some of the lowest in the whole structure (18 Å² on average, compared with the average B -factor of 38 Å²), which further underscores the stability of this interface. With the aim of producing monomeric TurboGFP, we mutated several clusters of residues within the first interface. Invariably, all mutants were completely insoluble and non-fluorescent on expression in *E. coli*, confirming the importance of the first dimer interface in the context of the overall TurboGFP structure. The second, less extensive dimer interface (monomers A|D and B|C) occludes 703 Å² of molecular surface per monomer and is composed of a mixture of polar and hydrophobic interactions involving residues I88, K90, V96, H98, S100, M117, T119, G146, H173 and H175 (Fig 3G).

In general, the crystal structure of TurboGFP has validated the assumptions made during the design of non-aggregating ppluGFP2 derivatives. The addition of charged termini shields the positively charged patch on the TurboGFP surface and probably contributes to mutual electrostatic repulsion of the C termini. Most of the other surface mutations loosen the weaker dimer–dimer interface, with the exception of L200D, which probably contributes to the overall reduction in protein hydrophobicity.

Least-squares superposition of atomic coordinates of TurboGFP monomer with those of GFP, DsRed, pocilloporin, eqFP611, KFP1 and zFP538 results in $C\alpha$ r.m.s.d. of 1.25, 1.07, 1.04, 1.00, 1.08 and 1.11 Å, respectively. As expected from the sequence alignment, the geometry of the loops (residues 29–32, 66–72, 78–82, 92–95, 105–110, 143–151, 160–165, 175–179, 199–207) that join the conserved structural elements of TurboGFP differ in other members of the GFP family, whereas the core secondary structure elements themselves superimpose well—the exceptions are the extra helix present only in TurboGFP (residues 180–188) and the N-terminal capping helix of GFP (residues 2–9).

Electron density maps indicate complete maturation of all four chromophores in the tetramer. The geometry of the mature TurboGFP chromophore is identical to that of GFP. It is noteworthy that the coloration of the frozen protein crystal completely disappeared after a brief exposure to X-ray radiation. To test this further, a new crystal was tested with the intensity of the beam attenuated to the lowest transmittance ($\sim 1\%$), and still the coloration was eliminated in the course of a single 1-s exposure. As atomic motion is highly restricted in a crystal at 100 K, it is likely that only an electronic configuration rearrangement is taking place, possibly initiated by excitation of outer shell electrons of the S atom of M34, located in the vicinity of the chromophore. Alternatively, a process similar to that described earlier (Matsui *et al*, 2002) might be responsible for this effect.

Overall, the milieu of the TurboGFP chromophore is similar to that of other GFPs, which show an abundance of buried charged side chains typical of internalized catalytic centres. The residues that contribute functional groups to this environment vary between the members of the family, but the chromophore-relative positions of important functional groups are conserved—for example, R87 is the equivalent of GFP R96 and E210 is the equivalent of GFP E222.

