Validating the GTP-cyclohydrolase 1-feedback regulatory complex as a therapeutic target using biophysical and in vivo approaches

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BACKGROUND AND PURPOSE
6R-L-erythro-5,6,7,8-tetrahydrobiopterin (BH₄) is an essential cofactor for nitric oxide biosynthesis. Substantial clinical evidence indicates that intravenous BH₄ restores vascular function in patients. Unfortunately, oral BH₄ has limited efficacy. Therefore, orally bioavailable pharmacological activators of endogenous BH₄ biosynthesis hold significant therapeutic potential. GTP-cyclohydrolase 1 (GCH1), the rate limiting enzyme in BH₄ synthesis, forms a protein complex with GCH1 feedback regulatory protein (GFRP). This complex is subject to allosteric feed-forward activation by L-phenylalanine (L-phe). We investigated the effects of L-phe on the biophysical interactions of GCH1 and GFRP and its potential to alter BH₄ levels in vivo.

EXPERIMENTAL APPROACH
Detailed characterization of GCH1–GFRP protein–protein interactions were performed using surface plasmon resonance (SPR) with or without L-phe. Effects on systemic and vascular BH₄ biosynthesis in vivo were investigated following L-phe treatment (100 mg·kg⁻¹, p.o.).

KEY RESULTS
GCH1 and GFRP proteins interacted in the absence of known ligands or substrate but the presence of L-phe doubled maximal binding and enhanced binding affinity eightfold. Furthermore, the complex displayed very slow association and dissociation rates. In vivo, L-phe challenge induced a sustained elevation of aortic BH₄, an effect absent in GCH1(fl/fl)-Tie2Cre mice.

CONCLUSIONS AND IMPLICATIONS
Biophysical data indicate that GCH1 and GFRP are constitutively bound. In vivo, data demonstrated that L-phe elevated vascular BH₄ in an endothelial GCH1 dependent manner. Pharmacological agents which mimic the allosteric effects of L-phe on the GCH1–GFRP complex have the potential to elevate endothelial BH₄ biosynthesis for numerous cardiovascular disorders.
Abbreviations

BH₄, 6R-L-erythro-5,6,7,8-tetrahydrobiopterin; Bₘₐₓ, maximal binding; F-GCH1, native/full length GCH1; GCH1, GTP cyclohydrolase 1; GFRP, GCH1 feedback regulatory protein; Kₛ, equilibrium dissociation constant; kₐᵣ, dissociation rate constant; kᵣₒ₃, association rate constant; L-phe, L-phenylalanine; NOS, nitric oxide synthase; PKU, phenylketonurea; SPR, surface plasmon resonance; T-GCH1, truncated GCH1

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<td>NOS, nitric oxide synthase</td>
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These Tables list key protein targets and ligands in this article which are hyperlinked to corresponding entries in http://www.guidetopharmacology.org, the common portal for data from the IUPHAR/BPS Guide to PHARMACOLOGY (Pawson et al., 2014) and are permanently archived in the Concise Guide to PHARMACOLOGY 2013/14 (Alexander et al., 2013).

Introduction

GTP-cyclohydrolase 1 (GCH1) catalyses the committing and rate limiting step in the production of 6R-L-erythro-5,6,7,8-tetrahydrobiopterin (BH₄), an essential cofactor for aromatic amino acid hydroxylases (Kaufman, 1963), nitric oxide synthase (NOS) (Tayeh and Marletta, 1989) and alkylglycerol mono-oxygenase (Watschinger et al., 2010). The products of these enzymes have widespread functions (Thöny et al., 2000) and hence GCH1, via its control of BH₄ biosynthesis, regulates a number of diverse physiological systems.

Numerous clinical studies have shown that intravascular BH₄ administration, at high concentrations, significantly improves endothelial function in patients with a wide range of cardiovascular disorders, by increasing nitric oxide (NO) bioavailability and/or reducing oxidative stress (Heitzer et al., 2002; Higashi et al., 2002; Wyss et al., 2005; Mayahi et al., 2007). Unfortunately, BH₄ is very unstable and, when orally administered, has limited efficacy likely due to oxidation during absorption (Cunnington et al., 2012). Therefore, other approaches to directly raise endogenous BH₄ biosynthesis hold therapeutic potential – one such approach involves the pharmacological activation of GCH1, within the vascular endothelial cells themselves.

Interestingly, mammals have evolved an endogenous system to dynamically regulate GCH1 activity, whereby the enzyme’s activity can be inhibited by BH₄ (via end product feedback inhibition), or activated by L-phenylalanine (L-phe). However, this dynamic post-translational regulation only occurs when GCH1 is bound to GCH1 feedback regulatory protein (GFRP) (Harada et al., 1993). In vitro studies have demonstrated that when GCH1 and GFRP are bound, BH₄ mediates allosteric feedback inhibition of GCH1, in a non-competitive manner, whilst L-phe can reverse this effect – stimulating GCH1 activity (Harada et al., 1993; Maita et al., 2004). Importantly, this allosteric regulation of GCH1 activity has an absolute requirement for GFRP, as BH₄ and L-phe are unable to influence the activity of purified GCH1 protein in isolation (Harada et al., 1993).

