A Globin Domain in a Neuronal Transmembrane Receptor of Caenorhabditis elegans and Ascaris suum

MOLAR MODELING AND FUNCTIONAL PROPERTIES

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Background: GLB-33 is a putative neuropeptide receptor in C. elegans with a globin domain.

Results: Recombinant globin domain displays a ferric hydroxide-ligated form. When reduced, it can bind CO or O2 and reduce nitrite to NO.

Conclusion: The globin domain may serve as an oxygen sensor or nitrite reductase.

Significance: Oxygen-sensing mechanisms are relevant for neuropeptide receptor binding.

We report the structural and biochemical characterization of GLB-33, a putative neuropeptide receptor that is exclusively expressed in the nervous system of the nematode Caenorhabditis elegans. This unique chimeric protein is composed of a 7-transmembrane domain (7TM), GLB-33 7TM, typical of a G-protein-coupled receptor, and of a globin domain (GD), GLB-33 GD. Comprehensive sequence similarity searches in the genome of the parasitic nematode, Ascaris suum, revealed a chimeric protein that is similar to a Phe-Met-Arg-Phe-amide neuropeptide receptor. The three-dimensional structures of the separate domains of both species and of the full-length proteins were modeled. The 7TM domains of both proteins appeared very similar, but the globin domain of the A. suum receptor surprisingly seemed to lack several helices, suggesting a novel truncated globin fold. The globin domain of C. elegans GLB-33, however, was very similar to a genuine myoglobin-type molecule. Spectroscopic analysis of the recombinant GLB-33 GD showed that the heme is pentacoordinate when ferrous and in the hydroxide-ligated form when ferric, even at neutral pH. Flash-photolysis experiments showed overall fast biphasic CO rebinding kinetics. In its ferrous deoxy form, GLB-33 GD is capable of reversibly binding O2 with a very high affinity and of reducing nitrite to nitric oxide faster than other globins. Collectively, these properties suggest that the globin domain of GLB-33 may serve as a highly sensitive oxygen sensor and/or as a nitrite reductase. Both properties are potentially able to modulate the neuropeptide sensitivity of the neuronal transmembrane receptor.

Nematodes often inhabit low oxygen environments where parameters such as food supply, temperature, and oxygen pressure vary. These fluctuations force the nematode to adapt its behavior to survive (1). Globins can play a role in this adaptation. The globin family is a heterogeneous group of proteins sharing the globin fold and burying a heme cofactor in a hydrophobic pocket, where diatomic gaseous ligands like O2 can bind to the heme iron atom. In silico analysis of the genome of the free-living nematode Caenorhabditis elegans resulted in the identification of a family of 33 globin genes, labeled glb-1 to glb-33, that are all transcribed. The encoded globins, GLB-1 to GLB-33, are localized in a specific subset of cells (2–5). Characterization of some of these globins indicated that they are involved in a wide variety of cellular processes. For instance, GLB-1 may function in regulating facilitated O2 diffusion in a small subset of cells (6). GLB-6 serves a role as redox sensor responding to oxidative stress, either directly or indirectly by the change in O2 levels (7). GLB-5 is an O2 sensor that interacts with the NPR-1 receptor as part of a foraging strategy (8, 9). GLB-26 is a myristoylated redox protein, involved in the defecation cycle under stress conditions (6, 10). The role of all other globins in C. elegans remains unknown. One of the C. elegans globin genes, glb-33, is expressed in neuronal cells and has been predicted to code for a 542-amino acid-long chimeric protein, GLB-33.

The N-terminal domain of GLB-33 (residues 1–372) is composed of 7 hydrophobic α-helices and represents a 7-transmembrane domain (GLB-33 7TM)1 typical of a G-protein-coupled receptor. The globin domain may serve as an oxygen sensor or nitrite reductase.
Characterization of the Globin Domain in a Neuronal Receptor

In Vivo Localization—The translational reporter for glb-33 was constructed using fusion PCR, as described by Hobert (35). The reporter contained the glb-33 target gene and 2.95 kb upstream and 0.653 kb downstream of the target gene to include the endogenous promoter region and 3' UTR gene regulating elements, respectively. The gfp gene was amplified from the pPD95.75 vector and fused at the 3' side of the glb-33 gene, thereby preceding the 3' UTR region. A coding region for a GA linker between GLB-33 and GFP was also included. The structure and characteristics of the primers used were as follows: 5'-TAGACAGTTGCA-CATCCT-3' (forward primer of the promoter of glb-33); 5'-GGGG-TACACCTTCAAAAAAC-3' (nested forward primer of the glb-33); 5'-TCCAGCGCTTCAGCTCCTGG-ACATTGCACAAATGTG-3' (reverse primer of the glb-33 gene); 5'-GAGTGGAATCAATATACAAATGACTTTTCTG-CTAGATCTCGC-3' (forward primer of the 3' UTR region of glb-33); 5'-TGTTCCTTCCCGATATGGTAN-3' (reverse primer of the 3' UTR region of glb-33); 5'-GATCCGGTACAATCTTC-CTC-3' (nested reverse primer of the 3' UTR region of glb-33); and 5'-GGAGCTGTTGACGGGCTGACCCGTGCTTGGACAAAGAGGAAAC-3' (forward primer of gfp), and 5'-TTTTGTATGTCCATCCATGC-3' (reverse primer of gfp).

In addition, the unc-119 gene, including a 2.189-kb upstream region and a 1.228-kb downstream region of the unc-119 isoform, was amplified with forward primer 5'-TCAGTAAAAAAGATGAAAT-3' and reverse primer 5'-GAATTTTAA-CATACCTTC-3'. The PCR product was used as a co-injection marker and rescued the locomotion defect of the unc-119(ed3) strain, the strain used for micro-injection. The final PCR products were injected into the gonads of young adult hermaphrodites using an Axiovert 135 (Zeiss) microscope and FemtoJet microinjection system (Eppendorf), at a concentration of 50 ng/µl for the glb-33 reporter and 20 ng/µl for the unc-119 gene. Transformed lines were analyzed using a Nikon Eclipse TE2000–5 confocal microscope.

In Silico Identification of Orthologs—The protein sequence of GLB-33 (accession number AAK68603) was used to search the nonredundant database of NCBI for orthologs using the protein-protein Basic Local Alignment Tool (BLAST) algorithm (BLASTP) (36, 37). Additional similarity searches were performed on the A. suum CDS gene set (38) using the NemaBLAST algorithm available at Nematode.net version 3.0 (39). GLB-33 orthologs were also identified using the BLASTP algorithm implemented in WormBase (40). All significant hits were manually aligned with GLB-33 using GeneDoc (41), employing the procedure used earlier (42, 43) based on the myoglobin fold (33) and the pattern of predominantly hydrophobic residues at 37 conserved solvent-inaccessible positions.

