The sensitivity of a radical pair compass magnetoreceptor can be significantly amplified by radical scavengers

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Birds have a remarkable ability to obtain navigational information from the Earth’s magnetic field. The primary detection mechanism of this compass sense is uncertain but appears to involve the quantum spin dynamics of radical pairs formed transiently in cryptochrome proteins. We propose here a new version of the current model in which spin-selective recombination of the radical pair is not essential. One of the two radicals is imagined to react with a paramagnetic scavenger via spin-selective electron transfer. By means of simulations of the spin dynamics of cryptochrome-inspired radical pairs, we show that the new scheme offers two clear and important benefits. The sensitivity to a 50 μT magnetic field is greatly enhanced and, unlike the current model, the radicals can be more than 2 nm apart in the magnetoreceptor protein. The latter means that animal cryptochromes that have a tetrad (rather than a triad) of tryptophan electron donors can still be expected to be viable as magnetic compass sensors. Lifting the restriction on the rate of the spin-selective recombination reaction also means that the detrimental effects of inter-radical exchange and dipolar interactions can be minimised by placing the radicals much further apart than in the current model.

Magnetoreception — the ability to sense magnetic fields — is widespread throughout the animal kingdom, but the underlying detection mechanisms are far from clear. There are two main hypotheses. One involves single-domain, ferrimagnetic or superparamagnetic iron-containing particles that are caused to move by their interaction with the Earth’s magnetic field and so influence the gating of mechano-sensitive or force-gated ion channels. The other is based on radical pairs formed by photo-induced electron transfer reactions in sensor proteins. The magnetic sensitivity arises from a combination of the coherent quantum spin dynamics and the spin-selective reactivity of a pair of spin-correlated radicals, which cause the yield of a signalling state to depend on the intensity and direction of the external magnetic field.

The radical pair mechanism is “quantum” not just in the trivial sense that it involves a non-classical property (spin angular momentum), but more interestingly because quantum coherences play an essential role. This aspect was highlighted by a recent suggestion that the angular precision of the magnetic compass in migratory birds can be understood in terms of avoided crossings of spin energy-levels in the radicals. Another quantum feature of the mechanism is that the electron spins in the radical pair are initially entangled although there is currently no reason to think that entanglement, as distinct from coherence, is necessary for the operation of the compass. It is also striking that the performance of a radical pair sensor may be enhanced by interactions with its fluctuating environment, a property apparently found in other areas of “quantum biology.”

Experimental and theoretical support for a radical pair mechanism of magnetoreception is accumulating (reviewed in ref. 2), in particular in the context of the avian magnetic compass. Magnetically sensitive radical pairs are thought to be formed by light-dependent electron transfer reactions in cryptochromes — blue-light photoreceptor flavoproteins — located in the retina. In vitro, purified cryptochrome from the plant Arabidopsis thaliana (AtCry1) and a closely related protein, E. coli photolyase (ECP1), both show light-dependent responses to weak magnetic fields (~1 mT). In these proteins, photo-excitation of the non-covalently bound flavin...
adenine dinucleotide (FAD) cofactor leads to the formation of radical pairs via sequential electron transfers along the “tryptophan-triad”, a chain of three conserved tryptophan residues within the protein25-28 (Fig. 1). This process reduces the photo-excited singlet state of the FAD to the anion radical, FAD•−, and oxidises the terminal, surface-exposed, tryptophan (Trp H) to give the cation radical, Trp H•+. Formed with conservation of spin angular momentum, the radical pair is initially in an electronic singlet state, [FAD•−Trp H•+]22, 23, 27. This form of the protein is a coherent superposition of the eigenstates of the spin Hamiltonian which comprises the Zeeman, hyperfine, exchange and dipolar interactions of the electron spins. As a consequence, the radical pairs oscillate coherently between the singlet and triplet states, a process that manifests itself in the yields of subsequent spin-selective reactions of the radicals. In particular, when the protein is immobilized, the anisotropy of the electron-nuclear hyperfine interactions causes the reaction product yields to depend on the orientation of the protein with respect to an external magnetic field.

Figure 1. Electron transfer pathway in cryptochromes. After photo-excitation of the FAD cofactor, three or four rapid sequential electron transfers along a triad or tetrad of tryptophan residues (Wα, Wβ, Wγ, Wδ) generate a spin-correlated radical pair [FAD•−TrpH•+] or [FAD•−TrpH•+]. The figure is based on the crystal structure of DmCry (PDB ID: 4GUS)25, 36.

