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On magnetic monopoles, the anomalous $g$-factor of the electron and the spin-orbit coupling in the Dirac theory

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1 Introduction

When Uhlenbeck and Goudsmit presented the concept of spin, Lorentz pointed out that the idea could not account for the magnetic dipole moment of the electron. Even if one were to put all the charge of a spherical electron on its equator, the current produced by the spinning motion would not be enough to match the magnitude of the anomalous Zeeman splitting observed. The algebra of the Dirac theory accounts very well for the measured values of this anomalous Zeeman splitting, but in the present paper we point out that the traditional physical interpretation of the mathematical formalism in terms of a magnetic dipole moment associated with the spin is at variance with the meaning of the algebra itself as it violates the built-in Lorentz symmetry. It interprets a vector as a scalar, which is a transgression that is similar to interpreting a tensor as a vector. The importance of such distinctions based on symmetry is well known. In relativistic quantum mechanics one discusses e.g. that the only bilinear Lorentz covariants that exist are one-component scalars, four-component vectors, six-component tensors, four-component axial vectors and one-component pseudo-scalars. Confusing a covariant of one type with a covariant of an other type is violating its symmetry. Using group theory we will propose an approach that respects the symmetry.

Section 2 contains an introduction to some aspects of the representation SU(2) for the rotation group and of the Dirac representation for the homogeneous Lorentz group, which will be used in the paper. This is a subject
matter that is considered to be “well known”. But we cover it from a very different, geometrical perspective than in its traditional treatment given in textbooks, which is far more abstract and algebraic. The insight gained from this different perspective will permit us to discern the error in the interpretation of the anomalous g-factor we mentioned above, and which we discuss in Subsection 6.1. Apparently this problem has escaped attention for more than 80 years, suggesting that the group theory is not as well understood as routinely assumed.

2 Group Representation Theory

2.1 SU(2) and the rotations of \( \mathbb{R}^3 \)

2.1.1 Principal idea

The general form of a SU(2) rotation matrix is:

\[
R = \left( \begin{array}{cc} \alpha_0 & -\alpha_1^* \\ \alpha_1 & \alpha_0^* \end{array} \right), \quad \text{with } \det R = \alpha_0 \alpha_0^* + \alpha_1 \alpha_1^* = 1.
\]

These matrices work on spinors:

\[
\xi = \left( \begin{array}{c} \xi_0 \\ \xi_1 \end{array} \right), \quad \text{with } \xi_0 \xi_0^* + \xi_1 \xi_1^* = 1,
\]

according to:

\[
\xi' = R \xi.
\]

The natural question is of course what the meaning of such a spinor is. In SO(3), the 3 \times 3 rotation matrices of \( \mathbb{R}^3 \) are working on 3 \times 1 matrices that represent vectors of \( \mathbb{R}^3 \), but in SU(2) the situation is different. Here the 2 \times 1 matrices, i.e. the spinors, represent rotations. The idea can be illustrated by considering the group multiplication table for an arbitrary group \((G, \circ)\):

\[
\begin{array}{ccccccc}
\circ & g_1 & g_2 & g_3 & \cdots & g_j & \cdots \\
g_1 & g_1 \circ g_1 & g_1 \circ g_2 & g_1 \circ g_3 & \cdots & g_1 \circ g_j & \cdots \\
g_2 & g_2 \circ g_1 & g_2 \circ g_2 & g_2 \circ g_3 & \cdots & g_2 \circ g_j & \cdots \\
\vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \cdots \\
g_k & g_k \circ g_1 & g_k \circ g_2 & g_k \circ g_3 & \cdots & g_k \circ g_j & \cdots \\
\vdots & \vdots & \vdots & \vdots & \ddots & \vdots & \cdots
\end{array}
\]

which illustrates that the group element \( g_k \) defines a function \( g_k \circ : G \to G; g_j \to g_k \circ g_j \). The notation \( g_k \circ \) for this function is somewhat arcane, but it has the advantage that the way it acts on a group element is obtained by mere juxtaposition of the symbols. In the specific case of the rotation group, we define then a rotation \( g_k \) no longer by all its function values \( g_k(r), \forall r \in \mathbb{R}^3 \), but by all function values \( g_k \circ g_j, \forall g_j \in G \). More rigorously, an arbitrary group element \( g_k \in G \) is identified with the function \( T_{g_k} \in F(G,G) \) that maps \( G \) to \( G \) according to:

\[
T_{g_k}(g_j) = g_k \circ g_j,
\]

where \( T_{g_k} \) is just a more standard notation for the function \( g_k \circ \) (We use here \( F(S_1,S_2) \) as a general notation for the set of functions from \( S_1 \) to \( S_2 \)). This identification implies that \( T_{g_k} \in F(G,G) \) represents \( g_k \in G \). Let us call this representation \( T_{g_k} \) of \( g_k \) the automorphism representation. The non-standard notation \( g_k \circ \) permits writing \( g_k \circ : g_j \in G \to g_j' = g_k \circ g_j \) and grasping more easily the idea of interpreting a rotation as a function that works on other rotations rather than on vectors. If we represent \( g_j \) by the SU(2) matrix \( X \), \( g_k \) by the SU(2) matrix \( R \), and \( g_j' \) by the SU(2) matrix \( X' \) then we have:

\[
g_j' = g_k \circ g_j = T_{g_k}(g_j) \quad \text{or} \quad g_j' = R X g_j.
\]

We see thus that we can represent \( T_{g_k} \) also by \( R \): In other words, the SU(2) matrices represent from this viewpoint two types of mathematical objects, viz. group elements \( g_j \in G \) and group automorphisms \( T_{g_k} \in F(G,G) \). This translates the fact that the automorphism group \( F(G,G,\circ) \) (where \( \circ \) is the composition of functions) of a group \((G,\circ)\), is isomorphic to \((G,\circ)\) under the mapping \( T \in F(G,F(G,G)) : g_k \to T_{g_k} \). In SU(2) it is possible to remove this
ambiguity between the representations of group elements and automorphisms by rewriting the second diagram in Eq. 5 as:

$$g_k \circ g_j = T_{jk} (g_j)$$

$$\xi' = \mathbf{R} \xi$$

by using the substitution:

$$X' = \mathbf{R} \mathbf{X}$$

where we recover Eq. 3 by defining the spinor $\xi$ as just a short-hand for the SU(2) rotation matrix:

$$\mathbf{X} = \begin{pmatrix} \xi_0 & -\xi_1^* \\ \xi_1 & \xi_0^* \end{pmatrix}, \quad \text{with } \xi_0 \xi_0^* + \xi_1 \xi_1^* = 1,$$

by taking its first column. In fact, all the information of the matrix $\mathbf{X}$ is already given by its first column. When we know the first column we know everything we need to know to write the second column. Moreover the first column of $\mathbf{RX}$ will be $\mathbf{R}\xi$. If we note the second column of $\mathbf{X}$ as $\eta$, then the second column of $\mathbf{RX}$ will be $\mathbf{R}\eta$ and it will be possible to derive $\mathbf{R}\eta$ from $\mathbf{R}\xi$ in the same way as we could derive $\eta$ from $\xi$, viz. $(\mathbf{R}\eta)_0 = -(\mathbf{R}\xi)_1^*$ and $(\mathbf{R}\eta)_1 = (\mathbf{R}\xi)_0^*$, as the SU(2) matrices constitute a group. A spinor $\xi$ in SU(2) can thus be considered as a set of parameters that define a rotation, i.e. a set of coordinates for a rotation. This rises the question how the information about the rotations occurs inside the parameter set $\xi$.