These feedback and feed-forward mechanisms ensure that BH₄ is kept within a tight physiological range in the body. As BH₄ is an essential cofactor for the metabolism of dietary L-phe by phenylalanine hydroxylase, the L-phe mediated feed-forward activation of GCH1 raises endogenous BH₄ and ensures that dietary L-phe is adequately metabolized. This is clinically important as L-phe can be neurotoxic above a certain concentration, leading to irreversible mental disability. Indeed, BH₄ supplementation is currently used to treat a subset of L-phenylketonuria (PKU) patients who are unable to metabolize L-phe (Heintz et al., 2013). The differential efficacy of oral BH₄ supplementation between coronary artery disease patents and those with PKU may be explained by the presence of enhanced oxidative stress, which subsequently leads to oxidative inactivation of BH₄ during absorption in the former, but not the latter, patient group (Cunnington et al., 2012).

The presence of a functional GCH1–GFRP complex has been demonstrated in humans, whereby oral administration of L-phe elicits an increase in plasma biopterin (a surrogate marker of BH₄) – an effect that is attenuated in patients carrying a loss of function GCH1 mutation (Saunders-Pullman et al., 2004). However, the effects of L-phe on biopterin and, more importantly, BH₄ levels in tissues have not been directly determined.

The crystal structure of the GCH1–GFRP complex has been solved, revealing GCH1 as a homodecamer (~280 kDa) sandwiched between two GFRP homopentamers (~50 kDa). These crystal structures revealed discrete binding pockets for L-phe and BH₄ located at the GCH1–GFRP interface and distinct from the GTP binding site (Maita et al., 2002; 2004), making these unique and rational drug targets to either enhance or limit BH₄ biosynthesis respectively. Upon binding to their respective pockets at the GCH1–GFRP interface, BH₄ and L-phe induce conformational changes in the remote GTP substrate binding pocket, impeding or facilitating GTP binding respectively (Maita et al., 2002; 2004). However, to date, crystallization studies have been limited to a truncated form of mammalian GCH1, lacking the first 42 amino acids...
due to N-terminal instability (Auerbach et al., 2000; Maita et al., 2002; 2004). Whilst these studies stated that this N-terminal region did not influence GFRP binding, feedback regulation or GCH1 activity, other studies have contradicted these findings, suggesting that the absence of a complete N-terminal region can alter GCH1 activity and further limits the capacity of GCH1 to bind to GFRP (Swick and Kapatos, 2006; Higgins and Gross, 2011).

We hypothesized that drugs which mimic the allosteric effects of L-phe on the GCH1–GFRP complex have the potential to elevate BH4 within vascular cells and restore endothelial function in numerous cardiovascular disorders, circumventing the limitations of oral BH4 treatment. However, to facilitate a rational drug discovery approach, a greater understanding of the GCH1 : GFRP complex and the potential limitations of the current crystal structures (which used an N-terminal truncation mutant) is required. Furthermore, in vitro and in vivo proof of concept studies validating the GCH1–GFRP axis as a tangible drug target to regulate endothelial BH4 are also lacking.

Therefore, in this study, we have quantified GCH1–GFRP protein interactions using surface plasmon resonance (SPR), comparing native/full length GCH1; GFRP, GCH1 feedback regulatory protein; T-GCH1, truncated GCH1.

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Methods

Cloning and construct formation

Human liver mRNA (AMS Biotechnology UK Ltd: M1234149) was reverse transcribed using the GE Healthcare First Strand cDNA Synthesis Kit. DNA encoding native GFRP, F-GCH1 and T-GCH1 (lacking the first 42 N-terminal amino acids) was amplified from the cDNA by PCR using KOD hot start polymerase (VWR International, Lutterworth, UK). PCR products (GFRP, F-GCH1 and T-GCH1) were individually purified, digested and ligated into the dual expression vector pDuet-1 encoding an N-terminal His6-tag in MCS1, using appropriate restriction enzymes. MCS2 was used for co-expression of non-tagged proteins. All PCR primers encoded a TEV cleavage site for removal of the His6 tag after protein purification (Table 1).

<table>
<thead>
<tr>
<th>Construct</th>
<th>Vector and multiple cloning site</th>
<th>Restriction enzyme ± TEV site</th>
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Each construct was generated by PCR amplification with corresponding oligonucleotide primers, as listed. A TEV site was incorporated for removal of the His6 tag. T-GCH1 (truncated mutant lacking the first N-terminal 42 amino acids).

F-GCH1, native/full length GCH1; GFRP, GCH1 feedback regulatory protein; T-GCH1, truncated GCH1.

Protein expression and purification

Overnight bacterial cultures were induced using 1 mM isopropyl β-D-1-thiogalactopyranoside at 25°C for 12 h. His6-tagged proteins, either alone or bound to non-tagged co-expressed proteins, were purified using Talon cell-thru metal affinity resin (Takara-Bio Europe/Clontech, Saint-Germain-en-Laye, France). For SPR experiments, purified pro-
teinase were incubated with TEV protease at 30°C overnight to cleave the His-tag, and then co-incubated with the affinity resin (4 h) to remove the cleaved His-tag. Finally, protein samples were run through a size exclusion chromatography Superdex column (GE Healthcare Ltd, Buckinghamshire, UK).