Cloning, Expression, and Purification of Recombinant GLB-33 GD and A. suum FMRF GD—C. elegans worms were grown as described previously (2). Young adult worms were collected, and total RNA and cDNA were prepared as described (6). To confirm the presence of a globin domain in the A. suum FMRF-amide receptor (accession number ADY45777.1), the cDNA coding for this globin domain was amplified and sequenced. A. suum cDNA (1365 bp) was kindly provided by Peter Geldhof and Edwin Claerebout (University of Ghent, Belgium) and was prepared as follows. Total RNA from L3 larvae

EXPERIMENTAL PROCEDURES

In Vitro Localization—The cDNA coding for GLB-33 (glb-33) is 1629 bp long and was cloned in the pEGFP-N1-vector with C-terminal GFP tag (Clontech) using BglII and BamHI restriction enzymes (Biolabs, Westburg, Netherlands) and T4 DNA ligase (Novagen). 1 µg of the expression construct was transfected into human neuroblastoma SH-SY5Y cells (ATCC® CRL-2266) with 3 µl of Lipofectamine 2000 (Invitrogen). Transfected, and investigated its spectroscopic characteristics, kinetics, and equilibrium of the binding process used earlier (42, 43) based on the myoglobin fold (33) and the pattern of predominantly hydrophobic residues at 37 conserved solvent-inaccessible positions.

Cloning, Expression, and Purification of Recombinant GLB-33 GD and A. suum FMRF GD—C. elegans worms were grown as described previously (2). Young adult worms were collected, and total RNA and cDNA were prepared as described (6). To confirm the presence of a globin domain in the A. suum FMRF-amide receptor (accession number ADY45777.1), the cDNA coding for this globin domain was amplified and sequenced. A. suum cDNA (1365 bp) was kindly provided by Peter Geldhof and Edwin Claerebout (University of Ghent, Belgium) and was prepared as follows. Total RNA from L3 larvae...
was extracted with TRIzol reagent. After treatment with DNase (Invitrogen), cDNA was generated from 1 μg of total RNA using the cDNA synthesis kit (iScript™, Bio-Rad). PCR was then performed with the gene-specific forward and reverse primers 5'-ATGGGCGCTAAAACACGGCCAC-3' and 5'-TGTTTCTTAAAGAAAATCTGTC-3', respectively. The PCR was performed in a final volume of 25 μl under the following conditions: 1× buffer, 1.5 mM MgCl₂ (Invitrogen), 0.2 mM of each dNTP, 1 μM of each primer, and 1 unit of Platinum® TaqPCR DNA polymerase (Invitrogen). The PCR was started at 4 min at 95 °C, which was followed by 35 cycles consisting of 30 s at 95 °C, 30 s at 55 °C, and 60 s at 72 °C. A final step of 10 min at 72 °C was added to the last cycle. The PCR product was analyzed on a 1.5% agarose gel.

The cDNAs coding for the GLB-33 GD (glb-33 gd, bp 1120 to 1365) were cloned in the pET23a vector with C-terminal His tag (Novagen) using Ndel and Xhol restriction enzymes (Biolabs). The sequences of the forward and the reverse primers that were used to amplify the cDNAs were as follows: 5'-CATAT-GGTTTCTCGAGAAGCTTTCGGCC-3' and 5'-GGCGTCGAG-TGCGATTTGGAGCAATTTGTGTC-3' for glb-33 gd and 5'-GGAGATCCATATGTCGATCTGAGCGCAGCAGGAATGATGATAGATT-CCGCTGCAGGTTTCTTAAAGAAAATCTGTC-3' for A. suum fmrf gd, respectively. Ligation of the cDNAs in the pET23a vector was performed using T4 DNA ligase (Novagen). The ligated product was sequenced on both strands using BigDye terminator chemistry and an ABI 377 sequencer (Applied Biosystems). The expression in BL21(DE3)pLysS was verified by UV-visible spectroscopy.

Continuous wave EPR spectra were recorded on a Bruker X-band spectrometer equipped with a helium cryostat (Oxford Inc.). All measurements were performed on a Bruker EleXsys instrument equipped with a helium cryostat (Oxford Inc.). The continuous wave EPR spectra were obtained with a modulation frequency of 100 kHz and a modulation amplitude of 0.5 millitesla. The electron spin echo (ESE)-detected EPR experiments were performed using the π/2-π-π-echo sequence with $t_{π/2} = 16$ ns, $t_{π} = 32$ ns; $τ$ was varied in steps of 8 ns from 88 to 3280 ns. The spectra recorded at the different $τ$ values were subsequently summed.

Characterization of the Globin Domain in a Neuronal Receptor—CO Association Kinetics by Flash Photolysis—CO-bound GLB-33 GD samples were prepared as shown in Equation 1,

$$k_{on,CO} \cdot [CO] + k_{off} \cdot [CO] = k_{on,CO} \cdot [CO] + k_{off} \cdot [CO] = 0$$

for the geminate recombination of CO to 400 ms for the bimolecular rebinding of the ligand at 412 nm. Exponential decays from the consecutive time scales were joined together to give the complete ligand-rebinding curve. All data were analyzed using the Matlab program (The Math Works Inc.). The rate of geminate rebinding ($k_{gen}$) was obtained by fitting a single exponential curve through the data points collected in the first 2000 ns after photodissociation shown in Equation 1,

$$\Delta OD_t = \Delta OD_{gen} \cdot \exp(-k_{gen} \cdot t) + \Delta OD_{obs}$$

After logarithmic resampling of the data points, pseudo-first order fast observed rebinding rate constants ($k_{obs,f}$) and slowly observed rebinding rate constants ($k_{obs,s}$) were determined by least square fitting of the decays with bi-exponential Equation 2,

$$\Delta OD_t = \Delta OD_{f} \cdot \exp(-k_{obs,f} \cdot t) + \Delta OD_{s} \cdot \exp(-k_{obs,s} \cdot t)$$

Fast and slow CO-rebinding association rate constants ($k_{on,f}$ and $k_{on,s}$, respectively) were calculated from the dependence of $k_{obs,f}$ and $k_{obs,s}$ on the CO concentration according to Equations 3 and 4 (45).
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Three-dimensional Models of GLB-33 and A. suum FMRF-GD—The three-dimensional structures of GLB-33 GD (residue 331–455), GLB-33 TM (residue 1–373), A. suum FMRF GD (residue 331–455), and A. suum FMRF 7TM (residue 1–330) were predicted using the I-TASSER server (52, 53) due 1–330) were predicted using the I-TASSER server (52, 53).

The autoxidation rate of GLB-33 GD (\( \sim 10 \mu M \) heme) was measured in air in 100 mM sodium acetate buffer, pH 4, and in 100 mM sodium phosphate, 0.5 mM EDTA, pH 7.4, at 25 °C and 37 °C, after previous reduction of the ferric protein with dithionite and desalting.

The decrease in absorbance at 577 nm was monitored spectrophotometrically every minute for the first 90 min and every 5 min for the next 60 min, using a Cary-5 UV-visible NIR spectrophotometer (Varian) in experiments performed at pH 4.0, and every 50 min using an HP 8543 UV-visible diode array spectrophotometer in experiments performed at pH 7.4. Absorbance traces were fitted according to a single exponential function.

