Ligand uptake in *Mycobacterium tuberculosis* truncated hemoglobins is controlled by both internal tunnels and active site water molecules [version 2; peer review: 2 approved, 1 approved with reservations]

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**Abstract**

*Mycobacterium tuberculosis*, the causative agent of human tuberculosis, has two proteins belonging to the truncated hemoglobin (trHb) family. Mt-trHBN presents well-defined internal hydrophobic tunnels that allow $O_2$ and *NO* to migrate easily from the solvent to the active site, whereas Mt-trHBØ possesses tunnels interrupted by a few bulky residues, particularly a tryptophan at position G8. Differential ligand migration rates allow Mt-trHBN to detoxify *NO*, a crucial step for pathogen survival once under attack by the immune system, much more efficiently than Mt-trHBØ. In order to investigate the differences between these proteins, we performed experimental kinetic measurements, *NO* decomposition, as well as molecular dynamics simulations of the wild type Mt-trHBN and two mutants, VG8F and VG8W. These mutations affect both the tunnels accessibility as well as the affinity of distal site water molecules, thus modifying the ligand access to the iron. We found that a single mutation allows Mt-trHBN to acquire ligand migration rates comparable to those
observed for Mt-trHbO, confirming that ligand migration is regulated by the internal tunnel architecture as well as by water molecules stabilized in the active site.

**Keywords**
Mycobacterium tuberculosis, hemoglobin, water molecules, ligand interaction

This article is included in the **Oxygen-binding and sensing proteins** collection.

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**Competing interests:** No competing interests were disclosed.

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Introduction

*Mycobacterium tuberculosis*, the causative agent of human tuberculosis, affects approximately two billion people worldwide, causing over three million deaths each year. The genome of this pathogenic organism includes two genes, *glbN* and *glbO*, which encode for two proteins, termed here truncated hemoglobin N (Mt-trHbN) and truncated hemoglobin O (Mt-trHbO), belonging to the truncated hemoglobin (trHb) family of heme proteins, widely distributed in eubacteria, cyanobacteria, microbial eukaryotes and plants.

The truncated hemoglobin family exhibits a three-dimensional structure similar to the common globin fold of myoglobin, but significantly smaller. The secondary structure of trHbs consists of four α-helices arranged in a two-over-two antiparallel sandwich instead of the common three-over-three helix globin fold. Phylogenetic analysis has distinguished three different groups of truncated hemoglobins, classified as groups I, II and III, also called N, O and P, respectively.

It has been shown that group I Mt-trHbN catalyzes the detoxification of NO in the presence of O₂. The first step of this mechanism involves NO migration and binding. Subsequently, NO migrates to the active site and reacts with the heme-bound O₂ to yield an unstable peroxynitrite adduct, which isomerizes to generate the relatively innocuous nitrate anion.

Several studies have examined the role of internal tunnels in ligand migration in trHbs. Three different internal tunnels have been characterized among the trHb members, in general one or two of these tunnels is found in each protein: a long tunnel (LT) topologically positioned between helices B and E, and two short tunnels, known as the E7 Gate (E7 gate) and the short tunnel G8 (STG8), which are roughly normal to the LT, as depicted in Figure 1. The E7 tunnel corresponds to the highly conserved E7 pathway widely studied in both myoglobin and hemoglobin. The STG8 tunnel is analogous to that found in Mt-trHbN, next to the key residues VG8 and IH11. Previous results indicate that WG8, an absolutely conserved residue in groups II and III truncated hemoglobins, is involved in hindering ligand migration in Mt-trHbO by blocking both STG8 and LT (Figure 1). In addition, the presence of a smaller residue at the G8 position in the Mt-trHbO mutant (WG8F) was observed to increase the small ligand association constant, although the molecular details of this process were not investigated.

By performing CO association kinetic constant measurements (*kₐ* CO), NO decomposition, and molecular dynamics (MD) simulations of Mt-trHbN, we addressed molecular mechanisms that control ligand association in *M. tuberculosis* truncated hemoglobins.

![Figure 1. Schematic representation of the two pathways for ligand migration presented in *M. tuberculosis* trHbN. The Long Tunnel (LT) and Short Tunnel G8 (STG8) are shown in orange.](image-url)

Materials and methods

**Site-directed mutant construction**

The trHbN G8 mutants (VG8W and VG8F) were prepared using the Stratagene QuickChange mutagenesis kit. The following primers were designed using Primer3: forward primer 5’–CACCTCACGCTG–3’ and reverse primer 5’–CAAGTTGGGCAGCAGCTG–3’; forward primer 5’–ACCACCTCACGCTG–3’ and reverse primer 5’–CAAGTTGGGCAGCAGCTG–3’. Polymerase chain reaction (PCR) amplification of pET9b carrying the glbN gene with the aforementioned primers was conducted following manufacturer’s instructions. The PCR mix consisted of 5 µl 10x reaction buffer, 5–50 ng double stranded DNA template, 125 ng oligonucleotide primer 1, 125 ng oligonucleotide primer 2, 1 µl dNTP mix, 1 µl PfuUltra HF DNA polymerase and double distilled H₂O to a final volume of 50 µl. The PCR reaction was 95°C for 30 s, followed by 16 cycles of: 95°C for 1 min and 68°C for 4 min. The reaction mix was then digested with DpnI to remove parental methylated DNA. Plasmid containing the mutated gene was then purified and used to transform Escherichia coli XL-1 Blue electrocompetent cells. Cells were provided by Invitrogen. Constructs were checked by sequencing.