Proteins were identified by SDS-PAGE and Western blotting using either a polyclonal primary GCH1 anti-peptide antibody (raised against amino acid residues 18–45) (Nandi et al., 2005), a commercial primary GFRP antibody (Santa Cruz Biototechnologies, Santa Cruz, CA) or a His-tag antibody (Abcam, Cambridgeshire, UK). Purified protein samples were combined and concentrated in 100 mM Tris pH 7.8, 100 mM NaCl or 50 mM HEPES pH 7.8, 100 mM KCl.

**GCH1 activity (HPLC).** GCH1 activity was assessed by HPLC as previously described (Howells et al., 1986), whereby neopterin content was quantified by isocratic HPLC and fluorescence detection. Quantification of neopterin was carried out by comparison with external standards and was normalized for sample protein content.

**GCH1 activity kinetic microplate assay.** An established kinetic microplate assay was modified and used to measure GCH1 activity in expressed proteins, in addition to HPLC (Kolinsky and Gross, 2004). This assay measures the accumulation of the intermediate reaction product, dihydronopterin triphosphate (H$_2$NTP), by monitoring an increase in $A_{340}$ over time. The assay was set up in a 96-well plate format as follows: 0.25 μM GCH1 protein (T or F) was combined with assay buffer (100 mM Tris–HCl to a final volume of 300 μL) and 100 μM GTP (Thermo Fisher Scientific, Hemel Hempstead, UK). Purified GFRP protein (1 μM) was added in certain experiments. Absorbance (340 nm) was measured using a Spectramax temperature controlled plate reader (Molecular Devices Ltd, Wokingham, UK) at 37°C until the reaction reached saturation. Absorbance units were expressed in mol H$_2$NTP as previously described (Kolinsky and Gross, 2004).

**SPR**

A Biacore™ T200 was used to conduct SPR experiments. F-GCH1, T-GCH1 or GFRP proteins obtained from single expression vectors were captured on a CM5 sensor chip surface using an anti-His-tag antibody (Biacore His-capture Kit; GE Healthcare Ltd). The sensor chip was activated using a 1:1 mixture of 50 mM N-hydroxysuccinimide and 200 mM of N-ethyl-N′-(diethylaminopropyl) carbodiimide. This was injected across two flow cells simultaneously, with the second flow cell acting as a control surface to identify any non-specific binding. A 50 μg·mL$^{-1}$ anti-His-tag antibody solution was injected over the experimental flow cell and both experimental and control surfaces were subsequently quenched with 1 M ethanolamine HCl (pH 8.5). His-tagged protein (100 nM) in running buffer (50 mM HEPES pH 7.4, 150 mM NaCl, 3 mM EDTA, and 0.005% (v/v) surfactant P20) was injected across both flow cells, allowing the protein to be captured by the anti-His-tag antibody immobilized on the surface of the experimental flow cell. Experiments were performed with F-GCH1 or T-GCH1 captured on a CM5 surface (with GFRP as the analyte), and reciprocally with GFRP captured on the surface (with either T-GCH1 or F-GCH1 as the analyte). For the analyte proteins, the His-tag was cleaved and the protein incorporated into the running buffer over a range of concentrations (100–2000 nM). Analyte proteins were injected across both flow cells. The surface was regenerated using an injection of 10 mM glycine (pH 1.5) following each analyte cycle. In separate experiments, L-phe (100 μM) or BH$_4$ (20 μM) was introduced along with the analyte protein into the running buffer. Flow rates were adjusted accordingly to enable equilibration (flow rate of 3 μL·min$^{-1}$ for 7000 s). All SPR assays were conducted at 25°C. Surface density after ligand immobilization was maintained at 1250–1500 RU for all experiments. Dose-dependent association experiments and binding characterization experiments were repeated four times.

**In vitro and vivo studies**

**Animal welfare and ethical statement.** All animals studies described in this paper were conducted following ethics approval and in accordance with the UK Home Office Animals (Scientific Procedures) Act 1986 [Amendment Regulations (SI 2012/3039)]. Experimental design and conduct were undertaken in accord with the ARRIVE guidelines (Kilkenny et al., 2010) and complied with The Basel Declaration and the Concordat on Openness on Animal Research. All techniques used for in vivo studies were as humane as possible. A total of 80 animals were used in the experiments described here.

**L-phe challenge in wild-type and GCH1(fl/fl)-Tie2Cre mice.** Three groups of mice were used for in vivo studies: (i) male C57BL/6 mice (12–14 weeks old), purchased from a commercial supplier (Harlan Laboratories, Loughborough, UK); (ii) GCH1(fl/fl)-Tie2Cre (KO) mice and (iii) GCH1(fl/fl) mice – hereafter referred to as wild-type littermates – generated by crossing male GCH1(fl/fl)-Tie2Cre and female GCH1(fl/fl) mice (Chuaiphichai et al., 2014). All animals were group housed with 12 h light/dark cycle and controlled temperature of 20–22°C and given access to a standard chow diet and water ad libitum. GCH1(fl/fl)-Tie2Cre mice have previously been shown to lack GCH1 in endothelial cells (Chuaiphichai et al., 2014).