Sequence Analysis of A. suum FMRF GD—Alignment of the protein sequences of GLB-33 and A. suum FMRF (24% identical and 44% similar) showed that the physicochemical properties of the majority of the amino acids are conserved (Fig. 1). In the alignment, the positions of the transmembrane helices and \( \alpha \)-helices are indicated as predicted by the three-dimensional model of these proteins. It is clear that the amino acids building up the transmembrane helices are very similar, differing only by a few amino acids, thus resulting in transmembrane regions of comparable length (Fig. 1A). However, in the globin-like domain of A. suum FMRF, the lack of the \( \alpha \)- and \( \beta \)-helices and similar fold, as it uses the alignment of hidden Markov models via HHsearch to significantly improve the accuracy and the detection rate (54). The quality of the models was evaluated with Procheck (55) and Verify3D (56). ProSA-web (57) was utilized to examine the energy of residue-residue interaction using a distance-based pair potential approach in which the energy is transformed to a Z score where residues with a negative score indicate reasonable side chain interactions. The overlays between the predicted models and the figures were made using the PyMOL Molecular Graphics System (58). The full-length predictions were generated with a combination of manual approach and docking techniques. First, the modeled N-terminal segment of the globin domains was approached to the C-terminal segment of the respective modeled 7TM domains. Second, the RosettaDock program (59) was launched to optimize the interface contacts of the two domains. A dimer of two GLB-33 GD models was also predicted using the known dimeric GLB-1 crystal structure (PDB code 2WTH) as a template. The interface of the dimer was optimized with the RosettaDock program.

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\begin{align*}
\text{O}_2 \text{ Equilibria—} & \text{O}_2 \text{ equilibrium curves of GLB-33 GD were determined at 25 °C using a modified thin layer diffusion chamber technique described elsewhere (46–48). Prior to measurements, the sample was reduced with dithionite and desalted, as described above. A 4-μL sample (} & \sim 100 \mu M \text{ GLB-33 GD in 100 mM sodium phosphate, 0.5 mM EDTA, pH 7.4) was equilibrated with water-saturated gas mixtures of varying O}_2 \text{ tension (PO}_2 \text{) created by Wösthoff gas mixing pumps. Reversible absorbance changes following changes in PO}_2 \text{ were recorded at 436 nm. Values for } P_{50} \text{ (PO}_2 \text{ at half-saturation) and } n_{50} \text{ (cooperativity coefficient) were calculated from the zero intercept and slope of Hill plots, log}(Y)/(1 – Y) \text{ versus logPO}_2, \text{ respectively, where } Y \text{ is the fractional saturation.}
\end{align*}
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Nitrite Reductase Activity—To study the reaction between deoxyglutinated GLB-33 GD and nitrite, a stock solution of nitrite (\( \sim 20 \mu M \) in 100 mM sodium phosphate, 0.5 mM EDTA, pH 7.4) and a sample containing GLB-33 GD (\( \sim 10 \mu M \) heme in 100 mM sodium phosphate, 0.5 mM EDTA, pH 7.4) were deoxyglutinated by equilibrating with \( N_2 \) gas for \( \sim 30 \) min. Nitrite solutions were freshly made every day, and nitrite concentrations were checked by the Griess method (49). An \( \sim 2–3\)-fold molar excess of a previously degassed dithionite solution was added (\( \sim 20–30 \mu M \) final concentration) to the \( N_2 \)-equilibrated GLB-33 GD sample, and absorbance spectra were taken to verify full heme deoxyglutination. The reaction between deoxyGLB-33 GD and nitrite (range 0.1–0.3 mM) was measured at 25 °C under pseudo-first order conditions in the presence of dithionite, as described previously (50, 51). Observed rates were obtained by fitting absorbance traces at 579 nm to a single exponential function, and the apparent second order rate constant was obtained from the slope of a linear plot of observed rates as a function of nitrite concentration.

Autoxidation Rate—The autoxidation rate of GLB-33 GD (\( \sim 10 \mu M \) heme) was measured in air in 100 mM sodium acetate buffer, pH 4, and in 100 mM sodium phosphate, 0.5 mM EDTA, pH 7.4, at 25 and 37 °C, after previous reduction of the ferric protein with dithionite and desalting.

The increase in absorbance at 577 nm was monitored spectrophotometrically every minute for the first 90 min and every 5 min for the next 60 min, using a Cary-5 UV-visible NIR spectrophotometer (Varian) in experiments performed at pH 4.0, and every 50 min using an HP 8543 UV-visible diode array spectrophotometer in experiments performed at pH 7.4. Absorbance traces were fitted according to a single exponential function.

Three-dimensional Modeling of GLB-33 and A. suum FMRF-amide Receptors—The three-dimensional structures of GLB-33 GD (residue 331–455), GLB-33 TM (residue 1–373), A. suum FMRF GD (residue 331–455), and A. suum FMRF 7TM (residue 1–330) were predicted using the I-TASSER server (52, 53) without structural restraints. The I-TASSER server builds three-dimensional models using multiple-threading alignments by LOMETS and iterative TASSER assembly simulations. The Phyre2 program was used to identify proteins with a
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the extended C-helix is striking when aligned to GLB-33 GD (Fig. 1, A and B), resulting in a possible modified truncated globin domain. This globin domain is preceded by a helical linker region that connects it to the 7TM domain (Fig. 1 A).

Therefore, we considered the possibility that the cDNA sequence of the globin domain of the A. suum FMRF reported in the database was incorrectly predicted, requiring a correction of the open reading frame. We therefore amplified the cDNA of A. suum FMRF and determined its sequence. Remarkably, the amplified cDNA was identical to the predicted one, supporting a correct protein sequence with an unusual tertiary structure.

In Vivo and in Vitro Localization of GLB-33—For the localization of GLB-33 in vivo, a translational reporter for the glb-33 gene was constructed including the endogenous promoter region and 3'UTR elements. The expression pattern of this reporter was in line with the results of a previously used transcriptional reporter (4) and indicated that GLB-33 was exclusively expressed in the nervous system (Fig. 2 A). The GLB-33 protein was present in a large number of neurons in the head and tail region, the nerve ring, the ventral and dorsal nerve cord, and several lateral nerve cords (Fig. 2, B and D). However, despite its wide expression in the nervous system, GLB-33 did not seem to be expressed in any of the amphid, cephalic, labial, or phasmid sensory neurons; the typical expression pattern for these neurons was not observed nor was any overlap seen between GFP expression and 1,1'-dioctadecyl-3,3,3',3'-tetramethylindocarbocyanine perchlorate (DiI) staining, a
compound that selectively stained several amphid and phasmid neurons (data not shown). This indicates that GLB-33 is expressed in motor neurons or interneurons, which are involved in locomotor behavior or information processing, and not in neurons that sense environmental cues. Finally, the expression pattern of the reporter was in line with the membrane-bound localization of the receptor (Fig. 2). The membrane localization of GLB-33 was confirmed by the transfection of \textit{glb-33-gfp} in human neuroblastoma SH-SY5Y cells (Fig. 2E).