**Protein purification**

All chemicals and reagents were obtained from Sigma Aldrich, unless indicated otherwise. The trHbN protein variants were purified using standard techniques reported for other bacterial globins. Briefly, mutated constructs were used to transform *E. coli* BL21 DE3 pLysS. Starter cultures grown overnight in LB supplemented with kanamycin (50 µg ml⁻¹) and chloramphenicol (35 µg ml⁻¹) were used to inoculate 6 batches of 1 L LB medium at 1% (v/v), supplemented with kanamycin and 3 µM FeCl₃. Once cultures reached an OD₆₀₀ of around 0.4, expression of trHbN was initiated by the addition of 1 mM IPTG and grown for a further 4 h. Cells were harvested by centrifugation at 5500 rpm for 20 min at 4°C and stored overnight at -20°C. After thawing, cells were resuspended in 40 ml buffer (10 mM TRIS-HCl (pH 7.0) with 1 mM EDTA, 10 mM DTT,
45 µg ml⁻¹ phenylmethylsulphonyl fluoride, 500 µg ml⁻¹ RNase and 50 µl DNase), homogenized using a Douce homogeniser and ultracentrifuged at 44,000 rpm for 1 h at 4°C. The supernatant, red in color, was loaded onto a 30 ml DEAE Sepharose Fast Flow column (Pharmacia Biotech) equilibrated with 10 mM TRIS-HCl (pH 7.0), washed with the same buffer until the UV trace returned to baseline, and eluted via a gradient from 0 to 1 M NaCl in 10 mM TRIS-HCl using an Akta purifier (GE Healthcare Bio-Sciences, Amersham Biosciences, U.K. Ltd.). Fractions that were most red in color were concentrated using a Vivaspin 20 concentrator (Sartorius Stedim Biotech) to around 5 ml and loaded onto a gel filtration Superdex 75 column, equilibrated with 0.15 M NaCl in 10 mM TRIS-HCl (pH 7.5); again, fractions with the most color were collected, combined and stored at -80°C. Purity was checked using gel electrophoresis and analysis of the heme-to-protein ratio (410 nm and 280 nm in the UV-visible absorption spectrum).

Kinetic stopped flow measurements of CO binding

Rapid mixing experiments were conducted with a thermostated stopped flow apparatus (BioLogic SFM-300). Kinetics of carbon monoxide (CO) binding to determine the $k_{\text{on}}$ CO were measured on the deoxy state of mutant and wild type globins at 20°C. Solutions containing 5 µM protein in a 100 mM sodium phosphate at pH 7.0 were degassed in a nitrogen atmosphere and reduced with an equimolar concentration of sodium dithionite and mixed with increasing CO concentrations. The observed pseudo first-order rate constant ($k_{\text{on}}$) was determined by fitting the absorbance decay resulting from association of the protein with CO, to a single exponential function. Kinetic rate constants ($k_{\text{on}}$ CO) were obtained from the slope of the plots of $k_{\text{obs}}$ as a function of CO concentration.

NO decomposition

To determine rates of nitric oxide (NO) decomposition by wild type and mutant Mt-trHbN proteins, NO was added, as ProliNONOate, to a solution of 50 mM KPi buffer with 50 µM EDTA (pH 7.5), 100 µM NADPH and 100 nM E. coli ferredoxin reductase inside a thermostated, magnetically stirred reaction vessel. Mt-trHbN (2 µM) was added at the apex of the signal response to 2 µM ProliNONOate and NO decay was followed until depleted using an NO electrode (World Precision Instruments). Rates of NO decay were calculated for each protein by determining the time taken for peak NO concentrations to decay by 0.5 µM and were expressed per µM heme, determined spectrally by the peak in the Soret region at 410 nm.

Set up of the simulations

The starting structure corresponds to Mt-trHbN crystal structure (PDB entry 1IDR, http://www.rcsb.org/pdb/explore/explore.do?structureId=1IDR), at 1.9 Å of resolution). The protonation state of the amino acids was assumed based on the environment of the residues in the crystal structure. All solvent-exposed His residues were protonated at the N-atom, as well as the proximal HisF8, because of its coordination to the heme iron. An octahedral box of 10 Å of radius, which corresponds to 5234 explicit water molecules was added to the system. TIP3P water molecules were used by tLEaP module of the AMBER12 package. The params99 Amber force field was used for all the aminoacid parameters except heme parameters which were developed in our group and strongly validated for being used in several studies of heme proteins.