L-phe (100 mg·kg$^{-1}$) in saline was administered orally (by gavage) to all three groups of mice. This dose is equivalent to ~200 g of beef in a healthy 70 kg adult (Uhe et al., 1997) and matches the dose used in L-phe loading studies in PKU patients (Saunders-Pullman et al., 2004). Mice were killed, by exsanguination under isoflurane anaesthesia, after 20 min, 1, 2, 4 or 8 h. Plasma and aortic tissues were collected and snap frozen for subsequent analysis of L-phe, biopterin and BH$_4$ levels. L-phe levels were measured using HPLC detection in plasma and tissue homogenates following sample preparation as described previously (Atherton and Green, 1988). Quantification of L-phe was performed by comparison with external standards (0–250 mmol·L$^{-1}$) and the lower limit of detection was 2.7 μM. Values were normalized for protein content using a standard Bradford assay.

**L-phe challenge in sEnd1 endothelial cells.** sEnd-1 cells (a stable murine endothelial cell line) were cultured in DMEM as previously described (Nandi et al., 2008) and used between passages 3 and 7 for all experiments. Human modified oxi-
dized lipoprotein (RP-048 – Source BioScience Life Sciences, Nottingham, UK) (100 μg·mL⁻¹) (Feldmann et al., 2013) was incubated with cells for 2 h to induce oxidative stress (Bowers et al., 2011). L-Phe (500 μM; 0.5 h) or vehicle control was subsequently added to cells, and the impact on nitrite accumulation (a correlate of nitric oxide) in the media and BH₄ in cell lysates was assessed.

**BH₄ measurement.** Bioperin and BH₄ were measured in cell lysates, tissue and plasma, as previously described using fluororescence and electrochemical detection following sample separation by HPLC (Howells et al., 1986; Starr et al., 2014). Quantification was performed by comparison with external standards after normalizing for sample protein content. All analyses were conducted in a blinded fashion and investigators were unblinded to treatment/genotype following completion of data analysis.

**Nitrite measurement.** Plasma or tissue homogenates were deproteinated and nitrite content was then quantified using a fluorometric method utilizing 2,3-diaminonaphthalene (Bryan and Grisham, 2007). The amount of nitrite in each sample, expressed as micromolar nitrite-per milligram protein was calculated from a linear calibration curve of known nitrite concentrations (linear range: 0.5–5 μM) and normalized for total amount of protein.

**Chemiluminescent measurement of superoxide anion.** Superoxide levels were quantified from sEnd1 cells using a Lucigenin chemiluminescence-based assay (Li and Shah, 2001). Briefly, cells were seeded at equal density in a 96-well microplate luminometer (Model Lucy 1, Rosys Anthos, Austria) and pretreated with human modified oxidized lipoprotein ± L-phe as described above. Media were removed and replaced by 100 μL Krebs solution (119 mM NaCl, 4.7 mM KCl, 1.5 mM MgSO₄, 2.5 mM CaCl₂, 25 mM NaHCO₃, 1.2 mM KH₂PO₄, 11 mM glucose and 100 μM L-arginine), pH 7.4, and were kept cold on ice. Immediately before recording chemiluminescence, NADPH (final concentration 100 μmol·L⁻¹) and lucigenin, bis-N-methylacridinium nitrate (10 μmol·L⁻¹) was added to tissues and superoxide dismutase (SOD, 200 units·mL⁻¹) was used as a positive control. Light emission was recorded as mean arbitrary light units/cycle over 60 min.

**Data analysis**

SPR data were analysed using the curve fitting software Origin 7.0 (OriginLab Corporation, Northampton, MA, USA) and Bioevaluation software to determine the kₘ and kₜr rate constants and binding parameters, using both first and second order kinetic models. Bₘₐₓ calculations were normalized for surface density when this differed between experiments. A global fitting approach using the Bioevaluation software was not adequate to fully describe and fit the binding curves. Therefore, individual curve fitting was conducted in order to calculate binding parameters and rate constants (Supporting Information Fig. S2). The representative data shown in the results (Figure 2) were best described using first-order kinetics; hence the values were determined using monophasic fits.

For HPLC based assays, data are presented as mean ± SEM (where n = number of animals). One-way analysis of variance was used to analyse data obtained from HPLC-based assays measuring BH₄, bioperin, nitrite and superoxide anion levels in vascular tissues, plasma, cells and media. For the real-time kinetic assays, data are presented as mean ± SEM (where n = number of cell pellets from individually grown cultures). Two-way analysis of variance was used to analyse activity data from kinetic assays. GraphPad Prism 5 (GraphPad Software Inc, La Jolla, CA, USA) was employed to analyse all assay data.