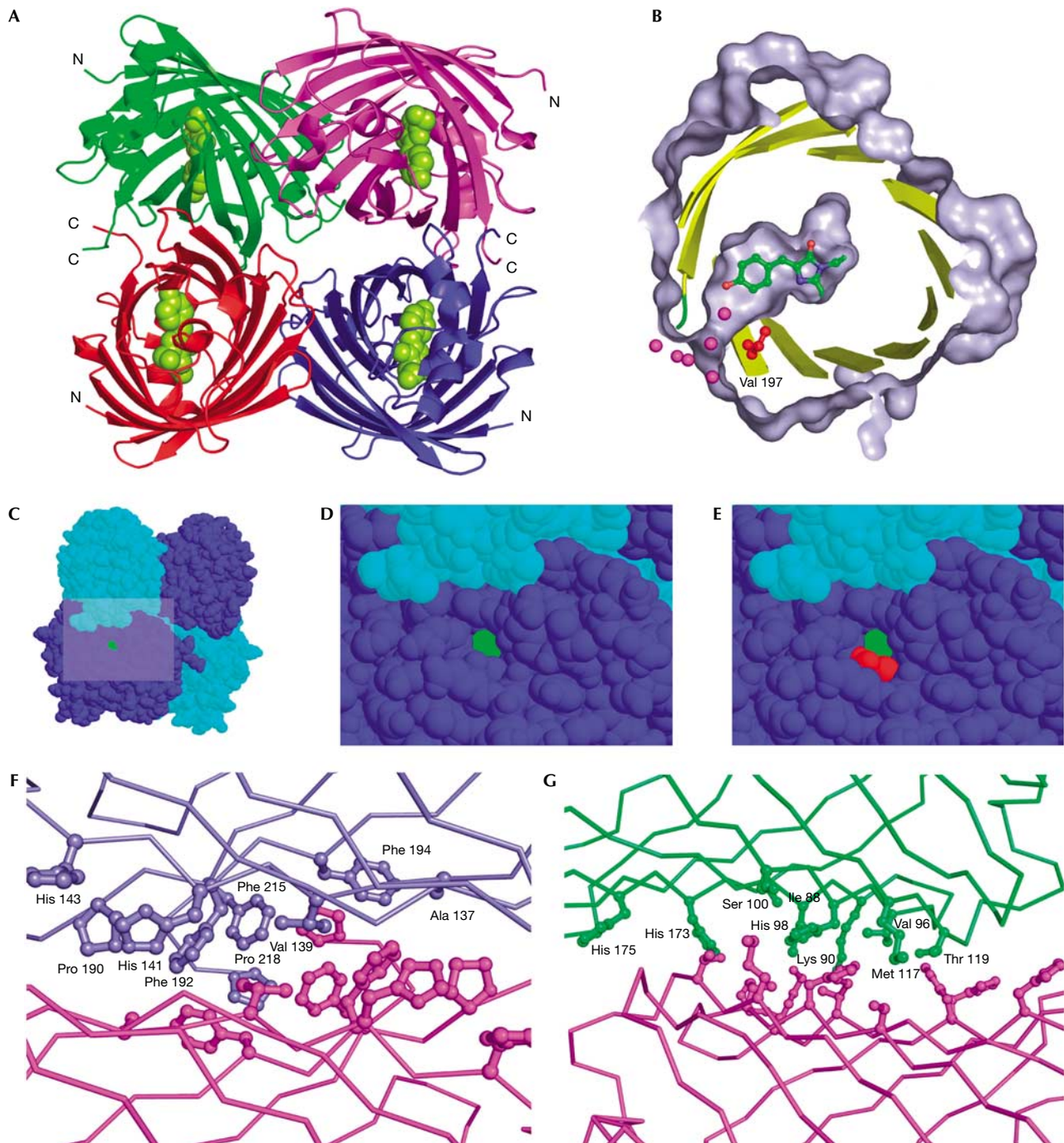


Fig 3 | TurboGFP crystal structure. (A) The overall structure of TurboGFP. α traces are in green, violet, red and blue. The chromophore is shown as green van der Waals spheres. (B–E) A pore leading to the TurboGFP chromophore. The chromophore is highlighted in green and Val 197 in red. (B) Protein surface (grey) is cut away to show the pore and the chromophore cavity. Sections of secondary structure elements are shown as yellow cartoons. Relevant water molecules are depicted as magenta spheres. (C–E) The pore remains unobstructed on tetramerization. (F,G) Contacts between TurboGFP monomers. Two TurboGFP monomers are shown as α traces, with contact residues shown as balls and sticks of the same colour as the particular monomer. For clarity, only one set of residues is labelled. (F) The more extensive contact area having mainly hydrophobic character is shown. (G) The less extensive contact area, composed of a mixture of polar and hydrophobic interactions is shown.

Table 1 | Spectral characteristics and parameters of refolding and refolding/maturation kinetics

Fluorescent protein	ϵ (λ_{\max})*	QY (λ_{\max}) [‡]	Refolding half-time (s)	Maturation half-time (s)	k_{ox} (10^{-4} s^{-1})
EGFP	55,000 (489)	0.60 (509)	90.6	3,915	1.77
Venus [§]	110,000 (515)	0.63 (527)	46.2	4,076	1.70
SYFP2 [§]	101,000 (515)	0.68 (527)	69.3	3,300	2.10
TurboGFP	70,000 (482)	0.53 (502)	11.0	1,468	4.72
TurboGFP-V197L	73,000 (482)	0.47 (502)	9.5	2,493	2.78

Protein refolding and maturation were followed by measuring the recovery of fluorescence at 25 °C. Maturation rate constants (k_{ox}) were determined by computer-fitting the kinetic data to the first-order exponential decay (Origin 6.0).

EGFP, enhanced green fluorescent protein; GFP, green fluorescent protein.

*Extinction coefficient ($\text{M}^{-1} \text{ cm}^{-1}$) with excitation maximum (nm) in parentheses.

[‡]Quantum yield with emission maximum (nm) in parentheses.

[§]Data from Kremers *et al* (2006).

The putative basis for fast TurboGFP maturation

Our data show that TurboGFP fluorescence appears significantly earlier than that of the commonly used EGFP under similar *in vivo* conditions. To understand the basis of this phenomenon, we compared protein-folding and chromophore maturation kinetics for EGFP and TurboGFP in renaturation and renaturation/maturation assays.

