J. Soils and Crops 25 (1) 13-20, June, 2015 ISSN 0971-2836 THE MOLECULAR STRUCTURE OF GREEN FLUORESCENT PROTEIN

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ABSTRACT

The research was done in department of Biochemistry and Cell Biology and the Reddick center for computational Biology, Edinburg university, Edinburg during year 2013 .The crystal structure of recombinant wild-type green fluorescent protein (GFP) has been solved to a resolution of 1.9 Å by multi-wavelength anomalous dispersion (MAD) phasing methods. The protein is in the shape of a cylinder, comprising 11 strands of -sheet with an helix inside and short helical segments on the ends of the cylinder. This motif with -structure on the outside and -helix on the inside, represents a new protein fold, which we have named the?-can. Two promoters pack closely together to form a dimer in the crystal. The fluorophores are protected inside the cylinders, and their structures are consistent with the formation of aromatic systems made up of Tyr⁶⁶ with reduction of its C - C bond coupled with cyclization of the neighboring glycine and serine residues. The environment inside the cylinder explains the effects of many existing mutants of GFP and suggests which side chains could be modified to change the spectral properties of GFP. Furthermore, the identification of the dimer contacts may allow mutagenic control of the state of assembly of the protein. It is concluded that, the study of protein dimer indices will helpful in identifying the new era for rice research which will help in developing protein rich rice which can provide a dietary supplement for people.

(Key words: Protein, wavelength, flurophore, amino acids)

INTRODUCTION

Green fluorescent protein, GFP, is a spontaneously fluorescent protein isolated from coelenterates, such as the Pacific jellyfish, Aequoria *victoria*¹. Its role is to transduce, by energy transfer, the blue chemiluminescence of another protein, aequorin, into green fluorescent light (Ward, 1979). The molecular cloning of GFP cDNA (Prasher et al., 1992) and the demonstration by Chalfie that GFP can be expressed as a functional transgene (Chalfie et al.,1994) have opened exciting new avenues of investigation in cell, developmental and molecular biology. Fluorescent GFP has been expressed in bacteria (Chalfie et al., 1994), yeast (Kahana et al.,1995), slime mold (Moores et al.,1996), plants (Casper and Holt, 1996; Epel et al., 1996), drosophila (Wang and Hazelrigg, 1994), zebrafish (Amsterdam et al., 1996), and in mammalian cells (Ludin et al., 1996; DeGiorgi et al., 1996). GFP can function as a protein tag, as it tolerates N- and C-terminal fusion to a broad variety of proteins many of which have been shown to retain native function (Moores et al., 1996; Cubitt et al.,1995; Olsen et al.,1995). When expressed in mammalian cells fluorescence from wild type GFP is typically distributed throughout the cytoplasm and nucleus, but excluded from the nucleolus and vesicular organelles (reviewed by Cubitt et al., 1995, LG Moss unpublished observations). However, highly specific intracellular localization including the nucleus, mitochondria (Rizzuto et al., 1996),

secretory pathway Kaether and Gerdes, 1995), plasma membrane (Marshall et al., 1995) and cytoskeleton (Kahana et al., 1995) can be achieved via fusions both to whole proteins and individual targeting sequences. The enormous flexibility as a noninvasive marker in living cells allows for numerous other applications such as a cell lineage tracer, reporter of gene expression and as a potential measure of proteinprotein interactions (Mitra et al., 1996). Green fluorescent protein is comprised of 238 amino acids. Its wild-type absorbance/ excitation peak is at 395 nm with a minor peak at 475 nm with extinction coefficients of roughly 30,000 and 7,000 M⁻¹ cm⁻¹, respectively(Kahana and Silver, 1996). The emission peak is at 508 nm. Interestingly, excitation at 395 nm leads to decrease over time of the 395 nm excitation peak and a reciprocal increase in the 475 nm excitation band (Cubitt et al., 1995). This presumed photoisomerization effect is especially evident with irradiation of GFP by UV light. Analysis of a hexapeptide derived by proteolysis of purified GFP led to the prediction that the fluorophore originates from an internal Ser-Tyr-Gly sequence which is posttranslationally modified to 4(phydroxybenzylidene)imidazolidin-5-one, structure (Cody et al., 1993). Studies of recombinant GFP expression in E. coli led to a proposed sequential mechanism initiated by a rapid cyclization between set⁶⁵ and Gly⁶⁷ to form a imidazolin-5 one intermediate followed by a much slower (hours) rate-limiting oxygenation of the Tyr⁶⁶ side chain by O₂ (Heim et al., 1994). Combinatorial

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mutagenesis suggests that the Gly⁶⁷ is required for formation of the fluorophore (Delagrave *et al.*,1995). While no known co-factors or enzymatic components are required for this apparently auto-catalytic process, it is rather thermosensitive with the yield of fluorescently active to total GFP protein decreasing at temperatures greater than 30 C(Lim *et al.*,1995). However, once produced GFP is quite thermostable.