**Materials**

Bacterial culture reagents, plasmid vectors and competent cells were purchased from Novagen, VWR International. Oligonucleotide primers were synthesized by Sigma-Aldrich (Dorset, UK). All other reagents were purchased from Sigma-Aldrich unless otherwise stated.

**Results**

**Expression and activity of native/full length and T-GCH1 with GFRP**

Soluble human recombinant T-GCH1, F-GCH1 and GFRP proteins were successfully expressed individually in BL21 (GCH1) or Rosetta (GFRP) cells (Figure 1A). In dual expression cultures, metal affinity purification of His₅-T-GCH1 or Hisᵢ-F-GCH1 revealed that GFRP was able to bind to both forms of GCH1 and could be co-isolated (Figure 1B).

Western blotting using specific antibodies revealed immunoreactive bands for GFRP and all His-tagged proteins (Figure 1C, i and ii). The GCH1 antibody (which recognizes an N-terminal epitope) correctly identified native but not T-GCH1 (Figure 1C, iii), confirming an intact N-terminal region on native GCH1 and demonstrating that the two forms of GCH1 could be readily distinguished from one another.

Activity assays and native PAGE gels run immediately before SPR analysis, demonstrating that proteins were predominantly in an oligomeric form (Figure 1D–F) (Maita et al., 2001). Consistent with previous reports (Higgins and Gross, 1986; Starr et al., 2014), F-GCH1 displayed lower levels of activity compared to T-GCH1 (Figure 1E), an effect that was reversed when F-GCH1 was co-incubated with GFRP (Figure 1F).

**Protein–protein interactions between native or T-GCH1 with GFRP**

Representative binding profiles illustrate differences between GFRP binding to F-GCH1 (Figure 2A, left) or T-GCH1 (Figure 2A, right). Binding parameters and determined kₘ and kₜr values are listed in the associated table. Binding was observed between both forms of GCH1 with GFRP, in the absence of additional L-phe, BH₄, or GTP. The analysis yielded Kᵢ values of 8 nM for F-GCH1 : GFRP and 17 nM for T-GCH1 : GFRP in the absence of ligands (tabulated in Figure 2A).

**Effects of L-phe on F-GCH1–GFRP protein–protein interactions**

L-phe changed both the association and dissociation rate constants with both forms of GCH1 and GFRP resulting in an eight-fold increase in binding affinity for F-GCH1–GFRP interactions: 1 nM Kᵢ, and 10-fold increase in binding affinity.
Figure 1
Expression and activity of human recombinant native/full length GCH1 (F-GCH1), truncated GCH1 (T-GCH1) and GTP cyclohydrolase 1 (GCH1) feedback regulatory protein (GFRP). (A) F-GCH1 expression in uninduced (lane 1) and IPTG induced (lane 2) cells. T-GCH1 expression from IPTG induced (lanes 3 and 4) cells. GFRP expression in uninduced (lane 5) and IPTG (isopropyl β-D-1-thiogalactopyranoside) induced (lanes 6 and 7) cells. Products were resolved on a 4–15% SDS-PAGE gradient gel. (B) Elution of purified proteins from dual expression cultures: His-T-GCH1 and GFRP bands are observed at ∼25 and ∼10 kDa, respectively; His-F-GCH1–GFRP bands observed at ∼28 and ∼10 kDa respectively. (C) (i): Immunoreactive bands for GFRP at ∼10 kDa were observed using a commercially available GFRP antibody; (ii) immunoreactive bands for both GFRP and T-GCH1 using a commercially available His-tag antibody were observed at ∼10 and ∼24 kDa, respectively; (iii) immunoreactive band at ∼28 kDa for F-GCH1 using an N-terminal GCH1 antibody; no immunoreactive bands for T-GCH1. (D) Native 5% PAGE gels run prior to surface plasmon resonance (SPR) analysis. Predominant bands highlighted for F-GCH1, T-GCH1 and GFRP as indicated, from two independent experiments. (E) Activity of purified F-GCH1 and T-GCH1 was quantified by HPLC detecting neopterin production; mouse liver homogenate and empty vector control were used for comparison (n = 6, mean ± SEM). *P < 0.05, ***P < 0.001, significantly different from control. (F) GCH1 activity measured by microplate assay in the presence of 250 nM GCH1, 1 μM GFRP and 100 μM GTP (n = 10, mean ± SEM). ***P < 0.01, significantly different from F-GCH1 without GFRP).
Figure 2
Surface plasmon resonance sensorgrams and tabulated data for His-tag captured native/full length GCH1 (F-GCH1) or truncated GCH1 (T-GCH1) interacting with GCH1 feedback regulatory protein (GFRP) analyte in the absence and presence of L-phenylalanin (L-phe). (A) Representative sensorgrams comparing F-GCH1–GFRP binding curves (left) and T-GCH1–GFRP binding curves (right). GTP-cyclohydrolase 1 (GCH1) is captured at a surface density of $\sim 1300 \text{ RU}$ via His-capture, and GFRP is introduced in varying concentrations (12.5–400 nM) with tabulated first-order dissociation constants ($K_D$) and on- and off-rate constants ($k_{on}$, $k_{off}$) ($n = 4$). (B) Representative sensorgrams for GFRP binding to F-GCH1 in the presence of L-phe. GCH1 immobilized at a surface density of $\sim 1500 \text{ RU}$ via His-capture, and GFRP and L-phe are introduced in varying concentrations (12.5–400 nM), with tabulated first-order dissociation constants ($K_D$) and on- and off-rate constants are tabulated ($n = 4$). (C) Comparison of binding kinetics; binding profiles for T-GCH1 and F-GCH1 in the presence and absence of ligands with 400 nM GFRP; F-GCH1 + GFRP; T-GCH1 + GFRP; F-GCH1 + GFRP + L-phe.
for T-GCH1–GFRP interactions: 1.7 nM \(K_d\). Representative sensorgrams for F-GCH1–GFRP + L-Phe (Figure 2B); T-GCH1–GFRP + L-Phe (Supporting Information Fig. S1) and corresponding values for F-GCH1 and T-GCH1 are summarized in the associated table (Figure 2B). In addition to these findings, a clear, two-fold rise in maximal binding (\(B_{max}\)) was observed in F-GCH1–GFRP interactions in the presence of L-phe. A comparison of SPR derived data and binding kinetics are summarized in Figure 2C.