Following Ritz et al.9, most authors (e.g. refs 14, 28–33) have envisaged spin-selective reactions of both singlet and triplet radical pairs. The latter requires there to be a triplet product state that is energetically accessible from the radical pair. As no such species exists in cryptochrome, we base our treatment here on the more plausible scheme shown in Fig. 2a which we henceforth refer to as the “current” model22, 23, 34. In Fig. 2a, the spin-selective reaction channel is charge recombination (rate constant kR) within the singlet state of the radical pair (RP) which regenerates the ground state of the protein, G. As found experimentally for the flavin-tryptophan radical pair in AcCry and EcPL25, 26, the RP state also undergoes proton transfer reactions, which occur with equal rate constants (kPT) for singlet and triplet pairs, to produce a secondary, long-lived radical pair state, S. In vivo, S is assumed to be, or to lead to, the biochemical signalling state of the protein2. The interaction of the electron spins with the magnetic field can induce a significant change in the yield of S if kPT1 (the characteristic time of singlet recombination) is comparable to or shorter than kPT−1 (the time required for the formation of S), small compared to the electron spin relaxation time (≈1 µs or possibly longer) and longer than the coherent singlet-triplet interconversion time. These conditions mean that the radicals must not be too far apart, otherwise charge recombination will be too slow. This restriction is satisfied for the EcPL and AcCry1, in which the edge-to-edge separation of the aromatic rings of FAD and Trp H is ≈1.47 nm35–37. In the cryptochrome from the fruit fly, Drosophila melanogaster (DmCry), however, there is an additional electron donor, Trp H beyond Trp H (Fig. 1)38. The edge-to-edge distance between FAD and TrpH in DmCry is 1.70 nm35, 36 which is large enough that direct charge recombination in [FAD•−Trp H•+] cannot compete effectively with electron spin relaxation, at least for the purified protein in vitro, explaining the weak magnetic field effects observed for DmCry19. Sequence alignments suggest that avian cryptochromes also have a fourth tryptophan which could be involved in radical pair formation; we return to this point below.

All of the above has been gleaned from spectroscopic observations of purified cryptochromes in vitro22, 23, 39, 40. The same proteins may behave differently in a cellular environment as a result of their interactions with (for example) signalling partners and whatever structures immobilise and align them as direction sensors. No studies of magnetic field effects have yet been reported for any of the four avian cryptochromes. There is therefore scope to speculate about alternative radical pairs that might undergo different photo-reactions to those in Fig. 2a41.

Here we describe a modified reaction scheme in which the magnetic compass sensitivity is enhanced by a spin-selective reaction of one of the constituents of the radical pair with an additional paramagnetic molecule. We refer to this molecule as a ‘scavenger’, a term defined as “a substance that reacts with (or otherwise removes) a trace component [...] or traps a reactive reaction intermediate”42. The scavenging process can lead to large magnetic field effects, even in the limit of very slow charge recombination (kPT → 0) where the current model (Fig. 2a) predicts no magnetic field effects at all. Such a reaction scheme could allow radical pairs with separations much larger than 2 nm to operate as efficient magnetic compass sensors including, but not restricted to, those containing a fourth tryptophan residue in the electron transfer chain.

Spin-selective scavenging. We start by showing how a spin-conserving reaction with a paramagnetic scavenger can affect the coherent spin dynamics of a radical pair. Our treatment is a generalization of Letuta and Berdinskii43. We consider a pair of radicals (A and B) in a spin-correlated singlet state and determine how the spin state changes when one of the radicals (A) undergoes spin-selective reactions with a paramagnetic scavenger C. Molecules A, B and C have spin quantum numbers, S, equal to ½, ½, and J ≥ ½, respectively. The scavenging
reactions convert A into a diamagnetic species (S = 0), for example by the transfer of an electron to or from C. We focus on organic molecules with weak spin-orbit coupling such that the reactions conform to Wigner’s spin conservation rule; as a consequence, the spin of C must change by ±½. Thus, in general, there are two parallel, spin-allowed reactions with distinct rate constants, \( k_C^+ \) and \( k_C^- \), and distinct products:

\[
\begin{align*}
2A + 2j+1C & \quad \xrightarrow{k_C^+} A + 2j+2C \\
2A + 2j+1C & \quad \xrightarrow{k_C^-} A + 2jC 
\end{align*}
\]

(the superscripts are the spin multiplicities, 2S + 1).