Physical and chemical studies of purified GFP have identified several important characteristics. It is very resistant to denaturation requiring treatment with 6 M guanidine hydrochloride at 90 C or pH of <4.0 or >12.0. Partial to near total renaturation occurs within minutes following reversal of denaturing conditions by dialysis or neutralization (Ward and Bokman, 1982). Circular dichroism predicts significant amounts of sheet structure that is subsequently lost on denaturation. (Ward and Bokman, 1982) Over a non-denaturing range of pH, increasing pH leads to a reduction in fluorescence by 395 nm excitation and an increased sensitivity to 475 nm excitation (Ward et al., 1982). Reduction of purified GFP by sodium dithionite results in a rapid loss of fluorescence that slowly recovers in the presence of room air. While insensitive to sulfhydryl reagents such as 2-mercaptoethanol, treatment with the sulfhydral reagent di-thiobisnitrobenzoic acid (DTNB) irreversibly eliminates fluorescence(Inouve and Tsuji, 1994).

The availability of E. coli clones expressing GFP has led to extensive mutational analysis of GFP function. Truncation of more than 7 amino acids from the C-terminus or more than the N-terminal Met lead to total loss of fluorescence (Dopf and Horiagan, 1996). All non-fluorescent mutants also failed to exhibit absorption spectra characteristic of the intact fluorophore, implying a possible defect in post-translational processing. Screens of random and directed point mutations for changes in fluorescent behavior have uncovered a number of informative amino acid substitutions. Mutation of Tyr⁶⁶ in the fluorophore to His results in a shift of the excitation maximum to the UV (383 nm) with emission now in the blue at 448 nm (Heim et al., 1994). A Tyr⁶⁶Trp mutant is blue-shifted albeit to a lesser degree. Both changes are associated with a severe weakening of fluorescence intensity compared to wild type GFP. Mutation of Ser⁶⁵ to Thr, Ala, Cys or Leu causes a loss

of the 395 nm excitation peak with a major increase in blue excitation (Delagrave et al., 1995 ; Heim et al.,1995). When combined with Ser⁶⁵ mutants, mutations at other sites near the fluorophore such as Val68Leu and Ser72Ala can further enhance the intensity of green fluorescence produced by excitation at 488 nm (Delagrave et al., 1995 ; Cormack et al., 1996). However, amino acid substitutions significantly outside this region also affect the protein's spectral character. For example, Ser²⁰²Phe and Thr²⁰³Ile both cause the loss of excitation in the 475 nm region with preservation of 395 nm excitation (Chalfie et al., 1994 ; Heim et al., 1994 ; Ehrig et al., 1995). Ile¹⁶⁷ Thr results in a reversed ratio of 395 to 475 nm sensitivity (Cubitt et al., 1995), while Glu²²² Gly is associated with the elimination of only the 395 nm excitation (Ehrig et al., 1995). Another change, Val¹⁶³ Arg, not only enhances the magnitude of the Ser⁶⁵ Thr mutant, but also increases the temperature tolerance for functional GFP expression (Kahana and Silver, 1996). Molecular evolution techniques have been reported to improve GFP fluorescence (Crameri et al., 1996). Unfortunately, a roster of substitutions associated with complete loss of function has not been published.

Because GFP in crystallum exhibits a nearly identical fluorescence spectrum and lifetime to that for GFP in aqueous solution (Perozzo et al., 1988) and fluorescence is not an inherent property of the isolated fluorophore, the elucidation of its threedimensional structure will help provide an explanation for the generation of fluorescence in the mature protein, as well as the mechanism of autocatalytic fluorophore formation. Furthermore, the development of fluorescent proteins with additional emission and excitation characteristics would dramatically expand their biological applications. Color vision is based on the fact that spectral properties of a common fluorophore, cisretinal, are altered as a function of protein environment within red, blue, or green opsins (Merbs and Nathans, 1992). The GFP from the sea pansy, Renilla reniformis, which exhibits a single major excitation peak at 498 nm, apparently utilizes an identical core fluorophore to that of A. victoria GFP. These findings taken together with the spectral changes exerted by substitutions in amino acids over

100 residues from the GFP fluorophore suggest that a rational strategy to modify and expand the fluorescence behavior of GFP based on protein structure may be possible. Here we report the X-ray diffraction structure derived from a crystal of wild-type, recombinant *A. victoria* green fluorescent protein.