Periodic boundary conditions were used for all the simulations performed with a 9 Å cutoff. Particle mesh Ewald (PME) summation method for treating the electrostatic interactions. The SHAKE algorithm was used to keep constant the non-polar hydrogen equilibrium distance. Temperature and pressure were kept constant with Langevin thermostat and barostat, respectively, as implemented in the AMBER12 program. The equilibration simulation protocol was performed as follow: (i) slowly heating the system from 0 to 300K for 20 ps at constant volume, by using harmonic restraints of 80 kcal/mol Å² for all C atoms and (ii) pressure equilibration of the whole system during 1 ns at 300K with restrained atoms in (i). (iii) Unconstrained 100 ns molecular dynamics simulation at constant temperature (300K) was performed.

In silico mutant proteins were built by using tLEaP module of AMBER12 package, and underwent the same protocol used for wild type protein. Root Mean Square Deviation (RMSD) was used as structure stability controls. All structures were observed to be stables during the time scale of the simulation (Figure S1).

Ligand migration free energy profiles

The free energy profile for the CO migration process inside the protein tunnel/cavity system was computed by the Implicit Ligand Sampling (ILS) approach that post-processes, using a probe molecule, an MD simulation performed in the absence of the ligand. This method was thoroughly tested for heme proteins. ILS calculations were performed on a rectangular grid (0.5 Å resolution) that includes the whole simulation box (i.e. protein and the solvent) and the probe used was a CO molecule. Calculations were performed on 5000 frames taken from the last 90 ns of simulation time. The values for grid size, resolution and frame numbers were tested in a previous study. Analysis of the ILS data was performed using an ad-hoc fortran-90 program available upon request. Moreover, ILS has been shown to yield quantitative results for ligand migration processes when compared with more costly free energy methods that treat the ligand explicitly.

Results

CO association kinetic constant measurements

Although CO is not the natural ligand of the hemeproteins, it is widely used as a probe for ligand association studies due to its ease of use. In order to address the molecular determinants controlling ligand migration we performed CO ligand association constant measurements of wild type Mt-trHbN and two mutants: VG8F and VG8W. Kinetic traces for CO binding were measured through the absorbance of wild type Mt-trHbN and two mutants: VG8F and VG8W. Kinetic traces for CO binding were measured through the absorbance decay resulting from association of the protein with CO, to a single exponential function. The CO association kinetic constants ($k_{\text{on}}$) were obtained from the slope of Figure 3.

Association of CO is well described by a single exponential decay, whose rate constant ($k_{\text{obs}}$) depends linearly on CO concentration and the slope can be interpreted as $k_{\text{on}}$ CO. A significant $k_{\text{on}}$ CO decrease for VG8F (715 ± 27 mM⁻¹s⁻¹), and an even larger decrease for VG8W (48 ± 1 mM⁻¹s⁻¹) was observed in relation to that observed for the wild type protein (4495 ± 357 mM⁻¹s⁻¹) (Figure 3). Table 1 summarizes the measured $k_{\text{on}}$ CO values for wild type and mutant Mt-trHbs O and N, and is presented alongside literature data.

Molecular dynamics simulations

Small ligand association in the tHb family is presumably regulated by two main processes: i) ligand migration from solvent bulk to the
protein distal site cavity, ii) displacement of water molecules from the distal site cavity \(15-17, 35\). With this in mind, we performed classical MD simulations as they allow us to investigate both processes involved in ligand association. Ligand migration was studied using ILS calculations for the wild type, as well as both VG8F and VG8W mutant proteins. Displacement of retained water molecules in the distal site was considered by performing classical MD simulations and analyzing the solvation structure in each active site.

The wild type Mt-trHbN presents two tunnels available for ligand migration, the LT and the STG8 (Figure 4A). On the one hand, the LT connects three internal cavities: \((\text{trHb : CO})_1\), \((\text{trHb : CO})_2\), and \((\text{trHb : CO})_3\). The STG8, on the other hand, has the distal site cavity \((\text{trHb : CO})_1\) connected to both the cavity \((\text{trHb : CO})_2\) and the solvent, although the cavity \((\text{trHb : CO})_2\) plays only a secondary role, due to the fact that it does not alter the energy migration profile along the STG8 straight from the distal cavity to the solvent. The VG8F mutant conserves both tunnels, although they are constrained compared to those in the wild type. In the VG8W case, however, the energy profiles suggest a completely blocked STG8 and a LT for which the accessibility to the iron heme is partially reduced.