**Spectral Characterization of GLB-33 GD and \textit{A. suum} FMRF GD**

**UV-visible Spectra**—The UV-visible absorption spectra of GLB-33 GD in the as-expressed, the dithionite-reduced (ferrous deoxy), the ferrous CO-bound, and the ferrous oxy form, prepared at pH 8.5 and the aquo-met form, prepared at pH 4, are shown in Fig. 3. The ferrous deoxy form has the Soret band at 433 nm and a Q band maximum at 558 nm. These maxima are typical for pentacoordinate high spin ferrous deoxyheme proteins (45). The absorption spectrum of the CO form of GLB-33 GD is similar to those of sperm whale myoglobin (swMb), with peaks at 420, 539, and 568 nm (45). After reduction with dithionite and desalting, the ferrous oxy form has peaks at 413, 543, and 581 nm (Fig. 3A). Although the absorption spectrum of as-expressed GLB-33 GD (pH 6–8.5) is apparently very similar to that of a ferrous oxyglobin, with the Soret band located at 413 nm and the Q bands at 544 and 577 nm, these maxima are also similar to those of the ferric hydroxide-ligated form of swMb (61) and to those of low spin heme oxygenase at alkaline pH (62). EPR (see below) confirms that the ligated state of the as-expressed GLB-33 GD can be assigned to the hydroxide-ligated ferric form. Note that in the as-purified samples at pH 8.5, there was still a small fraction of ferrous oxy form present capable of reversible O₂ binding.

The UV-visible spectra of the protein remained practically unchanged during the purification process, after several freeze-thaw cycles and after prolonged exposures to air, indicating that the hydroxide-ligated ferric derivative is very stable. Dialysis of GLB-33 GD against 10 mM sodium acetate, pH 4, resulted in the aquo-met form (Fig. 3B), with absorption maxima comparable with those of hexacoordinate high spin heme oxygenase (62). This acidic transition was completely reversible, because redialysis to buffers with pH 6–8.5 reconverted GLB-33 GD to the ferric hydroxide-ligated form, thus confirming that the as-expressed protein was not in the ferrous oxy state (data not shown).

In Fig. 3C, the UV-visible absorption spectra of the as-expressed and deoxy-reduced \textit{A. suum} FMRF GD are depicted as prepared in 50 mM Tris-HCl, pH 7.5. The as-expressed protein has the Soret band at 412 nm and Q band maxima at 534 and 558 nm, indicative of ferrous hexacoordinate globins, in contrast with the ferrous pentacoordinate heme of the deoxy-GLB-33 GD. The absorption maximum at 661 nm, typical for free biliverdin (63), in both spectra indicates the presence of denatured protein as the intensity of the peaks remained unchanged after reduction of the heme iron atom (Fig. 3B). Indeed, \textit{A. suum} FMRF GD was very unstable, and extensive precipitation occurred during the whole purification process, as well as during attempts to bind other ligands to the heme iron atom, as could be deduced from the disappearance of the Q band maxima (data not shown). Stable liganded species could not be produced for this reason.

**Resonance Raman Spectra**—Fig. 4A shows the high frequency region of the RR spectra of GLB-33 GD in the as-expressed, the ferrous, and the CO-bound ferrous forms in 50 mM Tris-HCl, pH 8.5. The high frequency region of the RR spectra of heme proteins contains a number of so-called “marker bands” that depend on the oxidation state, the coordination state, and the spin state of the heme iron atom.
RR spectra confirmed that GLB-33 GD is expressed in the hydroxide-ligated ferric form under the conditions used (Fig. 4A(a)), with a first set of marker bands at $\nu_4 = 1376 \text{ cm}^{-1}$, $\nu_3 = 1502 \text{ cm}^{-1}$, and $\nu_2 = 1581 \text{ cm}^{-1}$. The $\nu_2$ line at 1376 cm$^{-1}$ and the $\nu_3$ line at 1503 cm$^{-1}$ are typical of hexacoordinate low spin ferric heme complexes (64). The frequencies of $\nu_3$ and $\nu_2$ are very close to the hydroxide-ligated form of met-globins, which are characterized with $\nu_3$ at ~1505–1507 cm$^{-1}$ and $\nu_2$ at ~1585 cm$^{-1}$ (65–67). A second set of marker bands is located at $\nu_3 = 1476 \text{ cm}^{-1}$ and $\nu_2 = 1558 \text{ cm}^{-1}$, which likely stems from a small fraction of a high spin ferric form (68, 69).

The high frequency region of dithionite-reduced, deoxy-GLB-33 GD (Fig. 4A(b)) has a single set of marker bands at $\nu_4 = 1356 \text{ cm}^{-1}$, $\nu_3 = 1470 \text{ cm}^{-1}$, and $\nu_2 = 1560 \text{ cm}^{-1}$, typical for a pentacoordinate high spin ferrous heme. Two $\nu_4$ bands at 1371 and 1354 cm$^{-1}$ are found in the RR spectra of the CO-bound ferrous protein recorded at low laser power (0.5 milliwatt) (Fig. 4A(c)). These stem from the CO-bound ferrous and photolysed form of the globin, respectively. The latter is confirmed by the fact that the intensity of the $\nu_4$ band at 1354 cm$^{-1}$ increases when the laser power is increased to 100 milliwatts (Fig. 4A(d)). This illustrates that the CO-bound ferrous form is photo-labile. The $\nu_4$ frequency of the CO-bound form is lower than that of the as-expressed ferric form, suggesting that the GLB-33 GD distinguishes –OH from CO as a ligand, as also found for a bacterial heme-based aerotactic transducer (HemAT-Bs(O$_2$) (70)).

FIGURE 3. UV-visible absorption spectra of GLB-33 GD (A) in the as-expressed form pH 8.5 (black) (Soret band, 413 nm; Q bands, 577 and 544), the ferrous deoxy form (blue) (Soret band, 433 nm; Q band, 558 nm), the CO-bound ferrous form (red) (Soret band, 420 nm; Q bands, 568 and 539 nm), and the ferrous oxy form (green) (Soret band, 413 nm; Q bands, 581 and 543 nm) at pH 8.5. B, GLB-33 in the as-expressed form at pH 8.5 (black) and pH 4 (red). The latter gives a spectrum for the aquo-met form (Soret band, 408 nm; Q bands, 503 and 633 nm). C, A. suum FMRF GD in the as-expressed form (black) (Soret band, 412 nm; Q bands, 565 and 534 nm) and the deoxy-ferrous form (red) (Soret band, 424 nm; Q bands, 558 and 534 nm).
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Fig. 4B shows the low frequency region of the RR spectra of the different GLB-33 GD ligated forms. The low frequency region of the RR spectra of heme proteins contains a number of bending modes of the heme peripheral vinyl and propionate substituents. The frequency of these bending modes depends on the strength of the interactions between these side chains of the heme and amino acid residues in the heme pocket of the protein. The assignment of the RR bands is based on the work of Spiro and co-workers on Mb (68). The propionate bending mode, $\delta (C_{pro}-C_{al})$, of as-expressed GLB-33 GD (Fig. 4B (a)) is located at 376 cm$^{-1}$ (broad signal) indicating a strong H-bond interaction between the heme propionates and nearby amino acid residues. A vinyl bending mode is found at 416 cm$^{-1}$. The broad profile of this band indicates the presence of a second vinyl mode at a higher frequency of $\sim$430 cm$^{-1}$. A broader signal is found in the 490–515-cm$^{-1}$ region where the Fe-OH signal is expected (71). For dithionite-reduced deoxy-GLB-33 GD (Fig. 4B (b)), the frequency of the propionate bending mode is located at a higher frequency of 382 cm$^{-1}$, indicating an even stronger interaction between the heme propionates and nearby amino acids. Furthermore, a larger separation between the two vinyl modes was observed. One is located at 414 cm$^{-1}$, and the second mode is located at 432 cm$^{-1}$. These data show significant interactions between the protein moiety and the heme that are dependent on the heme-ligation state.