### 2.1.2 Constructing the representation

The answer is that it is done by using the fact that any rotation can be obtained as a product of two spatial reflections in $\mathbb{R}^3$. The reflections with respect to the planes through the origin of $\mathbb{R}^3$ generate a group of rotations and reversals, of which the rotation group is a subgroup. It is easy to figure out how we write a $2 \times 2$ reflection matrix, and once we know the matrices for the reflections we can calculate the matrices for the rotations and reversals by making products. A reflection $A$ is defined by a unit vector $\mathbf{a}$ that is normal to its reflection plane. The coordinates of $\mathbf{a}$ can be expected to occur as parameters in the $2 \times 2$ matrices $A$ that defines the reflection $A$ but we do not know where. We therefore write the reflection matrix $A$ heuristically as $A = a_x \sigma_x + a_y \sigma_y + a_z \sigma_z$. The matrix $\sigma_x$ will tell us where and with which coefficients $a_x$ appears in $A$. The same is true, mutatis mutandis for $\sigma_y$ and $\sigma_z$. To find the matrices $\sigma_x, \sigma_y, \sigma_z$, express that $A^2 = \mathbf{I}$. We find that we can meet this requirement when the matrices $\sigma_j$ satisfy the conditions $\sigma_j \sigma_k + \sigma_k \sigma_j = 2 \delta_{jk} \mathbf{I}$. In other words, identifying them with the Pauli matrices will give us the representation searched for. By expressing a rotation as the product of two reflections, one can then derive the well-known Rodrigues formula:

$$\mathbf{R}(\mathbf{n}, \varphi) = \cos(\varphi/2) \mathbf{I} - i \sin(\varphi/2) [\mathbf{n}, \sigma],$$

for a rotation by an angle $\varphi$ around an axis defined by the unit vector $\mathbf{n}$. To derive this result it suffices to consider two reflections $A$ (with matrix $[a \cdot \sigma]$) and $B$ (with matrix $[b \cdot \sigma]$) whose planes contain $\mathbf{n}$, and which have an angle $\varphi/2$ between them, and to use the algebraic identity $[b \cdot \sigma] [a \cdot \sigma] = (b \cdot a) \mathbf{I} + i (b \wedge a) \cdot \sigma$. There is an infinite set of such pairs of planes, and which precise pair one chooses from this set does not matter.

### 2.1.3 A parallel formalism for vectors

By construction, this representation contains for the moment only group elements. Of course, it would be convenient if we were also able to calculate the action of the group elements on vectors. This can be done by developing a parallel formalism for the matrices $A$, wherein $A$ takes a different meaning and obeys a different kind of algebra. As the matrix $A$ contains the components of the vector $\mathbf{a}$ we can conceive the idea of taking the matrix $A$ also as the representation of the unit vector $\mathbf{a}$. This idea can be generalized to a vector $\mathbf{v}$ of arbitrary length, which is then represented by $\mathbf{V} = v_x \sigma_x + v_y \sigma_y + v_z \sigma_z$. We have then $\mathbf{V}^2 = -(\det \mathbf{V}) \mathbf{I} = v^2 \mathbf{I}$. This idea that within SU(2) a vector $\mathbf{v} \in \mathbb{R}^3$ is represented by a vector $\mathbf{v} \cdot \sigma$ according to the isomorphism:

$$\mathbf{v} = v_x \mathbf{e}_x + v_y \mathbf{e}_y + v_z \mathbf{e}_z \leftrightarrow v_x \sigma_x + v_y \sigma_y + v_z \sigma_z = \begin{pmatrix} v_x & v_y \\ v_y & -v_x \end{pmatrix} \mathbf{v} \cdot \sigma,$$

was introduced by Cartan [1]. From $\mathbf{V}^2 = v^2 \mathbf{I}$ it follows that: $\mathbf{V}_1 \mathbf{V}_2 + \mathbf{V}_2 \mathbf{V}_1 = 2 (v_1 \cdot v_2) \mathbf{I}$. To find out how the group acts on these representations of vectors, it suffices to observe that the reflection $A$, defined by the unit vector $\mathbf{a}$, transforms $\mathbf{v}$ into $A(\mathbf{v}) = \mathbf{v} - 2 (\mathbf{v} \cdot \mathbf{a}) \mathbf{a}$. Expressed in the matrices this yields: $\mathbf{V} \rightarrow -A \mathbf{V} A$. We see that this transformation law for vectors $\mathbf{v}$ is quadratic in $A$ in contrast with the transformation law for group elements $g$,
which is linear: $G \rightarrow AG$. Vectors transform thus quadratically as rank-2 tensor products of spinors, whereas spinors transform linearly.