The effect of BH\(_4\) (20 \(\mu\)M) on GCH1–GFRP interactions was also investigated, revealing increased binding affinity in its presence but no distinguishable differences between T-GCH1–GFRP and F-GCH1–GFRP interactions (Supporting Information Table S1).

**Effects of L-phe challenge on Ox-LDL pre-treated endothelial cells**

Addition of human modified oxidized lipoprotein to endothelial cells led to a significant elevation of superoxide anion (Figure 3A), reduction in nitrite (Figure 3B) and reduction in BH\(_4\) (Figure 3C) compared to baseline and consistent with published observations (Bowers et al., 2011). Application of L-phe (500 \(\mu\)M, 30 min) restored superoxide, nitrite and BH\(_4\) towards baseline levels (Figure 3A–C).

**Effects of oral L-phe challenge on biopterin and BH\(_4\) levels in vivo**

Oral L-phe challenge (100 mg·kg\(^{-1}\) bolus) in C57BL/6 mice led to a peak plasma L-phe concentration at 20 min which normalized to baseline by 4 h (Figure 4A). Similarly, the peak increase in plasma biopterin was rapid and normalized by 8 h (Figure 4B). In contrast, biopterin in aortic tissue followed a different profile, showing a more gradual and continual rise over the 8 h period (Figure 4C). Importantly, the functionally important pterin, BH\(_4\) (which behaves as an NOS cofactor and has vaso-protective properties), was also significantly elevated in aorta 4 h after administration of L-phe, normalizing by 8 h (Figure 4D). Finally, whilst administration of L-phe to GCH1 wild-type littermates stimulated biopterin/BH\(_4\) production in a similar manner to that observed in commercially purchased C57BL/6 mice (Figure 4B, E and F), L-phe had no significant effect in mice lacking endothelial GCH1 [GCH1(fl/fl)-Tie2Cre] (Figure 4E and F). These data suggest that L-phe stimulates endothelial GCH1, leading to a rise in BH\(_4\) in the aorta.

**Discussion and conclusion**

**Protein interactions of native and truncated GCH1 with GFRP**

Published structural studies of the GCH1–GFRP complex have used a truncated form of mammalian GCH1, lacking a large portion of the N-terminal region, suggesting that this region had no influence on either activity or binding to GFRP (Auerbach et al., 2000; Maita et al., 2002, 2004). This suggestion has, however, been challenged, as the N-terminal region has been shown to modulate GCH1 activity and to be essential for GFRP binding (Swick and Kapatos, 2006; Higgins and Gross, 2011). Furthermore, previous studies have suggested that the known ligands, L-phe and BH\(_4\), or substrate, GTP, are an essential requirement for GCH1–GFRP binding (Harada et al., 1993; Yoneyama and Hatakeyama, 1998).