In order to quantify the contribution of the single G8 mutation we computed free energy profiles for CO migration through both LT and STG8 tunnels (Figure 4B, 4C). The free energy was set to a value of 0 kcal/mol where CO ligand is fully solvated at 13 Å and 24 Å from the Fe atom, for STG8 and LT respectively. Wild type Mt-trHbN presents small barriers (~2 kcal mol\(^{-1}\)) for CO migration from the solvent to the active site cavity \((\text{trHb : CO})_1\) through both tunnels.

The active site water molecules occupancy was computed for all three systems by performing 200 ns of MD simulations with explicit water molecules. In each case a water molecule was able to access the active site and was stabilized by the iron and the distal site residues (Figure 5). Specifically, in both wild type and VG8F Mt-trHbN a water molecule was present for approximately 40% of the length of the simulation (Figure 5A, 5B). The VG8W mutant active site, on the other hand, is occupied by water molecules in 80% of the simulation time, probably due to the hydrogen bonding capacity of W (Figure 5C).

\[ \text{NO decomposition in the presence of M. tuberculosis HbN} \]

Mt-trHbN has previously been described as a dioxygenase, capable of \(O_2\)-dependent ‘NO consumption\(^{13,39}\). Consequently, ‘NO decomposition by purified Mt-trHbN and the VG8F, VG8W mutants was

### Table 1. Association kinetic constants for wild type and mutants of Mt-trHbN and Mt-trHbO.

<table>
<thead>
<tr>
<th>Protein</th>
<th>(k_{\text{on}}) CO (mM(^{-1})s(^{-1}))</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mt-trHbN</td>
<td>4495</td>
<td>This work</td>
</tr>
<tr>
<td>Mt-trHbN VG8F</td>
<td>715</td>
<td>This work</td>
</tr>
<tr>
<td>Mt-trHbN VG8W</td>
<td>48</td>
<td>This work</td>
</tr>
<tr>
<td>Mt-trHbO</td>
<td>13 (79%) - 180 (21%) *</td>
<td>33</td>
</tr>
<tr>
<td>Mt-trHbO WG8F</td>
<td>3700 (75%) - 1200 (25%) *</td>
<td>12</td>
</tr>
</tbody>
</table>

*major and minor rate contributions to a biphasic fitting are indicated between brackets.

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**Figure 2.** Stopped-flow time course for the reaction of reduced 5 μM wild type (A), VG8F (B), and VG8W (C) mutants in 100 mM phosphate buffer at pH 7. The reaction was monitored at 423 nm (grey dots) and the line shows the best first order fit.

**Figure 3.** Apparent rates \(k_{\text{on}}\) for CO binding to ferrous Mt-trHbN. Curves for wild type (green), VG8F (orange) and VG8W (violet) mutants as a function of CO concentration in stopped flow measurements are shown. The time courses are measured at different CO concentrations ranging from 10 to 200 μM (after mixing). Continuous line corresponds to linear fit of \(k_{\text{on}}\) rates.
Figure 4. CO ligand migration along possible pathways in Mt-trHbN. (A) Schematic representations of the residues involved in the heme distal site and tunnels, the two tunnels and cavities estimated with ILS for the wild type form. Free energy profiles over STG8 (B) and LT (C) connecting the solvent with the distal site through the cavities (trHb : CO)_1, (trHb : CO)_2 and (trHb : CO)_3, for wild type (green), VG8F (orange) and VG8W (violet) mutant Mt-trHbN. Circles represent calculated free energy values with the ILS method and lines correspond to a fitting estimation of these calculated values. The x coordinate represents the Fe-CO distance along the pathways.

Figure 5. Schematic representations of the distal site of Mt-trHbN. (A) wild type, (B) VG8F and (C) VG8W forms showing, on the basis of MD simulations, the hydrogen-bond network (dotted lines) stabilizing a water molecule above the iron heme. The percentages depicted as insets in the figure correspond to active site water occupancy during MD simulation.
determined in a reaction mixture containing buffer, NADPH and *E. coli* FdR, to enable cyclic restoration of heme iron to the oxyferrous state. Figure 6A shows that in the absence of protein (red trace) decay of the •NO signal was monophasic until •NO was exhausted. The decay of NO in the presence of Mt-trHbN (black trace) was biphasic, with an almost linear initial rapid rate in decay, which was used to compare the various Mt-trHbN derivatives, followed by a slower rate in decay. This suggests that •NO is not being turned over in a cyclic manner, but is simply binding available heme. The chemical step being measured in this assay is the reaction between •NO and the oxyferrous heme; once this reaction has concluded, we assume that the heme is restored from ferric back to ferrous. We are unsure why the reaction is single turnover but it could be due to (a) rapid binding of •NO to the ferrous complex before oxygen can bind, rendering it unable to bind oxygen and initiate the reaction or (b) due to slow reduction of Mt-trHbN by the non-native *E. coli* FdR. •NO consumption results show that the VG8F and VG8W mutants have a statistically significant reduced •NO binding capacity compared to HbN (Figure 6B).