The Fe-CO stretching mode can be found in the low frequency region of the RR spectra of CO-bound ferrous heme proteins (72). In the current case, a clear determination of the Fe-CO modes is hampered by the high photolability of the CO-ligated GLB-33 GD form (Fig. 4B (c and d)), although a broader signal is seen in the 489–504-cm$^{-1}$ region typical of the Fe-CO modes.

**EPR**—The continuous wave EPR (data not shown) and ESE-detected EPR (Fig. 5a) both confirm the presence of two low spin ferric forms in the as-purified GLB-33 GD sample at pH 8.5. The two low spin components have the following principal $g$ values: $(g_{1},g_{2},g_{3}) = (2.62,2.20,1.815)$ (Fig. 5b, red) and $(g_{1},g_{2},g_{3}) = (2.845,2.12,1.69)$ (Fig. 5b, green). The former $g$ values are similar to those of a low spin Antarctic fish hemoglobin (73) and point to the formation of a hydroxide-ligated ferric heme form (74). The $g$ values are also similar to those observed for alkaline cytochrome $c$ peroxidases (75, 76). The second set of $g$ values is very similar to that of horseradish peroxidases ($g_{1},g_{2},g_{3}$) $\sim$ (2.9,2.1,1.69) showing low spin hydroxide ligation of the ferric heme in alkaline conditions (77, 78). A recent study on *Thermobifida fusca* hemoglobin revealed the presence of two ferric forms with similar EPR parameters as those found here. The component with a $g_{1}$ value around 2.8 was assigned to the hydroxide-ligated ferric complex where the OH group is strongly hydrogen bonded, whereas the form characterized by $g_{1}$ at $\sim$2.6 is ascribed to a coordination mode in which the OH group is not so strongly H-bonded (79). Interestingly, also in *T. fusca* hemoglobin, the hydroxide-ligated complexes were still observable at pH 6.0. No significant contribution of a high spin ferric form (i.e. aquo-met) is found in the continuous wave EPR or ESE-detected EPR of GLB-33 GD at pH 8.5 (data not shown). The EPR spectra thus show that low spin ferric heme complexes are already dominantly formed at lower pH values than in most other globins. The UV-visible absorption and RR spectra are in agreement with this finding. The difference between the two low spin ferric forms most probably stems from a change in the hydroxide stabilization by the surrounding protein matrix.
Characterization of the Globin Domain in a Neuronal Receptor

TABLE 1

Rate constants of ligand binding of *C. elegans* globins compared with rate constants reported for other globins

| Ligand | GLB-33 GD | GLB-1 | GLB-26 | swMb | A. suum Hb |
|--------|-----------|-------|--------|------|------------|
| CO     |           |       |        |      |            |
| $k_{on}$ ($\mu$M$^{-1}$ s$^{-1}$) | 0.90 ± 0.10 | 23    | 0.51 ± 0.06 | 0.21 ± 0.02 |            |
| $k_{off}$ ($\mu$M$^{-1}$ s$^{-1}$) | 0.35 ± 0.01 |       | 0.55 ± 0.08 | 3.0 ± 0.2   |            |
| $k_{gem}$ $(\times 10^5)$ s$^{-1}$ | 9.87 ± 0.01 | 0.43 ± 0.006 | 0.45 ± 0.05 |           |
| $F_{gem}$ | 0.50 ± 0.05 |       |        |      |            |
| O$_2$  |           |       |        |      |            |
| $k_{on}$ ($\mu$M$^{-1}$ s$^{-1}$) | 0.23 | ND    | 14    | 3     |            |
| $k_{off}$ (h$^{-1}$) | 0.5 |       | 15    | 0.013 | 215         |
| $K_{O2}$ ($\mu$M$^{-1}$) | ≥8.55$^a$ | 46$^a$ | ND    | 1.1  | 0.0072      |
| $P_{50}$ (torr) | ≤0.07$^b$ | 0.003$^b$ |       | 1    |             |
| $k_{oxid}$ (h$^{-1}$) pH 4 | 1.81 (25°C) | 15 (37°C) |       |      |             |
| $k_{reduct}$ (h$^{-1}$) pH 7.4 | 0.09 ± 0.02 (25°C) | 0.054 (37°C) 0.005 (25°C)$^c$ |       |      |             |

$^a$ Data are from equilibrium experiments.

$^b$ Data are from kinetic experiments.

$^c$ pH 7.2, from Ref. 83.

Ligand Binding Kinetics of GLB-33 GD

CO Rebinding Kinetics—Flash photolysis was performed at pH 7 as described under “Experimental Procedures.” CO rebinding from within the protein matrix to the heme iron atom is referred to as the [CO]-independent geminate rebinding phase (occurs in the nanosecond time scale after flash photolysis). For GLB-33 GD, this geminate rebinding process occurred with a rate constant of 9.87 (±0.07) $\times 10^7$ s$^{-1}$ (Table 1), about 18 times as fast as the geminate rebinding of CO to swMb ($k_{gem} = 0.55 (±0.08) \times 10^5$ s$^{-1}$, see Table 1). About half of the CO molecules rebind immediately, resulting in a high fraction of geminate rebinding of 0.50 ± 0.05 (Table 1), an indication that the CO molecules cannot easily escape from the protein matrix in comparison with swMb ($F_{gem} = 0.043 ± 0.006$, see Table 1). On longer time scales, a fraction of the gaseous ligands can escape to the bulk solvent and rebind from outside the protein matrix in a [CO]-dependent way. This bimolecular rebinding phase in GLB-33 GD is a biphasic process, characterized by a fast and a slow rebinding rate constant as follows: $k_{on,f} = 0.90 (±0.10) \times 10^6$ M$^{-1}$ s$^{-1}$ and $k_{on,s} = 0.35 (±0.01) \times 10^4$ M$^{-1}$ s$^{-1}$, respectively (Table 1 and Fig. 6). The latter is comparable with the CO rebinding rate of swMb (Table 1).