In the combined Hilbert spin-space of A, B and C, these reactions are governed by projection operators that can be written in terms of the spin angular momentum operators, \( \hat{S}_K \) (see Supporting Information, Section S1):

\[
\hat{P}_{AC}^\pm = \frac{1}{2} \pm \frac{1}{2j+1} \left[ \frac{1}{2} + 2\hat{S}_A \cdot \hat{S}_C \right].
\]

where \( \hat{P}_{AB}^+ + \hat{P}_{AC}^- = \hat{1} \) and \( (\hat{P}_{AC}^+)^2 = \hat{P}_{AC}^- \). Following Haberkorn, the reactions give rise to the following equation of motion for the (concentration-weighted) density operator of the \([A,B,C]\) spin system:

\[
\frac{d}{dt}\hat{\rho}(t) = -\frac{i}{\hbar} [\hat{H}_{AB}, \hat{\rho}(t)]_{\pm} - \frac{1}{2}k_C^- [\hat{P}_{AC}^+, \hat{\rho}(t)]_{\pm},
\]

where \([.,.\)]_{\pm} denotes the anti-commutator.

To see most clearly the effect these reactions have on the surviving AB radical pairs, we temporarily ignore all spin interactions and all other reaction steps. We assume that the radical pair is initially in a singlet state, specified by the projection operator \( \hat{P}_{AB}^S \) and that C has no initial spin-correlation with A or B:

\[
\hat{\rho}(0) = \hat{P}_{AB}^S / \text{Tr} \left[ \hat{P}_{AB}^S \right] = \left( \frac{1}{4} - \hat{S}_A \cdot \hat{S}_B \right) / \text{Tr} \left[ \hat{P}_{AB}^S \right].
\]
Using equations (3)–(5), the fraction of the AB radical pairs that remain unreacted at time $t$ is the sum of two exponential decays (see Supporting Information, Section S2):

$$s(t) = \text{Tr}[\hat{\rho}(t)] = \frac{J + 1}{2J + 1} \exp(-\frac{1}{2}k_{c}t) + \frac{J - 1}{2J + 1} \exp(-\frac{1}{2}k_{c}t),$$

in which the two terms are the probabilities of the $S = J \pm \frac{1}{2}$ coupled angular momentum states of the AC subsystem.

We now make the simplifying assumption that, for energetic reasons, the spin-selective reaction that produces C in its higher spin state ($S = J + \frac{1}{2}$) is negligibly slow, i.e. $k_{c} = 0$. For example, if C in equation (2) is a doublet (i.e. a radical, with $J = \frac{1}{2}$), it can end up as either a singlet ($^1C, S = J = \frac{1}{2}$) or a triplet ($^3C, S = J + \frac{1}{2} = 1$). For organic molecules, the former is normally the ground state and the latter an excited state. Similarly if C is a triplet ($J = 1$), we consider only the formation of the doublet product, $^1C$, and exclude the higher energy quartet product, $^3C$. With this simplification, therefore, a fraction

$$s(t \to \infty)_{k_{c}=0} = \frac{J + 1}{2J + 1}$$

of the AB pairs is unreactive. In general, the probability that the surviving AB pairs are still in a singlet state at time $t$ is (see Supporting Information, Section S2):

$$P_{AB}^S(t) = \frac{\text{Tr} \left[ \rho^S_{AB} \hat{\rho}(t) \right]}{\text{Tr}[\hat{\rho}(t)]} = \frac{\left( \frac{J + 1}{2J + 1} \exp\left( -\frac{1}{2}k_{c}t \right) + \frac{J + 1}{2J + 1} \exp\left( -\frac{1}{2}k_{c}t \right) \right)^2}{s(t)},$$

so that in the limit of exclusive formation of the $S = J - \frac{1}{2}$ scavenging product:

$$P_{AB}^S(t \to \infty)_{k_{c}=0} = \frac{J + 1}{2J + 1}.$$

It is clear from equations (8) and (9) that the scavenging reaction has converted a portion of the surviving radical pairs into the triplet state. When C is a doublet ($J = \frac{1}{2}$), 3/4 of the AB pairs survive at $t \to \infty$ and 3/4 of these survivors are singlets (in the AB-manifold). Similarly, if C is a triplet ($J = 1$), 2/3 of the radical pairs survive and 2/3 of them are singlets. If C is a singlet ($J = 0$, i.e. not paramagnetic) there are no spin restrictions on the reaction, all A radicals react with C and there are no radical pairs left at $t \to \infty$. Figure 3 shows $s(t)$ and $P_{AB}^S(t)$ for scavengers with different spin quantum numbers, $J$.

Additional insight into the origin of singlet-triplet interconversion in AB as a result of the spin-selective AC scavenging reaction is presented in the Appendix (Supporting Information).