In contrast to these observations, using SPR, we have shown that T-GCH1 and F-GCH1 are able to bind to GFRP with nanomolar affinity, in the absence of known ligands and substrate. Furthermore, whilst the N-terminal region modestly enhanced the affinity of interaction with GFRP, high
affinity interactions were still observed between T-GCH1 and GFRP. This ability of both native and truncated forms of GCH1 to interact with GFRP was independently confirmed using a GCH1–GFRP dual expression plasmid, where untagged GFRP was co-isolated with both His6-F-GCH1 and His6-T-GCH1. These findings are inconsistent with yeast two hybrid studies, in which N-terminal deletion diminished GCH1–GFRP interactions by 80% (Swick and Kapatos, 2006) or where the truncated enzyme displayed a relative inability to engage in higher ordered complexes with His6-GFRP (Higgins and Gross, 2011). The reasons for this discrepancy probably reflect differences in protein expression methodologies and the enhanced sensitivity of SPR to detect protein–protein interactions. A noticeable difference in maximal binding (Bmax) between F-GCH1–GFRP and TGCH1–GFRP was, however, detected, suggesting altered stoichiometry between the native and truncated complexes – a finding that requires further investigation but beyond the scope of the current study. Furthermore, absence of the N-terminal region conferred higher measured activity consistent with published observations (Higgins and Gross, 2011). It has been suggested that the N-terminal region may exert an auto-inhibitory effect leading to the observed activity changes mediated via a direct peptide bond, rather than a transient obstruction of the active site by a mobile N-terminal region (Higgins and Gross, 2011). Furthermore, instability of the full-length enzyme during purification has been reported by several groups (Yim and Brown, 1976; Auerbach et al., 2000), and it may equally be feasible that the enhanced activity of T-GCH1 can be, in part, attributed to greater T-GCH1 stability compared to F-GCH1 following purification. Together, our in vitro and biophysical data indicate that the GCH1 N-terminal region is not essential for GFRP binding, but that GCH1–GFRP binding kinetics and activity are altered when the N-terminal region is deleted – indicating a functionally important role for this region.

Figure 4
Effects of oral L-phe challenge on systemic and vascular biopterin levels in wild-type mice and GCH1(fl/fl)-Tie2Cre (KO) mice. (A) Plasma L-phenylalanine, (B) plasma total biopterins, (C) aortic total biopterins and (D) aortic 6R-L-erythro-5,6,7,8-tetrahydrobiopterin (BH4) levels detected by HPLC, over an 8 h time course, following 100 mg·kg⁻¹ oral L-phe challenge in wild-type mice. n = 6–12 for plasma and n = 4–8 aorta, mean ± SEM. ***P < 0.001, **P < 0.01, *P < 0.05: significantly different from saline control. (E) Aortic total biopterins and (F) aortic BH4 levels in GCH1(fl/fl)-Tie2Cre (KO) mice and wild-type littermates, 1 and 4 h after 100 mg·kg⁻¹ oral L-phe challenge. n = 6–8, mean ± SEM. **P < 0.01, significantly different from WT saline.
**Effects of L-phe on GCH1–GFRP interactions**

The interactions between GFRP and both forms of GCH1 were subsequently quantified in the presence of the allosteric effector molecule, L-phe. Whilst both forms of GCH1 were able to bind to GFRP in its absence, L-phe changed both the association and dissociation rate constants between both forms of GCH1 and GFRP, resulting in an 8- to 10-fold increase in binding affinity – an effect that was mimicked by another allosteric modulator, BH₄. The increase in binding affinity induced by L-phe between the truncated and native forms of GCH1 with GFRP was indistinguishable. These changes in binding affinity are consistent with the structural changes reported in the stimulatory crystal structure of the rodent T-GCH1–GFRP complex (Maita et al., 2002).

For the physiologically expressed form (F-GCH1) a clear two-fold rise in maximal binding (B_max) values was also observed upon the addition of L-phe, indicating a different stoichiometry to that previously suggested (Maita et al., 2002). This unexpected observation is, however, consistent with a previous report that native GCH1 binding to GFRP generates a very high molecular weight band which exceeds the size that would have been predicted for a GCH1 homodecamer bound to a GFRP pentamer (Higgins and Gross, 2011). However, further studies are required to fully understand this observation.

We thus demonstrate that SPR can readily detect the changes in GCH1–GFRP protein interactions, induced by small effector molecules such as L-phe. The very slow dissociation rates between GCH1 and GFRP observed using SPR indicate that within a physiological context, the two proteins would likely remain tightly complexed to one another.

**L-phe challenge on BH₄, nitric oxide and superoxide anion levels in vitro**

We used an *in vitro* model of endothelial dysfunction and eNOS ‘uncoupling’ (Xie et al., 2012) to establish the effects of L-phe on BH₄ and nitrite levels. Consistent with previous reports, incubation of endothelial cells with human oxidized lipoprotein led to a significant elevation of superoxide anion levels and concomitant reduction in nitrite and BH₄. Incubation with L-phe reversed this effect, indicating that stimulation of endothelial GCH1 by L-phe analogues has the potential to reverse endothelial dysfunction.

**L-phe challenge on biopterin and BH₄ levels in vivo**

A single 100 mg·kg⁻¹ L-phe oral dose led to a significant increase in plasma and aortic biopterin/BH₄ levels, in all wild-type mice – an effect that was not observed in aortas from GCH1(fl/fl)-Tie2Cre mice. Based on published evidence and data obtained in this study, we conclude that the raised biopterin/BH₄ levels following oral L-phe administration occur via activation of the GCH1 : GFRP complex in endothelial cells (Harada et al., 1993; Saunders-Pullman et al., 2004).