O$_2$ Equilibria—The apparent $P_{50}$, measured with the diffusion chamber method was 0.07 ± 0.01 torr at 25°C, pH 7.4 (data not shown). However, we cannot exclude that at these very low levels of PO$_2$, the protein within the diffusion chamber was partially oxygenated even when equilibrated with pure nitrogen (i.e. to obtain zero saturation). Therefore, it is possible that the true $P_{50}$ is lower than 0.07 torr. The equilibrium association constant ($K_{O2}$, M$^{-1}$) for O$_2$ derived from the $P_{50}$ value is ≥8.5 $\mu$M$^{-1}$ when assuming an O$_2$ solubility in water of 1.67 $\mu$M torr$^{-1}$ at 25°C (80). A Hill coefficient close to 1 (~0.9) indicated absence of cooperativity (data not shown).

Autoxidation Rate of GLB-33 GD—Autoxidation of oxy-GLB-33 GD at pH 4 (25 and 37°C) and at pH 7.4 (25°C) showed a single exponential process with a rate of 1.81 h$^{-1}$ (25°C) and 15 h$^{-1}$ (37°C) at pH 4, ~278 times faster than native swMb (0.054 h$^{-1}$) at pH 7 and 37°C (81), and ~62 times faster than the swMb H(E7)68E mutant (0.24 h$^{-1}$) at pH 7 and 37°C (82). At pH 7.4 the rate constant is 0.09 ± 0.02 h$^{-1}$. As the absorbance peaks of the oxy- and hydroxide-ligated ferric forms are almost identical (Fig. 3A), the spectral change over time was small. However, the fact that isosbestic points (e.g. 531, 555, 568, and 585 nm at pH 7.4) were maintained over time (data not shown) is consistent with a single oxy to ferric transition at both pH 4.0 and 7.4. For pig myoglobin, a larger increase of the oxidation rate constant (several thousands) (82) was found when going from pH 7 to pH 4 in comparison with GLB33GD. However, this smaller increase can be due to the fact that there is a conversion from the hydroxide-ligated form to the aquo-met form for the latter.

Nitrite Reductase Activity of GLB-33 GD—The nitrite reductase activity was measured in the presence of a slight excess of dithionite. The data show that deoxygenated GLB-33 GD was able to convert nitrite to NO and the nitrite-induced conversion of deoxyheme to NO-bound heme was confirmed spectrophotometrically (Fig. 7A). The visible absorption spectrum changed over time from that of deoxy-GLB-33 GD (Fe$^{2+}$) with a single peak at 558 nm to that of a NO-bound globin (Fe$^{2+}$-NO) with peaks at 545 and 570 nm, indicating that the
reaction followed the general scheme for nitrite reduction by deoxyglobins in the presence of dithionite (51, 84). When plotting the fitted observed rates ($k_{obs}$) as a function of nitrite concentration, the second order rate constant was 36.3 M$^{-1}$ s$^{-1}$ (Fig. 7B), which is ~10-fold higher than that for Mb (2.9 M$^{-1}$ s$^{-1}$) (85).

Three-dimensional Modeling of GLB-33 GD and GLB-33 7TM Domain—The structural model of the N-terminal GLB-33 7TM domain was predicted using the I-TASSER software (52, 53). The best predicted model shows a C score of ~0.30 and an expected TM score of 0.67 ± 0.12. Both these values indicate a satisfactory model with reliable topology. The stereo-chemical quality of the GLB-33 7TM model was analyzed with Procheck. The resulting Ramachandran plot displayed 91.9% of all amino acid side chain angles in allowed regions and 8.1% in disallowed regions. Verify3D analysis showed about 50% of the residues with a score over 0.15, and ProSA-web evaluation confirmed this result with an overall calculated Z score of −2.52. Phyre2 identified the human $\beta_2$-adrenergic G-protein-coupled receptor ($\beta_2$-AR, PDB code 2RH1 (86)) as the most similar protein of known structure, with 41% similarity with the query sequence, and which was chosen as reference structure for the comparison. An overlay of the model with the $\beta_2$-AR structure was generated using PyMOL (Fig. 8A). A root mean square deviation (r.m.s.d.) value of 1.80 Å was calculated over 243 Ca atom pairs, suggesting the presence of several differences. The position of every helix of the modeled GLB-33 7TM is very well superimposed on the $\beta_2$-AR structure throughout the whole length, with exceptions in correspondence of numerous loops as follows: N- and C-terminal extremities as well as the loop connecting helix 5 to 6 are not present in the PDB file of $\beta_2$-AR (evidence of strongly flexible regions); the GLB-33 7TM model shows a 17-residue longer loop connecting helix 3 to 4 compared with the reference structure; and the amino acid stretch that connects helix 4 to 5 presents a short ordered helix in the $\beta_2$-AR structure, although it was predicted as disordered in our model. The most significant distances in the Ca positions of the $\alpha$-helices are between a minimum of 1.8 Å (N-terminal of helix 5) and a maximum of 6.5 Å (N-terminal of helix 1).

Similarly, a prediction model was generated for GLB-33 GD. The C score (~0.62) and the expected TM score (0.63 ± 0.13) indicate the good quality of the best model predicted. Procheck evaluation indicated that the geometry of 97.6% of the residues is in allowed regions and as little as 2.4% is in disallowed regions. Verify3D showed that all the residues have a score higher than 0.2, and ProSA-web results confirm this finding through a negative Z score for every residue and a calculated Z score of ~7.44 for the global molecule. All these data indicate a reliable model. For comparison, an overlay of the predicted model on the known high resolution three-dimensional structure of GLB-1 (PDB code 2WTG (6)) was made (Fig. 8B). The r.m.s.d. of the overlay resulted in 2.30 Å calculated on 134 Ca atom pairs. The 3-over-3 globin fold is conserved in the GLB-33 GD model; nevertheless, the different orientation of the modeled helices compared with the reference structure causes a global noncorrespondence of the Ca backbone. A closer look at the amino acid residues in the distal site of the GLB-33 GD model highlights its exceptional hydrophobic composition as follows: Leu-41(CD3), Ile-69(E7), Arg-72(E10), and Ile-73(E11) (Fig. 8C). This highly apolar environment leaves Arg-72(E10) as the only residue available for ligand stabilization. Likewise, C. elegans GLB-1 displays a majority of hydrophobic residues in its distal pocket (Phe-34, Phe-38, Phe-49, Phe-66, Ile-73, and Phe-114), but O$_2$ is directly stabilized by an unusual combination of the distal residues Tyr-35(B10) and Gln-69(E7) (6).