**Simulation methods.** The cryptochrome photocycle in Fig. 2a has been modified to include a scavenging reaction (Fig. 2b). We restrict the discussion to a scavenger with spin $J = \frac{1}{2}$; preliminary calculations show that qualitatively similar effects can be expected when $J > \frac{1}{2}$. The states of the protein, G (ground state), RP (magnetically sensitive radical pair) and S (signalling state) are unchanged. To these is added a fourth state, X, formed by a spin-selective reaction of the FAD$^{\cdot+}$ radical in the RP state with a scavenger radical C. From equation (2), with $J = \frac{1}{2}$, X comprises the fully oxidised, diamagnetic flavin molecule and the tryptophan radical. The other product of the scavenging reaction is C$^{\cdot+}$, a diamagnetic form of the scavenger. Both S and X eventually return to the ground state, G. Although the primary radical pair in Fig. 2b is shown as containing WH$^{\cdot+}$, this radical could be further removed from FAD$^{\cdot+}$ than is the terminal tryptophan of the triad/tetrad in cryptochrome and it need not be a tryptophan radical. In principle, the scavenging reaction could also involve this radical instead of the FAD$^{\cdot+}$.

The key quantities we wish to calculate are the quantum yields of the reaction intermediates, S and X, as a function of the direction of an Earth-strength magnetic field (50 $\mu$T). The complete equation of motion for the spin density operator is:

$$\frac{d}{dt} \hat{\rho}(t, \Omega) = -i[\hat{H}(\Omega), \hat{\rho}(t, \Omega)] + \frac{\hat{K}}{\hbar} \hat{\rho}(t, \Omega).$$

Here, $\hat{H}(\Omega)$ is the spin Hamiltonian which is the sum of the individual spin Hamiltonians, $\hat{H}_{k}(\Omega)$, of the three paramagnetic molecules A ($= \text{FAD}^{\cdot+}$), B ($= \text{WH}^{\cdot+}$) and C, and their electron-electron exchange and dipolar couplings. The $\hat{H}_{k}(\Omega)$ operators contain the Zeeman interactions with the external magnetic field and hyperfine interactions with surrounding nuclear spins (in angular frequency units):

$$\hat{H}_{k}(\Omega) = -\gamma_B \text{B}_k(\Omega) \cdot \text{S}_k + \sum_{j} N_k \text{A}_{kj} \cdot \text{I}_{kj}.$$

$\text{I}_{kj}$ and $\text{A}_{kj}$ are, respectively, the angular momentum operator and the hyperfine tensor of nuclear spin $j$ in radical $k$. The sum runs over all $N_k$ magnetic nuclei in radical K. $\Omega$ denotes the polar and azimuthal angles specifying the direction of the field in the coordinate frame of the protein. The Larmor frequency, $\nu_0$, is related to the strength of the external magnetic field by $2\pi\nu_0 = \gamma_B |\text{B}_k(\Omega)|$. When $|\text{B}_k(\Omega)| = 50 \mu$T, $\nu_0 = 1.4 $MHz. The geomagnetic field is weak enough that, for organic radicals, the differences in g-values of the three electron spins can safely be neglected.
The reactions of the radical pair are accounted for by the superoperator \( \hat{K} \) in equation (10). Specifically, \( \hat{K} \) describes the spin-selective scavenging reaction that forms \( X \) (equation (4)), the spin-independent formation of \( S \), and the charge recombination reaction of the singlet configuration of \( A \) and \( B \):

\[
\hat{K} = k_{C}^{+} \hat{P}_{C}^{+} + k_{C}^{-} \hat{P}_{C}^{-} + k_{B} \rho \Omega.
\]

As in the previous section, we ignore the scavenging reaction that would produce an excited triplet state of \( C^{-} \) (equation (1)). \( k_{C} \) is independent of the spin interactions which are all much smaller than the thermal energy \( k_{B} T \). For the singlet initial condition in equation (5), the yield of \( S \), once all radicals have reacted, is given by

\[
Y_{S}(\Omega) = k_{t} \int_{0}^{\infty} \text{Tr}[\rho(t, \Omega)]dt.
\]

We define two quantities as measures of the performance of the radical pair as a magnetic direction sensor: the absolute (\( \Delta_S \)) and the relative (\( \Gamma_S \)) anisotropy of the yield of the signalling state \( S \):

\[
\Delta_S = \max_{\Omega} \{Y_{S}(\Omega)\} - \min_{\Omega} \{Y_{S}(\Omega)\},
\]

\[
\Gamma_S = \frac{\Delta_S}{\langle Y_{S} \rangle} \quad \text{where} \quad \langle Y_{S} \rangle = \frac{1}{4\pi} \int Y_{S}(\Omega)d\Omega.
\]

It is not clear which of \( \Delta_S \) and \( \Gamma_S \) corresponds more closely to what the birds perceive when they take a magnetic compass bearing. We therefore present calculations of both in the following. The corresponding quantities, \( \Delta_X \) and \( \Gamma_X \), for the yield of the scavenging product \( X \) can be calculated in a similar fashion (see Supporting Information, Figs S1–S4).