MATERIALS AND METHODS

Green fluorescent protein was purified from *E. coli* strain BL21(DE3)pLysS (Novagen) containing plasmid pTu58, bearing the wild-type Aequorea victoria green fluorescent protein (Chalfie et al., 1994). For the seleniomethionine protein, the plasmid was moved to E. coli methionine auxotroph strain B834(DE3)pLysS (Novagen). The purification involved cell lysis, centrifugation of cell debris, and four column chromatography steps: DEAE anion exchange column (Sigma, CL-6B) with a zero to 1M NaCl gradient in 10 m M phosphate, 2 m M EDTA, 2 m M DTT, pH 7; a hydrophobic interaction column (Sigma, CL-4B) with a 0.1 to zero M phosphate gradient in 2 m M EDTA, 2 m M DTT, pH 7; an HPLC anion exchange column (Bio-Rad, Bio-Gel DEAE-5PW) with a zero to 1M NaCl gradient in 10 mM phosphate, 2 m M EDTA, 2 m M DTT, pH 7; and an HPLC gel filtration column (Bio-Rad, Bio-Gel SEC-125) with 0.1 M phosphate, 2 m M EDTA, 2 m M DTT, pH 7. Gel filtration columns run at 10 m M phosphate showed predominately a 2-fold higher molecular weight species. Matrix-assisted laser desorption ionization mass spectrometry was performed by the University of Texas Health Sciences Center analytical chemistry service.

The protein was crystallized in sitting drop vapor diffusion wells (Hampton Research) at room temperature using 58% 2-methyl-2,4-pentanediol (Aldrich), 50mM morpholino ethane sulfonic acid, 0.1% sodium azide at pH 6.8. The protein concentration varied, but was typically 20-30 mg ml⁻¹. Crystals grew as green fluorescent square bipyramids up to 0.5 mm on a side. The space group was determined to be P41212 or its enantiomorph, with a=b=87.15 Å and c=119.85 Å at cryogenic temperatures, and a=b= 89.23 Å and c= 119.78 Å at room temperature. The unit cell also varies with changes in ionic strength, and this effect thwarted solution by multiple isomorphous replacement. Packing density calculations suggested that there were probably two molecules per asymmetric unit.

Multi-wavelength anomalous dispersion (MAD) data were taken at Brookhaven beam line X4A at four wavelengths. The wavelengths to be used were determined by reference and crystal absorption scans. The data were taken at liquid nitrogen temperature using inverse-beam geometry in wedges of four degrees and processed using DENZO (Otwinowski, 1993). Native and selenio-methionine data sets were also taken in the laboratory on an R-AXIS IIC detector with CuK radiation. The native data set used in the refinement had a R_{merge} of 7.7% to 1.9 Å resolution (99+% complete in all shells with 5fold average redundancy). Selenium atoms were located initially by standard difference Patterson maps between selenium-substituted and native protein using SHELX96 (Sheldrick et al., 1993) and HEAVY (Terwilliger et al., 1987) and confirmed by Patterson maps using the MAD data. MADSYS software (Yang et al., 1990) was used to give the anomalous diffraction differences shown in table 2 and to extract Fa, Ft, and phase information.

The resulting MAD-phased map was solvent flattened and two-fold averaged based on the selenium sites using CCP4 (Anonymous, 1994), skeletonized using the program O (Jones et al., 1991), and immediately revealed two 11-stranded cylindrical -barrels. The polypeptide chain was traced for one of the barrels beginning from the seleniomethionines and extending the structure in each direction, helped by the recognition of the modified tyrosine in the middle of the barrel as Tyr⁶⁶, the nucleus of the fluorophore. The correct enantiomorph is space group $P4_12_12$, as confirmed by the handedness of the ?-barrel and the ?-helices. Refinement has been started using the program X-Plor (Brunger, 1992) using the native data collected at room temperature; the current R-factor at 1.9 Å is 0.21 with an R-free of 0.26, with good geometry (rms) bond and angle deviations from ideality of 0.013 Å and 1.8, respectively) and tight restraint of the noncrystallographic symmetry. All measured data were included in the refinement. Coordinates and structure factors have been deposited at the Brookhaven Protein Data Bank under accession numbers 1GFL and R1GFLSF, respectively.