**Figure 6.** •NO decomposition by Mt-trHbN at ambient oxygen concentrations (approx. 200 µM, not measured). (A) •NO decay was monitored amperometrically in the absence (red trace) and the presence (black trace) of Mt-trHbN added at the apex of the signal response to 2 µM ProlinONOate. Data are representative of 3 technical repeats. (B) Mean rates of •NO decay in the presence of wild type Mt-trHbN or site-directed mutants from 3 technical repeats ± S.E.M * P < 0.05, unpaired t-test.
Grant information
This work was supported by Framework program 7 NOstress Grant, CONICET, University of Buenos Aires, and Agencia Nacional de Promoción Científica y Tecnológica, National Institutes of Health Grant R01AI095173 and Universidad de la República (CSIC, Uruguay) to R. R. Additional funding to SC and RR was provided by PEDECIBA (Programa de Desarrollo de Ciencias Básicas, Uruguay) and CeBEM (Centro de Biología Estructural del Mercosur). IB and JPB hold CONICET PhD fellowships. LB is a Pew Latin American Fellow. LB, DAE and MAM are members of CONICET.

I confirm that the funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Acknowledgements
FJ Luque is acknowledged for useful discussions and suggestions. Mehrnoosh Arrar is acknowledged for close reading of the manuscript.

Supplementary material

Figure S1. Root Mean Square Deviation during MD simulations for all the Mt-trHbN studied: wild type (green), VG8F (orange) and VG8W (violet).

References


Open Peer Review

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Version 2

Reviewer Report 30 September 2015

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This article shows that the previously observed structural and functional features of trHbO, using its G8 mutants, can be duplicated in trHbN. As such, the critical role of the G8 residue and LT and STG8 channels, and E7 gate in functions of trHbO as well as trHbN is demonstrated.

Some technical questions to be answered are:

1. In Fig. 2, the identical concentrations (5μM) of trHbNs are used. However, values of ΔAbsorbance of Figs. 2A (an estimated value of ΔA=0.05), 2B (an estimated value [extrapolated to t = 0] of ΔA=0.55), and 2C (an estimated value of ΔA=0.25). Why do such large discrepancies in ΔAs exist among Figs. 2A, B, and C, though sample concentrations are identical (5μM)? Why is the noise level of the trace in Fig. 2A is much smaller than those of Fig. 2B and 2C, though the latters have much larger ΔA values of 0.55 and 0.25, respectively?

2. The interpretation of the results of the Fig. 6A experiments is not convincing, because not enough data are presented. The NO decay assays were performed under the background of rapid spontaneous decay of NO (approximately 5.4 μM/100sec) to determine Δ[NO] (between with and without trHbN) = approximately 0.5 μM. Kinetic curves shown are extremely smooth to indicate a very large time constant of the NO assay instrument. Therefore, quantitative assay of Δ[NO] from the time courses of kinetic curves are unreliable. In order to prove the proposed one-cycle hypothesis, why didn't they measure Δ[NO] values as a function of [trHbN]? From these reasons, the error bars in Fig. 6B seem unusually small. Are these differences in the NO decay rate (X-axis), shown in Fig. 6B, statistically significant?

3. MD analyses are done under the condition of a single molecule of CO binding to trHbN or to its mutants. Since the hemes themselves in trHbN and mutants have low-affinity for CO, though the apparent affinity for CO of trHbN is known to be high, which I believe, binding of a single molecule of CO to the heme of deoxy trHbN is an energetically unfavorable up-hill process. How are the energy levels of the initial stage (deoxy trHbN + CO), relative to the final stage (trHb: CO) shown
in Fig. 4B, determined? Are they initially assumed? Or they are results of MD analysis? Shouldn’t the MD analyses be done under conditions where solvents contain large numbers of CO molecules, rather than a single CO molecule, thermodynamically speaking?

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 25 September 2015

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Michael Wilson
School of Biological Sciences, University of Essex, Colchester, UK

The authors have responded positively to my earlier comments and in my view have improved the manuscript.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Version 1**

Reviewer Report 24 April 2015

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Michael Wilson
School of Biological Sciences, University of Essex, Colchester, UK

This is a succinct and generally well written paper reporting experimental data on ligand binding to Mt-trHbN and to two mutants (VG8F and VG8W). These latter have been designed to test hypotheses regarding the possible routes by which access is gained to the heme by small neutral gaseous molecules. The experiments and the molecular modelling that supports them, and which provides mechanistic
insights, have been carefully performed. The results are of interest to the field as they add to the body of accumulated evidence that proteins, including those with the function of binding small neutral molecules, provide specific, and often dynamic, channels to permit rapid access to the binding site. Furthermore the kinetics of binding are seen to be strongly influenced by single amino acid substitution in the access channels.

Although the results support the general conclusions drawn by the authors some clarification of a number of points would be helpful. These are given below.