**Results**

In the following, we explore the effect of scavenging reactions on \( \Delta_S \) and \( \Gamma_S \) using spin systems of progressively increasing complexity to mimic important aspects of the [FAD•− WH•+] radical pair. In every case, the hyperfine tensors and the relative orientation of the radicals were taken from ref. 11. The forward rate constant \( k_{f} \) was fixed at \( 0.1 \mu s^{-1} \), consistent with a recent study of spin relaxation in \( \text{AtCry} \), which suggested that magnetic field effects would be strongly attenuated if the radical pair lifetime exceeded \( \sim 10 \mu s \). The charge recombination rate constant, \( k_{b} \), was varied in the range \( (0, 10k_{f}) \) and its value reported as \( \phi = k_{b}/k_{f} \) (for purified \( \text{AtCry}^{23} \), \( \phi \) was found to be \( \sim 2 \)). The external magnetic field was \( 50 \mu T \) and spin relaxation was neglected. Unless stated otherwise, the scavenger reacts with the FAD• radical.

Our starting point is a model with just three \(^{14}\text{N}\) hyperfine interactions: the N5 and N10 nitrogens in FAD• and the N1 nitrogen in WH•. We begin with a diamagnetic scavenger \( (J = 0) \) to provide a basis for comparison.
with the more interesting \( J = \frac{1}{2} \) case. Figure 4a,b show the dependence of the anisotropic yield of \( S \) on the scavenging rate constant, \( k_c \), for several values of \( \phi \). The maximum relative anisotropy, \( \Gamma_S \), is found when \( k_c = 0 \) (no scavenging) and for \( k_l \) at the upper end of the range studied (\( \phi = 10 \), Fig. 4a). \( \Gamma_S \) decreases as \( \phi \) is reduced, and vanishes for \( \phi = 0 \), when charge recombination (\( RP \rightarrow G \)) ceases to compete with the forward reaction (\( RP \rightarrow S \)). Magnetic field effects are expected to be small when the radicals are so far apart that the direct conversion of \( RP \) to \( G \) is slow compared to typical spin relaxation times (for a 10 \( \mu s \) relaxation time, this distance is \( >1.7 \) nm\(^4\)). Both \( \Gamma_S \) and \( \Delta_S \) are strongly attenuated by fast scavenging (\( k_c > 2 \pi \nu_0 \approx 10 \mu s^{-1} \)), which allows insufficient time for the 50 \( \mu T \) magnetic field to affect the spin dynamics. The maximum \( \Delta_S \) (Fig. 4b) is observed in the absence of scavenging and for intermediate values of \( \phi \) (e.g. \( \Delta_S \approx 0.055 \) when \( \phi = 2 \)).

The situation changes dramatically when the scavenger is paramagnetic (\( J = \frac{1}{2} \)). Figure 4c,d show the behaviour of \( \Gamma_S \) and \( \Delta_S \) for the [FAD\(^+\) - WH\(^+\)] model under the same conditions as Fig. 4a,b (but note the logarithmic scale of the vertical axis in Fig. 4c). In this case, \( \Gamma_S \) has a 50 \( \mu T \) electron Zeeman interaction but no hyperfine interactions; more complex systems will be discussed below. The general features of Fig. 4c,d are as follows. (1) Scavenging rate constants well in excess of 10 \( \mu s^{-1} \) no longer abolish the magnetic field effect on the signalling state. This is a direct consequence of the unreactivity of the coupled high-spin state (\( S = 1 \)) of the flavin radical and the scavenger radical. (2) Large values of \( k_c \) lead to significant magnetic field effects even when \( k_l \) is small or zero. The scavenging reaction can therefore take over the role played by spin-selective recombination in the current model (Fig. 2a). (3) The scavenging reaction amplifies both \( \Gamma_S \) and \( \Delta_S \). For this simple model, relative anisotropies, \( \Gamma_S \), in excess of 100% are seen for scavenging rate constants in the range 1 to 1000 \( \mu s^{-1} \) (Fig. 4c). For \( k_l = 0 \), the maximum \( \Gamma_S \) (338%) occurs when \( k_c \approx 52 \mu s^{-1} \). While it is true that these huge anisotropies are accompanied by smaller mean reaction yields, \( \langle \gamma \rangle \), the maximum absolute anisotropies, \( \Delta_S \) (Fig. 4d), are still larger than those found with a diamagnetic scavenger or with no scavenger at all (Fig. 4b). Qualitatively similar effects under otherwise identical conditions are found when the scavenged radical is WH\(^+\) rather than FAD\(^+\) (see Supporting Information, Figs S1 and S4).