RESULTS AND DISCUSSION

The structure of GFP has been solved using seleniomethionyl-substituted protein and multiwavelength anomalous dispersion (MAD) phasing methods. The electron density maps produced by the MAD phasing were very clear, revealing a dimer comprised of two quite regular -barrels with 11 strands on the outside of cylinders (Figure 1,2,3). These cylinders have a diameter of about 30 Å and a length of about 40 Å. Inspection of the density within the cylinders reveals modified tyrosine side chains as a part of an irregular -helical segment (Figure 4). Small sections of -helix also form caps on the ends of the cylinders. This motif, with a single -helix inside a very uniform cylinder of ?-sheet structure, represents a new protein class, as it is not similar to any other known protein structure.

The fluorophore is highly protected, located on the central helix within a couple of Ångstroms of the geometric center of the cylinder. The pocket containing the fluorophore has a surprising number of charged residues in the immediate environment (Figure 5 and Table 1). The environment around the fluorophore includes both apolar and polar amino acid side chains. Phe⁶⁴ and Phe⁴⁶ are near the fluorophore and separate the single tryptophan, Trp⁶³ from direct contact with fluorophore (closest distance of 13 Å). A table of all atoms that come in contact with the fluorophore and their distances to the fluorophore is provided (Table 1).

The crystallographic contacts are all rather tenuous, consisting of a few amino acids side chains for each. The non-crystallographic symmetry is maintained by extensive contacts and thus is likely to be the source of the dimerization seen in solution studies (Figure 6). The dimer contacts are fairly tight and consist of a core of hydrophobic side chains from Ala²⁰⁶, Leu²²¹, and Phe²²³ from each of the two monomers and a wealth of hydrophilic contacts (Figure 5), including Tyr³⁹, Glu¹⁴², Asn¹⁴⁴, Asn¹⁴⁶, Ser¹⁴⁷, Asn¹⁴⁹, Tyr¹⁵¹, Arg¹⁶⁸, Asn¹⁷⁰, Glu¹⁷², Tyr²⁰⁰, Ser²⁰², Gln²⁰⁴, and Ser²⁰⁸. Contacts with other crystallographic molecules are not extensive, and the salt-dependence of this dimer interface and/or the loose contacts with neighboring molecules may explain the difficulties with isomorphism in initial heavy atom phasing studies.

Mass spectrometry studies of the bacterially expressed wild-type and selenio- methionyl protein show masses of 26836.1 (±0.9) and 27069.3 (±1.4) g mole⁻¹, respectively. The masses calculated for the known pTu58 gene sequence, including the original inadvertent Gln⁸⁰Arg PCR error during the cloning of the gene for GFP (Chalfie et al., 1994) and the cyclization and oxidation of the tyrosine are 26835.5 and 27070.0 for the seleniomethionine, respectively. The differences of 0.6 and -0.7 g mole⁻¹ are small and the results are therefore consistent with essentially complete fluorophore formation, including the loss of water after cyclization. The error limits do not allow accurate determination of the degree of oxidation of the dehydrotyrosine, however. These results indicate the starting material for the crystallization was essentially fully formed GFP and the lack of difference density in Fo-Fc maps in this region shows that the crystal contains fully cyclized GFP.

The remarkable cylindrical fold of the protein seems ideally suited for the function of the protein. The strands of -sheet are tightly fitted to each other like staves in a barrel, and form a regular pattern of hydrogen bonds. Together with the short -helices and loops on the ends, the 'can' structure forms a single compact domain and does not have obvious clefts for easy access of diffusable ligands to the fluorophore. This fold, taken with the observation that the fluorophore is near the geometric center of the molecule explains the observed protection of the fluorophore from collisional quenching by oxygen $(K_{hm} < 0.004 \text{ M}^{-1}\text{s}^{-1})^{34}$ and hence reduction of the quantum yield. Perhaps more seriously, photochemical damage by the formation of singlet oxygen through intersystem crossing is reduced by the structure. The tightly constructed -can would appear to serve this role nicely, as well as provide overall stability and resistance to unfolding by heat and denaturants.