1. Why in Figs 2A and B does time appear not to start at t=0?

2. In Fig and discussion is it proposed that the water molecule is bound to the iron (common for ferric but not for ferrous iron) or stabilised in that location only by hydrogen binding. It is presumed that for the modelling the iron is in the ferrous state as the authors are discussing CO binding.

3. Although the authors discuss the decomposition of NO catalysed by Mt-trHbN in the presence of oxygen the assays do not make it clear that this is the reaction under study. No mention of the oxygen concentration is made in the legend to Fig 6. It seems oxygen is present to account for the disappearance of NO. From Fig 6 it is stated that NO binds to the protein but is not degraded (e.g. to nitrate via peroxynitrite). Is this because regeneration of the reduced heme (necessary for oxygen binding) is so slow or is it because as ferric heme is reduced by the NADPH/ferredoxin system NO binds before oxygen and thus no turnover occurs as it is known that the NO-ferrous complex does not react with oxygen? In any case it is not made clear to what chemical step the measured kinetics refer. It would improve the manuscript if the authors clarified these points.

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 13 Jul 2015

Leónardo Boechi, Universidad de Buenos Aires, Buenos Aires, Argentina

1. **Why in Figs 2A and B does time appear not to start at t=0?**

   We thank the reviewer for noticing that specific issue. We decided to discard the first 8ms because they were too noisy.

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   We agree with the reviewer that the water bounds in general very weak to the ferrous iron. However, there is evidence showing that in a polar distal site (with polar residues), as in the case of truncated hemoglobins, water molecules remain inside stabilized by the polar


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No competing interests were disclosed.

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**Author Response 13 Jul 2015**

**Leónardo Boechi**, Universidad de Buenos Aires, Buenos Aires, Argentina

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   We thank the reviewer for noticing that specific issue. We decided to discard the first 8ms because they were too noisy.

2. **In Fig and discussion is it proposed that the water molecule is bound to the iron (common for ferric but not for ferrous iron) or stabilised in that location only by hydrogen binding. It is presumed that for the modelling the iron is in the ferrous state as the authors are discussing CO binding.**

   We agree with the reviewer that the water bounds in general very weak to the ferrous iron. However, there is evidence showing that in a polar distal site (with polar residues), as in the case of truncated hemoglobins, water molecules remain inside stabilized by the polar
2. **residues, and thus slow ligand binding to ferrous iron** (Olson and Phillips Jr, 1997; Ouellet *et al.*, 2008).

3. Although the authors discuss the decomposition of NO catalysed by Mt-trHbN in the presence of oxygen the assays do not make it clear that this is the reaction under study. No mention of the oxygen concentration is made in the legend to Fig 6. It seems oxygen is present to account for the disappearance of NO. From Fig 6 it is stated that NO binds to the protein but is not degraded (e.g. to nitrate via peroxynitrite). Is this because regeneration of the reduced heme (necessary for oxygen binding) is so slow or is it because as ferric heme is reduced by the NADPH/ferredoxin system NO binds before oxygen and thus no turnover occurs as it is known that the NO-ferrous complex does not react with oxygen? In any case it is not made clear to what chemical step the measured kinetics refer. It would improve the manuscript if the authors clarified these points.

As suggested by the review we modified the caption of Figure 6, and also we add a sentence.

The legend of the caption can be changed to:

Figure 6. •NO decomposition by Mt-trHbN at ambient oxygen concentrations (approx. 200 µM, not measured). (A) •NO decay was monitored amperometrically in the absence (red trace) and the presence (black trace) of Mt-trHbN added at the apex of the signal response to 2 µM ProliNONOate. Data are representative of 3 technical repeats. (B) Mean rates of •NO decay in the presence of wild type Mt-trHbN or site-directed mutants from 3 technical repeats ± S.E.M *P < 0.05, unpaired t-test.

The sentence added is:

The chemical step being measured in this assay is the reaction between •NO and the oxyferrous heme; once this reaction has concluded, we assume that the heme is restored from ferric back to ferrous. We are unsure why the reaction is single turnover but it could be due to (a) rapid binding of •NO to the ferrous complex before oxygen can bind, rendering it unable to bind oxygen and initiate the reaction or (b) due to slow reduction of Mt-trHbN by the non-native *E. coli* FdR.

**Competing Interests:** No competing interests were disclosed.
Marco Nardini
Department of Biosciences, University of Milan, Milan, Italy

This manuscript describes CO association kinetic constant measurements, NO decomposition, and molecular dynamics simulations on the wild type truncated Hb from *Mycobacterium tuberculosis* (Mt-trHbN) and two mutants (VG8F and VG8W) which introduce modifications in the two-tunnel system of the protein. The data are then compared to those from Mt-trHbO, suggesting that ligand migration is regulated by the internal tunnel architecture as well as by water molecules stabilized in the active site.