In view of the enhanced performance of this simple model system, it is important to know whether similar effects can be expected for other radicals. Figure 5 shows the dependence of \( \Gamma_S \) and \( \Delta_S \), on \( k_c \) and \( \phi \) for a radical pair, [FAD\(^+\) - Z\(^+\)], in which WH\(^+\) has been replaced by Z\(^+\), a radical with no hyperfine interactions. Radicals pairs of this type have been invoked to explain the reported disorientation of migratory birds exposed to Larmor-frequency magnetic fields\(^49\) and are expected to produce larger magnetic field effects than pairs in which both radicals have significant hyperfine interactions, e.g. [FAD\(^+\) - WH\(^+\)]\(^4\). Calculations were performed with N5, N10, H6, 3×H8α and one of the H3 protons in FAD\(^+\) and no hyperfine interactions in the paramagnetic scavenger C. The results (Fig. 5a,b) are qualitatively and quantitatively similar to those in Fig. 4c,d.
The next step was to test the sensitivity of $\Gamma_S$ and $\Delta_S$ to the presence of hyperfine interactions in the scavenging radical, C•. To do this we constructed two additional models using the 3-nucleus version of $[\text{FAD}^- \cdot \text{WH}^+]$ above (N5, N10 in FAD•−, N1 in WH•+). In the first, the scavenger was modelled on a freely diffusing ascorbic acid radical, which is characterised by a single dominant $^1$H hyperfine coupling and is known to be capable of oxidizing FAD•−. In the second, we assumed that the WH•+ radical is reduced by a second, uncorrelated FAD•−, which was again modelled by means of the N5 and N10 hyperfine interactions. The relative orientations of WH•+ and the FAD•− scavenger were chosen (arbitrarily) as those in the crystal structure of dimeric DmCry35,36. In both cases (Fig. 5c–f), the yield of the signalling state showed, once again, remarkably enhanced sensitivity to the direction of the external magnetic field.

Finally, we return to $[\text{FAD}^- \cdot \text{WH}^+]$ but now with several hyperfine interactions in both radicals. Figure 6a shows the dependence of $Y_S(\theta)$, the yield of the signalling state, on the direction of the field (θ) in the yz-plane of the flavin, for a model comprising N5, N10 and H6 in FAD•− and N1, H1, H4, Hβ and H7 in WH•+. In the absence of a spin-selective scavenging reaction (Fig. 6a, red line), there is a “spike” centred at $\theta = 90^\circ$ arising from avoided crossings of spin states with different singlet character. It has been suggested that such a spike could afford a much more precise compass bearing than a smoother, more gently varying $Y_S(\theta)$34 (see also ref. 34). In the presence of a scavenging reaction (Fig. 6a, black line), a much stronger spike is seen when the field is parallel to the z-axis of the flavin ($\theta = 0$; note the very different scales of the vertical axes). Figure 6b shows the dependence...
of the reaction yield is tiny. This is a consequence of the number and lack of symmetry of the hyperfine interactions in \( \text{WH}\) and \( \text{RP}\) have different protonation states (as indicated in Fig. 2b). For the sake of illustration, oxidation of flavin semiquinone radicals by \( \text{O}_2\) is \( 10^4\) times faster for the anionic radical than for the neutral radical. Clearly the details will depend on the identity and reactivity of the radical pair and the scavenger.

**Discussion Amplification.** The notion that magnetoreception could rely on quantum coherence in transient radical pairs has caught the imagination of scientists in a range of disciplines from zoology to theoretical physics. Although evidence in support of the hypothesis is accumulating, the sensor molecules have yet to be unequivocally identified. It is therefore appropriate to explore alternative photocycles. We have chosen to focus on cryptochrome because, 17 years after it was first proposed, it is still the only candidate magnetoreceptor for the avian compass and because FAD radicals seem to be near optimal as components of a direction sensor. A potential problem with the current model based on \([\text{FAD}^-\ \text{WH}^+]\) (Fig. 2a) is that the predicted anisotropy of the reaction yield is tiny. This is a consequence of the number and lack of symmetry of the hyperfine interactions in \( \text{WH}^+\) and \( \text{RP}\) have different protonation states (as indicated in Fig. 2b). For the sake of illustration, oxidation of flavin semiquinone radicals by \( \text{O}_2\) is \( 10^4\) times faster for the anionic radical than for the neutral radical. Clearly the details will depend on the identity and reactivity of the radical pair and the scavenger.