The location of certain amino acid side chains in the vicinity of the fluorophore also begins to explain the fluorescence and the behavior of certain mutants of the protein. At least two resonant forms of the fluorophore can be drawn, one with a partial negative charge on the benzyl oxygen of Tyr⁶⁶, and one with the charge on the carbonyl oxygen of the imidazolidone ring. Interestingly, basic residues appear to form hydrogen bonds with each of these oxygen atoms, His¹⁴⁸ with Tyr⁶⁶ and Gln⁹⁴ and Arg⁹⁶ with the imidazolidone. These bases presumably act to stabilize and possibly further delocalize the charge on the fluorophore. Most of the other polar residues in the pocket form an apparent hydrogen-bonding network on the side of Tyr⁶⁶ that requires abstraction of protons in the oxidation process. It is tempting to speculate that these residues help abstract the protons. As for the mutants, atoms in the side chains of Thr²⁰³, Glu²²², and Ile¹⁶⁷ are in van der Waals contact with Tyr⁶⁶, so their mutation would have direct steric effects on the fluorophore and would also change its electrostatic environment if the charge were changed, as suggested previously (Ehrig et al., 1995). A quantitative explanation will require further examination. It seems likely that other mutations of the residues identified to be near the fluorophore would also have effects on the absorption and/or emission spectra, and such experiments to change the electrostatic environment around the fluorophore are in progress. By virtue of their varied fluorophore environments and hence altered spectra, these mutants should lead to expanded uses of green fluorescent protein as gene markers, cell lineage markers, and encourage other uses in biotechnology.

Mutations in regions of the sequence adjacent to the fluorophore, i.e. in the range of positions 65-67, have been systematically explored (Delagrave et al., 1995), some having significant wavelength shifts and most suffer a loss of fluorescence intensity. For example, mutation of the central Tyr to Phe or His shifts the excitation bands but there is an overall loss of intensity. Secondary mutations to compensate for the deleterious intensity effects should also now be possible. The Ser⁶⁵Thr mutant is particularly interesting because of its reported increase in fluorescence intensity (Heim et al.,1994; Heim et al.,1995). The mechanism for increased fluorescence may be reduced collisional quenching, as the additional methyl group may make for better packing in the interior of the protein. On the other hand, the effect has been suggested to be through improved conversion of the tyrosine to dehydrotyrosine. However, the fact that we see significantly altered structure relative to standard protein conformations in the wild-type argues against a dramatic increase in cyclization and/or oxidation. This effect is most likely produced by increased expression and/or folding of the protein. The crystal structure of the Ser⁶⁵Thr mutant has also been solved (Ormo *et al.*,1996), and it will be interesting to compare the two structures for clues about the fluorescence and other differences. The report of improvements in GFP by DNA shuffling (Crameri *et al.*,1996), comprising mutations Phe¹⁰⁰Ser, Met¹⁵⁴Thr and Val¹⁶⁴Ala are difficult to explain based on the structure. Positions 154 and 164 are on the surface of the protein and may exert their effects through improved solubility and/or reduced aggregation. The Phe-Ser mutation at first glance would appear to destabilize the core of the protein and we have no idea how it would improve the system.

The mechanism of activation of the fluorophore from ordinary protein structure is consistent with a non-enzymatic cyclization mechanism like that of Asn-Gly deamidation (Wright, 1991) followed by oxidation of the tyrosine to dehydrotyrosine, as previously suggested. The role of molecular oxygen in this mechanism and in GFP fluorescence is paradoxical, however. Molecular oxygen is proposed to be needed for formation of the double bond between C and C on the tyrosine to form an extended aromatic system, but oxygen must also be excluded from regular interactions with the fluorophore or else collisional quenching of the fluorescence or damaging photochemistry will occur. The low bimolecular quenching rate suggests that the protein's design sacrifices efficient fluorophore formation for stability and higher quantum yields once fully formed.