First, the recovery of fluorescence during refolding of preheated urea-denatured fluorescent proteins was monitored (see Methods). Denaturation does not affect the chemical structure of the chromophore; therefore, such experiments only show protein-refolding kinetics. In these experiments, TurboGFP demonstrated extremely fast protein-folding kinetics, with a half-life of 11 s, whereas for EGFP the half-life was about 1.5 min (Table 1; Fig 4A). In general, chromophore formation, but not protein folding, is known to be the slowest and time-limiting step in fluorescent protein maturation. However, it is presumed that the fast and effective folding of TurboGFP contributes to the development of early fluorescent signal *in vivo*, by allowing a higher percentage of the protein to escape the protein degradation machinery of the cell. Notably, *in vivo*, faster folding of TurboGFP and ppluGFP2 is due to the absence of *cis*-prolines, which are reported to slow down the folding of *A. victoria* GFP (Enoki *et al*, 2004).

Next, we compared chromophore maturation rates for EGFP and TurboGFP. We used dithionite reduction of a chromophore of urea-denatured protein, commonly used to follow-up fluorescent protein maturation starting from the native polypeptide (Reid & Flynn, 1997; Nagai *et al*, 2002). Maturation curves fitted first-order exponential decay (Origin 6.0), enabling an estimation of maturation rate constants (k_{ox}). For EGFP, k_{ox} was found to be $1.77 \times 10^{-4} \text{ s}^{-1}$, close to the previously reported value (Reid & Flynn, 1997). At the same time, the maturation rate was significantly higher for TurboGFP, with k_{ox} of $4.72 \times 10^{-4} \text{ s}^{-1}$ (Table 1; Fig 4B).

Ultimately, we believe that both rapid protein folding and chromophore maturation contribute to early fluorescent signal development by Copepoda fluorescent proteins. Although protein folding depends on many parameters, faster chromophore maturation could be explained by the chromophore environment.

It is considered that chromophore maturation in GFP-like proteins proceeds in two stages: cyclization of amino acids 65–67

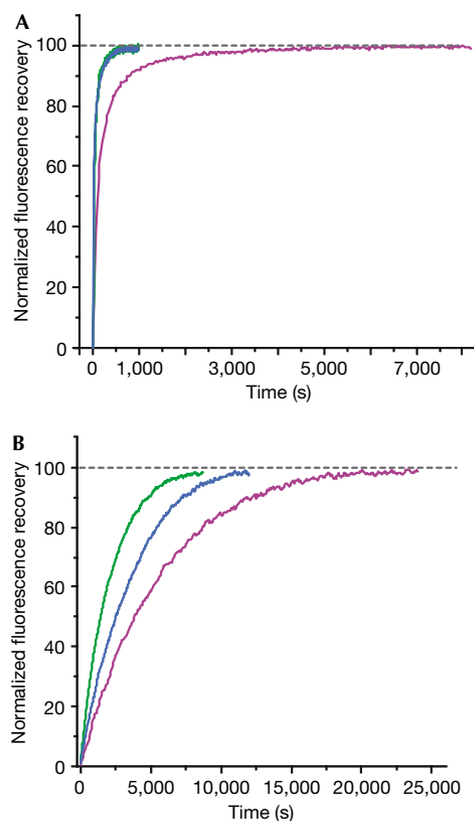


Fig 4 | Comparison of refolding and maturation speed *in vitro* of EGFP, TurboGFP and TurboGFP-V197L. Normalized fluorescence recovery plots are shown. EGFP, violet lines; TurboGFP, green lines; TurboGFP-V197L, blue lines. See Methods and Table 1 for details. (A) Refolding kinetics. (B) Chromophore maturation kinetics.

(using GFP residue numbers), followed by the slowest and thus time-limiting step, dehydrogenation of the Y66 C α -C β bond (Reid & Flynn, 1997). In the presence of molecular oxygen, this reaction proceeds spontaneously, although protein environment strongly influences the reaction kinetics (Kojima *et al*, 1998). The recently

proposed chromophore maturation mechanism (Rosenow *et al*, 2004) implies a dehydrogenation step initiated by Y66 C α deprotonation by base. Interestingly, TurboGFP contains a unique E89 residue, which could aid in proton abstraction (see supplementary information 4 online).

Another unique feature of TurboGFP compared with other fluorescent proteins is the water-filled pore, leading from the outside of the protein barrel to Y58 of the chromophore (Fig 3B–E). This pore remains unobstructed on tetramerization. Importantly, molecular oxygen is required for the rate-limiting step of chromophore dehydrogenation (Cody *et al*, 1993; Heim *et al*, 1994; Inouye & Tsuji, 1994). Therefore, it can be presumed that the pore facilitates oxygen conveyance to the premature chromophore, thus speeding up maturation.