A recent study suggested that the anisotropic yield of the state \( \text{S}^+\) radical pair. (a) The yield of \( \text{S}^+\), \( Y_{\text{S}}\) as a function of the direction of the magnetic field in the \( \text{yz}\)-plane of the flavin for \( k_c = 0\), \( k_b = 2k_f\) (red line) and \( k_c = 24.2\ \mu\text{s}^{-1}\), \( k_b = 0\) (blue line). Note the very different vertical scales for the two traces. (b) The relative yield anisotropy, \( \Gamma_\text{X}\), as a function of \( \phi\) and \( \phi\). The spin system comprised N5, N10 and H6 in \( \text{FAD}^+\) and N1, H1, H4, H3 and H7 in \( \text{WH}^+\). The scavenger had no hyperfine interactions.
in $Y_\phi(\Omega)$, predicted for certain directions of the field, could improve the precision of the compass bearing\(^{31}\) (see also ref. 34). Nevertheless, $\Gamma_\phi$ is not expected to be much larger than $\sim1\%$. A [FAD$^+$• Z•$^-$] radical pair, with no magnetic nuclei in Z•$, could give much stronger signals\(^{41}\), although the only obvious Z•$^+$ radical (superoxide, O$_2$•$^-$) is almost certainly unsuitable due to its exceedingly fast spin relaxation\(^{32}\). Furthermore, spikes in $Y_\phi(\Omega)$ are not expected for a [FAD$^+$• Z•$^+$] pair\(^{11}\) meaning that the compass precision would be inferior to that of a radical pair with hyperfine interactions in both radicals.

In any case, amplification of the primary signal will be an essential feature of the biological compass\(^{84}\). This could occur within the sensor itself and/or in the course of signal transduction. A cyclic kinetic scheme for amplification of the primary effect has recently been suggested\(^{42}\). It relies on the effects of slow radical termination reactions on the photo-stationary state of the continuously illuminated protein. Although this process has been demonstrated experimentally for a purified cryptochrome\(^{46}\), it is not clear how well it would work for nocturnally migrating birds whose magnetic compass appears to function at very low light levels\(^{6,73}\). The amplification scheme suggested here (Fig. 2b) is practically acyclic and hence not subject to the same limitations. Furthermore, the scavenging process gives rise to remarkably large anisotropy in the reaction yields (more than 100% in all the models considered here), which would drastically reduce (by a factor of 10$^3$ to 10$^6$) the number of integrated events required to elicit a directional response with the required signal-to-noise ratio\(^{11}\). As in the current model\(^{11}\), there is a spike in the reaction yield which becomes narrower and more pronounced as the lifetime of the radical pair is increased (see Supporting Information, Fig. S6). In contrast to the current model, this feature occurs when the field is parallel ($\theta=0$) rather than perpendicular ($\theta=90^\circ$) to the normal to the plane of the flavin moiety. Such spikes offer the possibility of highly precise compass bearings\(^{11,34}\). Furthermore, the shape of the reaction yield anisotropy is to a large extent independent of the details of the scavenging process, including the hyperfine interactions in all three radicals, such that similar responses can be expected under a variety of conditions (see Supporting Information, Fig. S5).

**Distance constraints.** A spin-selective reaction channel is required in order that the effect of the magnetic field on the coherent spin motion can alter the yield of the reaction product. In the scheme shown in Fig. 2a, this is the charge recombination step. The rate constant of this reaction, $k_{rc}$, must be larger than or comparable to the electron spin relaxation rate, otherwise the reaction yield will simply reflect the statistical ratio of singlet and triplet states, which is essentially independent of the direction of a weak magnetic field. Assuming this reaction occurs in a single step (rather than by sequential electron hopping), this requirement precludes radical pairs with edge-to-edge distances greater than $\sim1.7$ nm. According to the empirical "Moser-Dutton ruler", electron transfer rate constants as small as 10$^3$ s$^{-1}$ are expected for an edge-to-edge distance of 2 nm, even in the Marcus activation-less limit\(^{49}\). Furthermore, an increase of 0.33 nm in the separation of the radicals is expected to result in a 100-fold decrease in the rate constant of charge recombination. In principle, this restriction on the separation of the radicals could be relaxed if charge recombination occurred indirectly, either by reversible electron hopping along the Trp-triad/tetrad or via a second, independent, electron transfer chain. Given the strongly exergonic nature of the forward electron transfer (reflecting the steadily increasing solvent-exposure of the radical pair as the charge is propagated along the electron transfer chain)\(^{75}\) neither possibility seems likely.