The excited state dynamics of GFP have been studied using Stark, steady state, and time-resolved fluorescence spectroscopies (Chattoraj *et al.*,1996) (and Youvan and Michel-Beyerle, personal communication). The results suggest that proton transfer is involved in interconversions within two ground and two excited states. The extended set of polar interactions around the flourophore could easily accomodate proton rearrangements, with the most likely direct effects being associated with the His¹⁴⁸ with the hydroxyl of Tyr⁶⁶, Arg⁹⁶ interactions with the imidazolidone, or Glu²²² interactions with the hydroxyl of Ser⁶⁵. Since the Ser⁶⁵/Glu²²² mutants have both lost their native interactions together with their ~400 nm absorption bands, one possibility is that

the 400nm band arises from the abstraction of the Ser⁶⁵ hydroxyl proton by Glu²²². This is speculation however, but similar spectroscopic studies on mutants at these positions may be able to differentiate the roles of these sites in excited state dynamics.

The N- and C-termini truncation studies(Dopf and Horiagan, 1996) and the fluorescent fusion products(Moores et al., 1996;Cubitt et al.,1995;Olsen et al.,1995) are now understandable, given the structure of the protein. Since the Cterminus loops back outside the cylinder and the last seven or so amino acids are disordered it shouldn't be critical to have them present and further addition would seem to be easily tolerated. These residues do not form a stave of the barrel. The role of the Nterminus is a little less clear, as the first strand in the barrel does not begin until amino acid 10 or 11. Thus barrel formation does not require the N-terminal region. The N-terminal segment, is however, an integral part of the 'cap' on one end of the protein, and may be essential in folding events or in protecting the fluorophore. Again, extensions at the N-terminus would not disrupt the motif structure of the protein.

The chemical modification studies (Inouye and Tsuji,1994) using sulfhydral reagents can be partially explained. Reaction of one of the cysteines near one end of the cylinder, Cys⁷⁰, would appear to disrupt the packing of the cap on that end, and hence allow quenching of the fluorophore. Significant fluorescence intensity effects by the modification of Cys⁴⁸ on the exterior of the protein would not be expected, a priori. The structure determination of the dithionite-reduced, non-fluorescent species has not yet been studied, but should provide additional data on the nature of the fluorophore. The pH dependence of the excitation bands at 395 nm and 475 nm (Ward et al.,1982) is almost certainly due to His¹⁴⁸, whose N atom is 3.3 Å from the Tyr⁶⁶ hydroxyl oxygen atom of the fluorophore, although NMR pKa measurements or mutagenesis studies would be needed for confirmation.

The dimer we see as the asymmetric unit in

the crystal is likely to be the same one formed in solution, since the ionic strength of the crystallization buffer is low, and we see dimers at low (<100 mM) ionic strengths in solution. Thus, it is not surprising to us to see the large number of hydrophilic dimer contacts. The smaller hydrophobic patch could conceivably be involved in physiological interactions with aequorin, as there would be a natural advantage to close proximity for efficient energy transfer. It is not known at present whether dimers form in physiological circumstances, or what the effect of dimerization is on energy transfer, aside from the circumstantial inferences on the excitation spectra previously reported on the native protein(Ward et al., 1982). The dimer contacts should now be able to be modified in such a way to disrupt the formation of dimers without affecting stability and folding. Other nearby residues could also be converted to hydrophobic residues to enhance dimer stability if desired. Control of the dimerization will be important for fluorescence resonance energy transfer (FRET) studies of protein-protein interactions using GFP, as one would not want to induce association and hence resonance energy transfer between the differently colored GFP proteins by mechanisms other that of the target protein interactions. Mutants may also be developed for reduction of aggregation during expression and hence fewer problems with inclusion bodies.

Thus, the three-dimensional structure of GFP has provided a physico-chemical basis of many observed features of the protein, including its stability, protection of it fluorophore, behavior of mutants, and dimerization properties. The structure will also allow directed mutation studies to complement random and combinatorial approaches.

From above research it is concluded that, the study of protein dimer indices will helpful in identifying the new era for rice research which will help in developing protein rich rice which can provide a dietary supplement for people.