Although the topic of the structure and the dynamic behavior of protein matrix tunnels in truncated Hb, and in particular in Mt-trHbN, has been “squeezed” a lot during the past years, the data reported in this paper add some new information and might be of interest in the field. The paper is well written (with regards to the requirements of the journal) and describes a technically sound piece of scientific research with data that supports the conclusions. Indexing is recommended, if the (few) minor comments below are addressed.

Minor remarks:
- **Abstract: line 4**
  The authors write that “Mt-trHbO” possesses tunnels that are partially blocked.” In fact normally trHbOs are associated with internal discrete cavities and not with tunnels. The authors should rephrase the sentence.

- **Abstract: line 11**
  *Mt-trHbN* should not be in Italics.

- **Abstract: line 12**
  The sentence “mutations introduce modifications in both tunnel topologies” is quite cryptic and it is not clear what the author mean with “tunnel topologies”. The authors should rephrase the sentence to clarify it.

- **Introduction: page 3, first column, line 8**
  The authors might want to include a review on trHbs more recent than that indicated in reference (3). There are several of them published in the last few years.

- **Introduction: page 3, first column, line 23**
  The paragraph starting from line 23 is a bit misleading because the authors try to generalize the description of the protein matrix tunnels in trHbs by mixing what happens in trHbNs and trHbOs. This is confusing since it might give the impression that three tunnels co-exist in trHbs, which is not true. In this respect, Figure 1 contributes a lot to make confusion, since it is not clear which trHb protein represents and it seems that it has three co-existing tunnels. It is probably better to keep separate trHbNs and trHbOs, both in the text description and in Figure 1. The authors should describe the tunnel features in trHbN (short and long tunnel) and trHbO (cavities, small E7 residues an possible E7 gating), and show two panels in Figure 1 with depicted the tunnel/cavity systems in Mt-trHbN (panel A) and Mt-trHbO (panel B), possibly using a similar protein orientation and highlighting the role of the G8 residue in the two cases.

- **Introduction: page 3, second column, line 7**
  The sentence regarding the “internal water molecules” is too generic as it is written now, since it is not clear if the authors refer to globins, to trHbs or to Mt-trHbs. The authors should rephrase the
sentence to clarify this issue.

- Introduction: page 3, second column, line 10
  The authors should say that the experimental measurements and the MD simulations have been performed only on Mt-trHbN and mutants, and not, for instance, on Mt-trHbO.

- Materials and Methods: page 3, second column, line 40
  The purification paragraph seems to refer only to trHbN. What about its mutants? The authors should add a sentence to clarify this issue.

- Materials and Methods: page 4, first column, line 44
  The following sentence is not written fully correctly:
  “Amino acids protonation states were assumed based on environment of the residue in the crystal structure. All solvent-exposed His were protonated at the N-δ delta atom, as well as HisF8, because of its coordination to the heme iron”.

  One possibility is to rephrase it as follows:
  “The protonation state of the amino acids was assumed based on the environment of the residues in the crystal structure. All solvent-exposed His residues were protonated at the N-δ atom, as well as the proximal HisF8, because of its coordination to the heme iron”.

- page 5, second column, line 12
  It is not clear what the authors mean when they write that the STG8 “has only the distal site cavity, (trHb : CO)₁, …”, especially if this sentence is coupled with Figure 4A, where (trHb : CO)₁ seems to be connected to STG8 through (trHb : CO)₂.

- page 5, Title of Table 1
  It is probably better to change “..for wild type and mutants Mt-trHbs O and N” to “..for wild type and mutants of Mt-trHbN and Mt-trHbO”

- page 6, Figure 4 legend
  In the legend of Panel C there is no mention of the (trHb : CO)₃ site.

- References: page 9
  Reference (21) is missing the title

**Competing Interests:** No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Author Response 13 Jul 2015**

**Leonardo Boechi,** Universidad de Buenos Aires, Buenos Aires, Argentina

- **Abstract: line 4**
  The authors write that “Mt-trHbO” possesses tunnels that are partially blocked ..” In fact normally trHbOs are associated with internal discrete cavities and not with tunnels. The authors should rephrase the sentence.
We rephrase the sentence “whereas Mt-trHbO possesses tunnels that are partially blocked by a few bulky residues, particularly a tryptophan at position G8” by writing “whereas Mt-trHbO possesses tunnels interrupted by a few bulky residues, particularly a tryptophan at position G8”

- **Abstract: line 11**
  
  *Mt-trHbN should not be in Italics.*
  
  Corrected, thanks

- **Abstract: line 12**
  
  The sentence “mutations introduce modifications in both tunnel topologies” is quite cryptic and it is not clear what the author mean with “tunnel topologies”. The authors should rephrase the sentence to clarify it.