Both in the representation matrices $A = a \cdot \sigma$ for reflections $A$ and $V = v \cdot \sigma$ for vectors $v$, $\sigma_x$, $\sigma_y$ and $\sigma_z$ are thus the Pauli matrices, and the symbol $\equiv$ serves to flag the introduction of a (rather confusing) stenographic notation $\sigma = (\sigma_x, \sigma_y, \sigma_z)$. The Pauli matrices are thus the images of the basis vectors $e_x, e_y, e_z$ in the isomorphism $(e_i \leftrightarrow \sigma_j)$ defined by Eq. 9. The drawback of the convenient convention to use the shorthand $\sigma$ for $(\sigma_x, \sigma_y, \sigma_z)$ is that it may create the misleading impression that $v \cdot \sigma$ represents a scalar, which it does not. It just represents the counterpart in the isomorphism of what would be a pedantic notation $(v_x e_x, v_y e_y, v_z e_z)$ for $v = v_x e_x + v_y e_y + v_z e_z = v$.

The reader will notice that the definition $V = v \cdot \sigma$ with $V^2 = v^2 I$ is analogous to Dirac’s way of introducing the gamma matrices to write the energy-momentum four-vector as $E\gamma_1 + cp\gamma_2$ and postulating $(E\gamma_1 + cp\gamma_2)^2 = (E^2 - c^2 p^2) I$. In other words, it is the metric that defines the whole formalism, because we are considering groups of metric-conserving transformations (as the definition of a geometry in the philosophy of Felix Klein’s Erlangen program). For more information about the calculus on the rotation and reversal matrices, we refer the reader to reference [2]. Let us just mention that as a reflection $A$ works on a vector $v$ according to $V \rightarrow -AVA = -AVA^{-1}$, a rotation $R = BA$ will work on it according to $V \rightarrow BAVAB = RVR^{-1} = RVR^\dagger$. The identity $R^{-1} = R^\dagger$ explains why we end up with SU(2).

In summary, there are two parallel formalisms in SU(2), one for the vectors and one for the group elements. In both formalisms a matrix $V = v \cdot \sigma$ can occur but with different meanings. In a formalism for group elements, $v$ fulfills the role of the unit vector $a$ that defines the reflection $A$, such that we must have $|v| = 1$, and then the reflection matrix $V = A$ transforms according to: $A \rightarrow GA$ under a group element $g$ with matrix representation $G$. The new group element represented by $GA$ will then no longer be a reflection that can be associated with a unit vector like it was the case for $A$. In a formalism of vectors, $|v|$ can be different from 1 and the matrix $V$ (that represents now a vector) transforms according to: $V \rightarrow GVGV^{-1} = GVG^\dagger$. Here $GVG^\dagger$ can be associated again with a vector.

### 2.1.4 Other approaches

The approach outlined above is non-standard. The standard treatment follows in general a linearization procedure. One starts the development by establishing the quadratic formalism $V \rightarrow -AVA$ and $V \rightarrow RVR^\dagger$ for vectors. The way back to a linear formalism $X \rightarrow RX$ (for group elements) or $\xi \rightarrow R\xi$ (for their spinors) is then tricky and shrouded in mystery. It amounts to so say to defining a spinor as a kind of a square root of an isotropic vector.

This runs for instance as follows. One first considers a triad of normalized, mutually orthogonal basis vectors $(e_x, e_y, e_z)$. One then observes that $(e'_x, e'_y, e'_z) = (R(e_x), R(e_y), R(e_z))$ defines $R$ unambiguously. There is a one-to-one correspondence between the rotated triads $(e'_x, e'_y, e'_z)$ and the rotations $R$ that produced them by acting on the chosen reference triad $(e_x, e_y, e_z)$. In a second stage, one considers that there is also a one-to-one correspondence between isotropic vectors $e'_x + ie'_y$ and triads $(e'_x, e'_y, e'_z)$. By separating the real and imaginary parts in $e'_x + ie'_y$, one can reconstruct $e'_x$ and $e'_y$, while $e'_z = e'_x \wedge e'_y$.