Prot	ein	Fluoro	Distance		
Residue	Atom	Residue	Atom	(Å)	
Arg 96	NH2	Tyr 66	Ο	2.7	
Gln 94	NE2	Tyr 66	Ο	3.0	
His 148	ND1	Tyr 66	OH	3.3	
Gln 69	CD	Tyr 66	Ο	3.4	
Glu 222	OE2	Tyr 66	CE2	3.5	
Val 150	CG2	Tyr 66	CE1	3.6	
Phe 165	CE1	Tyr 66	CD1	3.6	
Thr 203	CG2	Tyr 66	CE2	3.6	
Ile 167	CD1	Tyr 66	OH	3.7	
Thr 62	CG2	Tyr 66	CG	3.7	
Tyr 145	CE2	Tyr 66	OH	3.7	
Ser 205	OG	Tyr 66	CE2	4.0	
Val 61	CG1	Tyr 66	CE2	4.4	
Gln 183	NE2	Tyr 66	Ο	4.8	
Val 68	CG2	Ser 65	С	4.9	

Table 1. List of amino acid side chains with close contacts (less than 5 Å) to the fluorophore. The fluorophore is defined as the 7 atoms of the phenol of Tyr^{66} , the 6 atoms of the imidazolidone, and the bridging methylene between the rings. The following amino acid side chains would be expected to have the most direct effects on fluorescence and perhaps fluorophore formation. The atom names are taken from the Brookhaven Protein Data Bank nomenclature

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Table 2. Anomalous diffraction differences and scattering factors for seleniomethionyl GFP at the four wavelengths used. Following the format used by Yang *et al.* (1990), Bijvoet differences ratios are given in diagonal elements with centric values in parentheses, and dispersive differences are given in off-diagonal elements. Scattering factors were chosen to minimize the cumulative errors in Bijvoet and dispersive terms

Wave-													scat	tering
	30 > d > 3.4 (Å)			3.4 > d > 2.7 (Å)		2.7 > d > 2.2 (Å)				5				
length														
(Å)	0.9879	0.9794	0.9792	0.9686	0.9879	0.9794	0.9792	0.9686	0.9879	0.9794	0.9792	0.9686	f	f"
0.9879	0.026	0.042	0.030	0.020	0.037	0.046	0.037	0.032	0.064	0.071	0.068	0.064	4.0	1.1
	(0.026)				(0.035)				(0.052)					
0.9794		0.050	0.026	0.045		0.058	0.035	0.049		0.088	0.065	0.077	- 10.5	3.9
		(0.029)				(0.039)				(0.060)				
0.9792			0.067	0.032			0.074	0.040			0.101	0.071	7.9	5.5
			(0.031)				(0.042)				(0.062)			
0.9686				0.049				0.059				0.089	3.4	3.9
				(0.027)				(0.039)				(0.064))	



Figure 1. The overall shape of the protein and its association into dimers. Eleven strands of -sheet (green) form the walls of a cylinder. Short segments of -helices (blue) cap the top and bottom of the '? -can' and also provide a scaffold for the fluorophore which is near geometric center of the can. This folding motif, with -sheet outside and helix inside, represents a new class of proteins. Two monomers are associated into a dimer in the crystal and in solution at low ionic strengths. This view is directly down the two-fold axis of the non-crystallographic symmetry. Figures 1 and 6 were produced with Ribbons (Carson, 1987).



Figure 2. Stereo view of a monomer, with colors that vary slowly as a function of the distance along the polypeptide chain. The termini and C atoms of every 20th amino acid are marked just to the upper right of each atom. Figure produced by RasMol(Sayle and Milner-White,1995).



Figure 3. A topology diagram of the folding pattern in GFP. The -sheet strands are shown in light green, ? -helices in blue, and connecting loops in yellow. The positions in the sequence that begin and end each major secondary structure element are also given. The anti-parallel strands (except for the interactions between stands 1 and 6) make a tightly formed barrel



Figure 4. Model of the fluorophore and its environment superposed on the MAD-phased electron density map at 2.2 Å resolution. The clear definition throughout the map allowed the chain to be traced and side chains to be well placed. The density for Ser⁶⁵, Tyr⁶⁶ and Gly⁶⁷ is quite consistent with the dehydrotyrosine - imidazolidone structure proposed for the fluorophore. Many of the side chains adjacent to the fluorophore are labeled. Figures 4 and 5 were produced with O^{43}



Figure 5. Stereo view of the fluorophore and its environment. His¹⁴⁸, Gln⁹⁴ and Arg⁹⁶ can be seen on opposite ends of the fluorophore and probably stabilize resonant forms of the fluorophore. Charged, polar, and non-polar side chains all contact the fluorophore in some way.



Figure 6. The dimer contact region. The two polypeptide chains associate over a broad area, with a small hydrophobic patch (in the yellow box) and numerous hydrophilic contacts. The two-fold symmetry axis is in the plane of the page, and is marked by the red arrow. The polar residues are marked with red atoms for oxygen and blue for nitrogen.

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