Interestingly, aortic BH₄ was still detectable in mice lacking endothelial GCH1, suggesting that cell types such as smooth muscle or adventitial fibroblasts may also generate BH₄. Our findings are in agreement with the initial characterization of these genetically modified mice, which demonstrated that endothelial GCH1 gene deletion or endothelial denudation in wild-type mice reduced, but did not abolish, aortic BH₄ levels (Chuaiphichai et al., 2014). Whilst we were unable to show a statistically significant difference in basal BH₄ levels within aortic tissue between GCH1(fl/fl)-Tie2Cre mice and WT littermates, we did observe a trend reduction in the baseline state – the differences observed between this study and that of Chuaiphichai et al. (2014) are likely to reflect subtle differences in tissue dissection and preparation.

Importantly, in the present study, L-phe administration did not stimulate a significant rise in aortic BH₄ in GCH1(fl/fl)-Tie2Cre mice, indicating that the GCH1–GFRP complex is primarily located in endothelial cells. As such, novel therapies activating the GCH1–GFRP axis are most likely to target this cell type. Indeed, the vascular endothelium is believed to be the primary source of BH₄ (d’Uscio and Katusic, 2006), and previous studies have demonstrated that changes in GCH1–GFRP interactions are a critical regulator of BH₄ and NO biosynthesis in endothelial cells, in response to laminar shear stress (Li et al., 2010).

The profiles of plasma L-phe and biopterin in mice mirrored that previously observed in humans challenged with oral L-phe (Saunders-Pullman et al., 2004), indicating that this is a suitable and clinically translatable model with which to investigate the GCH1–GFRP axis.

The observation that L-phe challenge stimulated aortic BH₄ levels for at least 4 h is encouraging, as it suggests that the activation of the GCH1–GFRP complex elicits a sustained elevation of vaso-protective BH₄ in target vascular tissues. These *in vitro* and *in vivo* findings thus provide mechanistic evidence to support published functional studies that have shown that L-phe administration restores endothelial function and attenuates the observed hypertension induced by administration of the GCH1 inhibitor di-amino-hydroxy.pyrimidine (Mitchell et al., 2004).

In conclusion, we have undertaken a detailed analysis of GCH1 and GFRP using complementary *in vitro* biophysical analysis with *in vitro* and *in vivo* murine studies. We have successfully expressed soluble human GCH1 and GFRP and, for the first time, quantified the binding rate constants between GCH1 and GFRP, using SPR. We have also demonstrated that the N-terminal region of GCH1 is not essential for GFRP to interact, but that deletion of this region alters the binding kinetics between the two proteins. Whilst GCH1 and GFRP were able to bind in the absence of known ligands, the presence of L-phe substantially elevated the maximal binding and the affinity of interaction – suggesting that, in an *in vivo* system (where ligands and substrate would be circulating), the two proteins would display high affinity interactions. Indeed, the rapid rise in plasma biopterin (observed within 20 min) coupled with slow GCH1–GFRP dissociation rates (obtained by SPR) supports the view that GCH1 and GFRP are likely to be constitutively bound *in vivo*, rather than binding in response to an acute elevation of circulating L-phe, following dietary intake.

Our biophysical and *in vivo* data suggest that the L-phe binding pockets on the GCH1–GFRP complex represent a rational drug target to raise vascular BH₄ for the treatment of endothelial dysfunction. It is important to note that L-phe itself is not a feasible chronic therapeutic intervention due to its diverse physiological functions and role in catecholamine biosynthesis. However, given the marked allosteric changes...
induced by L-phe, low MW small molecule mimetics that alter interactions between GCH1 and GFRP in a similar manner have the potential to regulate intracellular BH4 for therapeutic purposes. Indeed, the sustained effect on aortic BH4 levels following a single oral L-phe challenge is highly encouraging from a therapeutic standpoint as this could elevate endothelial NO levels and/or limit damaging reactive oxygen species, circumventing the limited bioavailability/efficacy of oral BH4. Such an agent would have use in a wide spectrum of cardiovascular diseases, underpinned by reduced NO bioavailability and/or enhanced oxidative stress.

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Author contributions

M. N. designed the research study. D. H., A. S. and L. H. performed the research. E. M. and K. M. C. provided access to and advice on GCH1(fl/fl)-Tie2Cre mice. P. R. B. directed all cloning and protein expression studies. B. J. S. and J. M. M. directed and contributed to analysis of all SPR data. D. H. and M. N. wrote the manuscript. All authors contributed to editorial changes in the manuscript.

Conflict of interest

The authors declare no conflict of interest.

References


Supporting information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site: http://dx.doi.org/10.1111/bph.13202

Figure S1 Binding profile for truncated GCH1 (T-GCH1) with GCH1 feedback regulatory protein (GFRP) in the presence of L-phenylalanine (L-phe). T-GCH1 is immobilized at a surface density of ∼1500 RU via His-capture, and GFRP and L-phe are introduced in varying concentrations (400–25 nM) (n = 4 for each experiment at varying immobilization patterns and surface densities). Representative sensorgrams are shown.

Figure S2 SPR data analysis by Origin software, curve fittings and residuals.

Table S1 Summary of the binding kinetics for both native/full-length GCH1 (F-GCH1) and GCH1 feedback regulatory protein (GFRP) as well as truncated GCH1 (T-GCH1) and GFRP in the presence and absence of BH.