This distance-constraint has important implications for the operation of a radical pair compass. It appears that animal cryptochromes (including those of birds) generally contain a fourth tryptophan, Trp DH, which extends the tryptophan triad to a tetrad. With an edge-to-edge distance of $\sim1.7$ nm, no magnetic field effects would be expected for [FAD$^+$• Trp$_2$H$^+$•] in the current model (Fig. 2a) if the radical lifetime is of the order of several microseconds. Recent studies of DmtCry\(^{38,39}\) and Xenopus laevis (6–4) photolyase\(^{36}\) suggest that this is indeed the case. Homology modelling of robin (Erithacus rubecula) Cry1a predicts an edge-to-edge distance of 1.96 nm for [FAD$^+$• Trp$_2$H$^+$•]. The absence of spin-selective recombination of these distant radical pairs on a microsecond timescale calls into question the current model (Fig. 2a). Our new scheme (Fig. 2b) not only liberates the model from the spatial constraints imposed by the charge recombination step, but also strongly amplifies the magnetic field effect. The scheme is even applicable when the partner of the FAD$^+$ radical is free to diffuse rather than attached to the cryptochrome. If such a small, rapidly tumbling radical reacted spin-selectively with a freely diffusing paramagnetic scavenger, there would be the added benefit that both would undergo slower spin relaxation than the protein-bound radicals. Modulation of the exchange interactions of freely diffusing paramagnetic species may cause singlet-triplet dephasing in the radical pair which could either attenuate or further amplify the anisotropy\(^{75}\).

Finally, lifting the restriction on the rate of the spin-selective recombination reaction also means that the detrimental effects of inter-radical exchange and dipolar interactions\(^{38,39}\) can be minimised by placing the radicals much further apart than would be permissible in the current model.

**Properties of the radicals.** In all the cases analysed above, the enhanced yields of the signalling state are largely independent of the hyperfine interactions in both the paramagnetic scavenger and WH$^+$•. In particular, the reduction in the magnetic field effect caused by the hyperfine interactions in WH$^+$• for the current model is not found for the scavenging reaction scheme. The mechanism does not require particular properties of the scavenger except that its spin relaxation is slow on the timescale of the spin dynamics and the recombination reactions. In practice, this suggests that the mechanism is feasible if spin relaxation in the scavenger occurs at a rate comparable to that in the primary radical pair. This restriction probably excludes certain rapidly relaxing species such as superoxide\(^{32}\) and many transition metal complexes. Disregarding this aspect for the moment, iron-sulphur clusters could in principle act as scavengers. We mention this in the light of the recent report of a multimeric complex of cryptochrome and a protein, IscA, containing [2Fe-2S] clusters\(^{77}\). While several aspects of this work are controversial, not least the claim that the complex possesses a permanent magnetic moment\(^{78}\), the reported structure is potentially interesting in the context of the mechanism suggested here. However, the
tryptophan triad/tetrad in cryptochrome is probably too far away from any of the [2Fe-2S] clusters for a sufficiently rapid electron transfer reaction.

In principle, molecules with electron spin greater than ½ (e.g. O₂⁻; J = 1) could act as scavengers. Although this again raises the question of rapid spin relaxation⁷², it opens the interesting prospect of directly linking the spin dynamics in the cryptochrome radical pair to redox-active ion channels, which have been implicated in neuronal firing in fruit flies⁶⁸,⁷⁹.

Conclusions

We have demonstrated that the directional response, both relative and absolute, of a radical pair to the Earth’s magnetic field can be significantly enhanced when one of the radicals can react with an external paramagnetic molecule. This scavenging reaction acts as a spin-selective recombination channel resulting in a field-dependent product yield even when spin-selective charge recombination in the radical pair is very slow. As a consequence, the radical pair mechanism is freed from the constraint that the constituents of the radical pair must be less than about 1.7 nm apart (edge-to-edge) in order that direct charge recombination is fast enough to compete with spin relaxation. We believe that our suggestion may have far-reaching implications for the detailed operation of the proposed quantum compass in birds.

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Author Contributions
D.R.K. conceived the idea behind this work, performed the research, and discussed the results with P.J.H. Both authors wrote the paper.

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