Until now, no three-dimensional structure of a globin associated with a transmembrane domain has been solved. As a consequence, homology modeling of the full-length protein was impossible. To visualize the full structure of GLB-33, a manual approach of the assembly of the GLB-33 GD model to the GLB-33 7TM model was attempted. The N terminus of the globin domain was located manually in the proximity of the C terminus of the 7TM domain. Subsequently, the docking program RosettaDock was used to optimize the interface interactions between the two models. The result is shown in Fig. 8D. Despite the limitation of this approach, the docking shows a
wide interaction surface between the two domains and a possible connection between the two terminal loops of the predicted models. Indeed, the interaction surface involves the N terminus of the A-helix, the E-F corner (i.e. C-terminal E-helix and N-terminal F-helix), 70% in length of the H-helix, and the C-terminal loop of the GD domain. Thus, these data suggest that any conformational change that occurs in the GD can be readily transmitted to the 7TM domain.

Three-dimensional modeling of A. suum FMRF 7TM and A. suum FMRF GD—Similarly to GLB-33, the full-length FMRF-amide receptor of A. suum was modeled separating first the N-terminal A. suum FMRF 7TM from the C-terminal A. suum FMRF GD and second rejoining manually the two predicted domains. The A. suum FMRF 7TM model predicted by the I-TASSER software (52, 53) shows a C score and an expected TM score (0.41 and 0.77 ± 0.10, respectively) that indicates a good quality model. The geometry analysis performed with Procheck resulted in 94.5% of the residues in allowed regions and in 5.5% in disallowed regions. The Verify3D program highlighted that about 38% of the residues shows a score over 0.15 and the Z score calculated with ProSA-web of −0.49 confirmed this result. As for GLB-33–7TM, Phyre2 identified the human β2-AR (86) as the most similar protein of known structure with 35% similarity. An overlay of the A. suum FMRF 7TM model with the GLB-33 7TM model was generated using PyMOL (Fig. 9A). Over 282 Ca atoms, the r.m.s.d. of 1.18 Å was calculated, suggesting a low level of variety. Indeed, the two models superimpose very well, except for longer N- and C-terminal loops in the GLB-33 7TM model, and for the longer interhelical loop between helices 3 and 4 already noticed in its comparison with the β2-AR structure, which is absent in the prediction of A. suum FMRF 7TM.
Strikingly, the three-dimensional model for the putative C-terminal A. suum FMRF GD indicated a globin fold not referable to any known globin family. It can be recognized as a classical 3-over-3 H/9251-helical globin fold, but lacking the A and D helices and exhibiting a longer C-helix (Fig. 9 B). For this model, we obtained a C score of 0.90 and an expected TM score of 0.56 0.14 with the I-TASSER software. The geometry quality assessment performed with Procheck indicated that 98.4% of the residues were in allowed regions and 1.6% were in disallowed regions. Verify3D and ProSA-web display positive concordant results (about 93% the residues have a score higher than 0.2, a negative Z score for every residue and a calculated Z score of 5.2 for the global molecule, respectively). All these data confirm the good geometry of this model. The Phyre2 similarity search identified the globin domain of GLB-6 from C. elegans (PDB code 3MVC (7)) as the most similar structure with 42% of similarity. Because A. suum FMRF GD lacks several helices, we modeled its structure with a variety of 2/2 truncated hemoglobin structures as a template to exclude bias in the modeling program. These tests did not result in a good model however, and suggested that the a-helical fold of A. suum FMRF GD is more related to the 3/3 classical globin fold than to the 2/2 truncated hemoglobins, even though some deviations can be detected (data not shown). Superimposing the models of A. suum FMRF GD and of GLB-33 GD (Fig. 9B), we obtained the r.m.s.d. value of 2.57 Å over 88 Ca atom pairs. Obviously, this comparison emphasizes the lack of the A-, B-, and D-helices in A. suum FMRF GD. The shorter F-helix of GLB-33 GD is also noticeable, as well as the similar open conformation of the E-helix in both predicted models. A detailed look at the amino acid composition of the heme pocket reveals that it is more polar than the heme pocket of GLB-33 GD, with Phe-10(B10), Glu-38(E7), Cys-41(E10), and Ile-42(E11) (Fig. 9C).

Finally, the full-length appearance of A. suum FMRF was investigated using the model (Fig. 9D). The two docked domains display a discrete interaction area and the possibility of a covalent connection between the two termini of the predicted models. The heme pocket of the docked A. suum FMRF GD is even more exposed to the solvent than in GLB-33 GD (Fig. 9D), as shown in Fig. 9D. Obviously, this is also due to the peculiar truncated globin fold of A. suum FMRF GD.

DISCUSSION

The similarity in primary and tertiary structures between the C. elegans GLB-33 and the A. suum FMRF-amide receptor, as indicated by modeling analysis (Fig. 9), suggests that they could
serve a similar role in vivo. The modeled N-terminal extracellular loops create a structural pocket where FLPs could bind thus leading to a conformational change, resembling that of other FMRF-amide receptors (Fig. 8A). Nothing is known, however, about the process regulated by these chimeric proteins, but some indications can be found by studying the in vivo transcription pattern. Analysis of transcriptome data of A. suum during development indicates that the FMRF-amide receptor is predominantly expressed in the L3 larval stage, and peaks when the larvae have just hatched from the egg, but it is unclear so far what role the FMRF-amide receptor could play in this developmental stage. In contrast, the C. elegans GLB-33 is expressed throughout all life stages of the nematode (3). Following the in vivo expression pattern of GLB-33, we detected this protein predominantly in inter- and motoneurons, and it might therefore be involved in movement following processing sensory cues (Fig. 2).

To explore the function of the globin domains of GLB-33 and of the A. suum FMRF-amide receptor, recombinant GLB-33 GD and A. suum FMRF GD were prepared and their biochemical properties investigated. It has to be noted that in A. suum the recombinant globin domain was very unstable, probably due to the lack of several helices as compared with genuine globins and the lack of the FMRF-amide receptor domain that might compensate this truncation. In the absence of various gaseous ligands, the globin domain of GLB33 showed an optical spectrum that is characteristic of a pentacoordinate ferrous globin, whereas the globin domain of A. suum is hexacoordinate. Raman and EPR spectra of C. elegans GLB-33 GD show tight interactions between the heme group or its ligand and the surrounding protein moiety (Figs. 4B and 5). Because changes in the ligation state of the heme iron atom are accompanied by conformational changes in the globin core, this suggests that the heme ligation and/or redox state of the globin domain could influence the conformational state of the coupled receptor domain. In this way, the globin domain could modulate the sensitivity of the effector domain for the binding of neuromodulators. Likewise, the binding of a neuromodulator to the receptor domain could introduce a conformational change in the globin domain, thus altering the ligand-binding properties of the latter. More arguments for the modulation of ligand affinity can be found in our modeling data. Docking the model of GLB-33 GD to that of GLB-33 7TM showed that the opening to the heme pocket is directed to the outer environment (Fig. 8D), suggesting that in the chimeric protein ligand binding and dissociation are still possible.