To verify this hypothesis, we performed site-directed mutagenesis of TurboGFP, introducing leucine instead of valine in position 197 (205 in GFP), shown in red in Fig 3B,E. This mutation should narrow the pore and therefore impair the proposed oxygen-conveyance mechanism. The resulting protein TurboGFP-V197L was brightly fluorescent, although its fluorescence signal was generated much more slowly when expressed in *E. coli*. Its renaturation kinetics was shown to be as fast as that of TurboGFP, indicating that this mutation does not impair protein folding (Table 1; Fig 4A). Also, the fluorescence characteristics of mutants, such as excitation/emission spectra, fluorescence quantum yield and extinction coefficient, were almost identical to those of the parent TurboGFP (Table 1), indicating that the chromophore environment was not significantly altered. The chromophore maturation kinetics of TurboGFP-V197L was markedly slower (Table 1; Fig 4B). This suggests that the pore found in TurboGFP/ppluGFP2 is essential for fast maturation of the chromophore, most probably as an access route for oxygen conveyance to the premature chromophore.

As an alternative hypothesis, it is proposed that water molecules coordinated in the pore can participate in proton abstraction from the Y66 C α atom. This facilitates the dehydrogenation step of chromophore maturation, in common with the model of Rosenow *et al* (2004), implying that the water molecule is the base responsible for proton abstraction.

Rapidly maturing fluorescent proteins are highly desirable tools for: monitoring the activity of promoters; studying the protein degradation machinery of the cell; and a wide range of applications in which a rapid generation of fluorescent signal is crucial. TurboGFP can be used *in lieu* of EGFP in applications that are not strictly dependent on the monomeric nature of the tag, but demand rapid signal detection. We believe that our findings could contribute to the rational design of rapidly maturing monomeric fluorescent proteins, suitable for fluorescence resonance energy transfer-based protein interaction studies and for fastidious fusions.

In general, the TurboGFP structure is intriguing, as it is the first reported non-Cnidarian fluorescent protein structure solved. We believe that it will have a significant impact on the further understanding of the evolution, functions and chromophore biophysics of GFP-like proteins.

METHODS

Construction of protein mutants. For bacterial expression, the DNA encoding the codon-optimized version of ppluGFP2 from

P. plumata (Shagin *et al*, 2004) was amplified using specific primers and cloned into Qiagen pQE30 vector using *Bam*HI–*Hind*III restriction sites. Site-directed mutagenesis was performed by overlap-extension PCR, with primers containing the appropriate target substitutions (Ho *et al*, 1989).

Expression in mammalian cell lines and imaging. For expression in eukaryotic cells, a PCR-amplified *Agel*–*Bgl*II fragment encoding TurboGFP or ppluGFP2 was swapped with EGFP in pEGFP-C1 vector (Clontech), resulting in TurboGFP-C1 or ppluGFP2-C1 plasmid. To generate TurboGFP– β -actin or ppluGFP2– β -actin fusion proteins, the corresponding gene was swapped with EGFP in a pEGFP–actin vector (Clontech). Fluorescence images of transfected HeLa cells were obtained using the Olympus CK40 inverted microscope equipped with the Olympus DP50 camera.

Protein purification. Proteins fused to the N-terminal polyhistidine tag were expressed in *E. coli* XL1-blue strain (Stratagene, La Jolla, CA, USA) and purified using TALON metal-affinity resin (Clontech). The polyhistidine tag was not removed.

Spectral measurements. Absorption spectra were recorded with a Beckman DU520 UV/VIS Spectrophotometer. The Varian Cary Eclipse Fluorescence Spectrophotometer was used for measuring excitation–emission spectra. For molar extinction coefficient determination, we relied on estimating the mature chromophore concentration. An alkali-denatured GFP chromophore absorbs at 446 nm with a molar extinction coefficient $44\,000\text{ M}^{-1}\text{ cm}^{-1}$. Molar extinction coefficients for the native state were estimated from the absorption of denatured proteins. For quantum yield determination, the fluorescence of the proteins was compared with equally absorbing EGFP (quantum yield 0.60; Patterson *et al*, 2001).

Refolding and maturation kinetics. Samples of fluorescent proteins were heated to 95 °C in denaturation solution (8 M urea, 1 mM dithiothreitol) for 4 min. Refolding reactions were initiated following 100-fold dilution in the renaturation buffer (35 mM KCl, 2 mM MgCl₂, 50 mM Tris pH 7.5, 1 mM dithiothreitol). In the maturation assay, 5 mM freshly dissolved dithionite was added to the denaturation solution (Reid & Flynn, 1997). Owing to the instability of dithionite at high temperatures and to provide for complete chromophore reduction, the sample was cooled to 25 °C and the addition of 5 mM dithionite followed by heating to 95 °C were repeated. Protein refolding and maturation were followed by measuring the recovery of fluorescence using the Varian Cary Eclipse Fluorescence Spectrophotometer, with the chamber temperature maintained at 25 °C.

Supplementary information is available at *EMBO reports* online (<http://www.emboreports.org>).

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