  We rephrase the sentence “These mutations introduce modifications in both tunnel topologies and affect the incoming ligand capacity to displace retained water molecules at the active site.” By writing “These mutations affect both the tunnels accessibility as well as the affinity of distal site water molecules, thus modifying the ligand access to the iron”

- **Introduction: page 3, first column, line 8**
  
  The authors might want to include a review on trHbs more recent than that indicated in reference (3). There are several of them published in the last few years.

  As suggested by the review we modified the references by more recent ones: Davidge & Dikshit (2013).

- **Introduction: page 3, first column, line 23**
  
  The paragraph starting from line 23 is a bit misleading because the authors try to generalize the description of the protein matrix tunnels in trHbs by mixing what happens in trHbNs and trHbOs. This is confusing since it might give the impression that three tunnels co-exist in trHbs, which is not true. In this respect, Figure 1 contributes a lot to make confusion, since it is not clear which trHb protein represents and it seems that it has three co-existing tunnels. It is probably better to keep separate trHbNs and trHbOs, both in the text description and in Figure 1. The authors should describe the tunnel features in trHbN (short and long tunnel) and trHbO (cavities, small E7 residues an possible E7 gating), and show two panels in Figure 1 with depicted the tunnel/cavity systems in Mt-trHbN (panel A) and Mt-trHbO (panel B), possibly using a similar protein orientation and highlighting the role of the G8 residue in the two cases.

  As suggested by the review, we modified the sentence to clarify.

  “Three internal tunnels were found in the truncated hemoglobin family;” by “Three different internal tunnels have been characterize among the trHb members, in general one or two of these tunnels is found in each protein;”

  We also change Figure 1 and its caption as suggested.
Caption Figure 1. Schematic representation of the two pathways for ligand migration presented in *M. tuberculosis* trHbN. The Long Tunnel (LT) and Short Tunnel G8 (STG8) are shown in orange.

- Introduction: page 3, second column, line 7
  The sentence regarding the "internal water molecules" is too generic as it is written now, since it is not clear if the authors refer to globins, to trHbs or to Mt-trHbs. The authors should rephrase the sentence to clarify this issue.

  As suggested by the review we clarify the sentence: “It has also been noted that in myoglobin, *M. Tuberculosis* trHbN as well as in *T. fusca* trHbO, internal water molecules were observed to block the heme accessibility, thus delaying ligand binding”

- Introduction: page 3, second column, line 10
  The authors should say that the experimental measurements and the MD simulations have been performed only on Mt-trHbN and mutants, and not, for instance, on Mt-trHbO.

  As suggested by the review we clarify the sentence by adding explicitly the name of the protein studied “By performing CO association kinetic constant measurements (…) of Mt-trHbN, we addressed molecular mechanisms that control ligand association in *M. Tuberculosis* truncated hemoglobins”

- Materials and Methods: page 3, second column, line 40
  The purification paragraph seems to refers only to trHbN. What about its mutants? The authors should add a sentence to clarify this issue.

  We clarify this by modifying “The trHbN protein was” by “The trHbN protein variants were”

- Materials and Methods: page 4, first column, line 44
  The following sentence is not written fully correctly: “Amino acids protonation states were assumed based on environment of the residue in the crystal structure. All solvent-exposed His were protonated at the N-δ delta atom, as well as HisF8, because of its coordination to the heme iron”.

  One possibility is to rephrase it as follows: “The protonation state of the amino acids was assumed based on the environment of the residues in the crystal structure. All solvent-exposed His residues were protonated at the N-δ atom, as well as the proximal HisF8, because of its coordination to the heme iron”.

  We thank the reviewer for the suggestion, the phrase was modified as suggested.

- page 5, second column, line 12
  It is not clear what the authors mean when they write that the STG8 “has only the distal site cavity, (trHb : CO)1, …”, especially if this sentence is coupled with Figure 4A, where (trHb : CO)1 seems to be connected to STG8 through (trHb : CO)2.

  We modified the sentence by adding information:

  On the one hand, the LT connects three internal cavities: (trHb : CO) 1, (trHb : CO) 2 and
(trHb : CO) 3. The STG8, on the other hand, connected to both the cavity (trHb : CO) 2 and the solvent.

- **page 5, Title of Table 1**
  It is probably better to change “..for wild type and mutants Mt-trHbs O and N” to “..for wild type and mutants of Mt-trHbN and Mt-trHbO”

  We thank the reviewer for the suggestion; the phrase was modified as suggested.

- **page 6, Figure 4 legend**
  In the legend of Panel C there is no mention of the (trHb : CO) 3 site.

  We thank the reviewer for the suggestion, we rephrase as “Free energy profiles over STG8 (B) and LT (C) connecting the solvent with the distal site through the cavities (trHb : CO) 1 , (trHb : CO) 2 and (trHb : CO) 3 , for wild type (green), VG8F (orange) and VG8W (violet) mutant Mt-trHbN.”

- **References: page 9**
  Reference (21) is missing the title

  We thank the reviewer; the reference was modified

**Competing Interests:** No competing interests were disclosed.