The reactivity of a globin domain toward ligands depends mostly on the amino acid residues in the heme pocket that can stabilize the bound ligand. The distal B10, E7, E10, and E11 residues are known to be involved in this stabilization in several globins (87–89) and are occupied by a different set of residues in the two globin domains investigated here (Fig. 1B): Ala-31, Ile-69, Arg-71, and Ile-73 in GLB-33 GD, and Phe-10, Glu-38, Cys-41, and Ile-42 in A. suum FMRF GD, respectively. Interestingly, neither globin domain contains the distal His(E7). Few examples of globins that contain Ile as the E7 distal residue, as GLB-33 GD, are known, including the truncated Hb of the cold adapted bacterium Pseudomonas haloplanktis TAC125 (Ph/2/2HbO), where the B10, E7, E10, and E11 positions are occupied by Tyr, Ile, Val, and Phe, respectively (90). The electronic absorption spectra of Ph/2/2HbO at physiological pH and low temperature (12 K) are very similar to those of GLB-33 GD at the same pH at room temperature, with absorption maxima in the Q band region at 535 and 570 nm, indicating a hexacoordinate low spin ferric species with a hydroxide group as the 6th ligand. In the case of Ph/2/2HbO, this ligand is donated to the heme iron by the Tyr(B10) residue (90). At room temperature, the absorption spectrum of Ph/2/2HbO indicates the additional presence of a hexacoordinate high spin His–Fe–H2O form (90) and is similar to the spectrum of GLB-33 GD at pH 4 (Fig. 3B). Also T. fuscum hemoglobin, with Tyr, Ala, Arg, and Leu on positions B10, E7, E10, and E11, exhibits hydroxide-ligated forms, which are in equilibrium with an aquo-met form at pH 6 (79). In the latter protein, EPR contributions of hydroxide-ligated heme forms with similar principal g values as for the GLB-33 GD case were observed, with the exception that the g1 ~2.6 contribution was only a minority species. In the case of GLB-33 GD, both components contribute approximately equal to the overall EPR spectrum. This is in line with the amino acid residues in the distal pocket of the two proteins. In T. fuscum hemoglobin, the hydroxide group bound to the ferric heme is stabilized by Tyr(B10), Tyr(CD1), and Trp(G8) in agreement with the dominance of the g1 ~2.8 contribution in the EPR spectrum indicative of strong H-bonding to OH−. In contrast, in GLB-33 GD, the B10 position is occupied by an Ala residue unable to stabilize the hydroxide ligand. The only polar side chain in the largely apolar heme pocket is Arg(E10). This explains the larger fraction in which the heme-bound hydroxide is more weakly H-bonded to the protein than in the T. fuscum hemoglobin, as shown by the larger EPR g1 ~2.6 contribution (Fig. 5). Because no amino acid residues are present in the distal pocket of GLB-33 GD that can donate a hydroxide group to the ferric heme iron (Fig. 8C), it is most likely that the hydroxide ligand originates from a water molecule as is also the case for T. fuscum hemoglobin. This hypothesis is supported by the observation that the conversion between the high spin aquo-met species and the low spin hydroxide-ligated form is completely reversible upon pH change through dialysis (data not shown).

Interestingly, a similar heme environment is found in the distal pocket of the Per-Arnt-Sim (PAS) domain of the O2 sensor Fix-L of Bradyrhizobium japonicum (B. japonicum Fix-L). Although the structures of the globin domain and of the PAS domain differ significantly, they both bind a heme cofactor, and some comparisons can be made. In Bradyrhizobium japonicum Fix-L, ligand interaction is possible through the guanidinium group of G122Arg (91). This residue has been shown to assist in the sensing of O2. This is not accomplished by stabilizing the oxy form however; instead, the side chain is involved in triggering a conformational switch upon binding of regulatory ligands (92). In GLB-33 GD, the Arg(E10) residue could be involved in a similar mechanism. It appears plausible that when O2 binds with high affinity to the ferrous heme, it will induce a confor-

7 Peter Geldhof (University of Ghent) and Robin Gasser (University of Melbourne), personal communication.
mational switch of the Arg-72(E10) residue upon formation of the oxy derivative, which may then act as a trigger to transduce the signal to the transmembrane receptor domain. The observation of two ferric hydroxide-ligated forms observed in EPR spectra (Fig. 5) suggests the existence of two different protein conformations with potentially different roles in signal activation or deactivation.

Another functionally important property of ferrous GLB-33 GD is its higher reactivity with nitrite, compared with other globins. This may be due to the positively charged Arg-72(E10) residue, which may help in facilitating electron transfer from the heme to the bound nitrite. The relatively fast conversion of nitrite to NO mediated by the ferrous protein could also be involved in efficient neuronal signaling and affect local activities of nearby neurons. Moreover, it has been shown that NO mediates the inhibitory effects of an FMRF-amide-like neuropeptide in A. suum, linking NO with neuropeptide signaling (93). Clearly, the effects of such reaction on the effector domain of the full-length protein remain to be investigated, particularly because C. elegans does not express nitric-oxide synthase enzymes and could then in principle be more reliant on nitrite reductase activities for NO signaling.

The covalent structural coupling of a globin domain and a 7TM domain is unique and has not been observed before. However, a functional noncovalent interaction has been shown for the O₂ sensor GLB-5, one of the globins encoded in the genome of C. elegans, and NPR-1, an FMRF-amide receptor activated by the binding of the neuropeptides FLP-18 and FLP-21 (8, 9, 94). GLB-5 modulates the same O₂-sensing neurons as NPR-1 (AQ, PQ, and URX), which also expresses GCY-35 and GCY-36, subunits of soluble guanylate cyclases that also act as O₂ sensors (9, 95–100). It was proposed that GLB-5 acts as a signaling molecule, inhibiting neuronal activation when the O₂ concentration drops below 21% (9). Interestingly, soluble guanylate cyclases can also be activated by NO, emphasizing a possible similar mechanism between GLB-33 and GLB-5 (83). Because of its very high affinity for O₂, GLB-33 may function as a highly sensitive O₂ sensor that would modulate neuronal sensitivity to neuropeptides and allow C. elegans to move toward nutrient-rich substrates in the hypoxic soil. Furthermore, the ability of GLB-33 GD to generate NO from nitrite (largely available in the soil) under hypoxic conditions would then add another level of complexity to this response.

CONCLUSION

Overall, we can conclude that GLB-33 is composed of a 7TM domain, similar to FMRF-amide receptor domains, covalently coupled to a globin domain that is able to bind external ligands at the heme group. Our modeling data show that the conformational switch of GLB-33 7TM upon ligand binding, likely a neuropeptide ligand, can result in a conformational change of the globin domain and conversely that reactions in the heme pocket may modulate the affinity of neuropeptides for the 7TM domain.

Characterization of GLB-33 GD showed that it possesses characteristics typical of an O₂ sensor when present in the ferrous and ferric forms. Furthermore, the ferric heme occurs as a combination of two hydroxide-ligated ferric forms under physiological conditions. In addition, because nitrite reductase activity is fast compared with that of other globins, GLB-33 GD could also be involved in local NO production from available nitrite when in the ferrous unliganded state, provided that close targets of NO bioactivity and heme-reducing pathways are present. We identify Arg-72(E10) as the most likely trigger for a conformational switch that could be transmitted to the 7TM domain and as such activate yet unknown neuronal signaling pathways.

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