The isotropic vector is thus a parameter set that defines a rotation in a one-to-one fashion. It is thus a set of complex coordinates for a rotation. The coordinates of the isotropic vector $(x', y', z') = e'_x + ie'_y$ are thus not position coordinates but rotation coordinates. They do not define a position in $R^3$ because they were not introduced to do so. The only real point $(0, 0, 0) \in R^3$ that belongs to the isotropic cone and could define a position does not define a triad. The complex coordinates define nevertheless an object in real Euclidean space, viz. the triad. Therefore spinors, which (as we will see) represent the information about these isotropic vectors and the corresponding triads, do turn in Euclidean space, despite the widespread opinion that they should be considered as defined in some abstract internal space like in the example of isospin $[3]$.

---

1 This viewpoint was criticized by Cartan who stated [1]: “Certain physicists regard spinors as entities which are, in a sense, unaffected by the rotations which classical geometrical entities (vectors etc.) can undergo, and of which the components in a given reference frame are susceptible to undergo linear transformations which are in a sense autonomous. Cartan qualified this idea, which amounts to a parallel interpretation of the mathematics, as “startling”. In fact, the claim that spinors do not turn in physical space could be attributed to a lack of understanding of the geometrical meaning of spinors, such that the isospin-inspired viewpoint is then ad hoc and neither compelling nor unique. Based on the principle of Occam’s razor, we should not introduce the isospin-inspired assumption unless it is strictly necessary. The viewpoint based on the analogy with isospin contains even more exceptional assumptions than the isospin model itself as in the spin operator $S_z$, the index $z$ refers to physical space despite the denial that the spinor would not turn in physical space. In the isospin operator $I_z$, the index $z$ does not refer to physical space, such that the postulates of this formalism are less demanding. As will transpire from the main text, the approximation to describe the electron as a point charge leads to the introduction of many mathematical quantities that should permit one to keep treating the physics of a finite-size object that one has shrunk to a point in a mathematically self-consistent way. In this mathematical procedure, one “pinches” a part of the true physical reality out of existence. One
As $e'_x + ie'_y$ is a vector it transforms according to the rule $V \rightarrow RV^\dagger$. One can then discover spinors by noting that
\[ \det [(e'_x + ie'_y)\sigma] = 0 \] because $(e'_x + ie'_y)^2 = 0$. The rows and columns of $[(e'_x + ie'_y)\sigma] = 0$ must therefore be proportional. This permits us to write:
\[ [(e'_x + ie'_y)\sigma] = \sqrt{2} \left( \begin{array}{c} \xi_0 \\ \xi_1 \end{array} \right) \otimes (-\xi'_1 \xi'_0) \sqrt{2}, \tag{10} \]
where
\[ \begin{array}{c} \xi_0 \\ \xi_1 \end{array} = R \begin{array}{c} \xi_0 \\ \xi_1 \end{array} \quad \text{and} \quad (-\xi'_1 \xi'_0) = (-\xi_1 \xi_0) R^\dagger. \tag{11} \]
The numbers $\sqrt{2}$ in Eq. 10 are introduced to satisfy the normalization condition $\xi_0\xi'_0 + \xi_1\xi'_1 = 1$. This way one linearizes the quadratic formalism $V \rightarrow RV^\dagger$ for vectors in terms of a linear formalism $\xi \rightarrow R\xi$ for spinors. This is then analogous to the way Dirac linearized the Klein-Gordon equation. The approach enhances our understanding of the formalism, as it permits us to see how the information about the rotated basis that defines the rotation is hidden inside the spinor. But by using it as the starting point for deriving the formalism, a spinor in SU(2) remains a mysterious object, a kind of square root of an isotropic vector, while the essential point, that it is just a rotation, remains hidden. It is conceptually much easier to understand the idea that a vector is a tensor quantity of rank 2 in the viewpoint voiced by Biedenharn.

2.2 The homogeneous Lorentz group

Also here the basic idea is that a spinor should be a set of coordinates for a group element. The conditions the analogues of the Pauli matrices will have to satisfy are now $\gamma_{\alpha}\gamma_{\nu} + \gamma_{\nu}\gamma_{\alpha} = 2g_{\alpha\nu}I$. There is no fourth $2 \times 2$ matrix that would anti-commute with all the Pauli matrices and therefore could be used to represent all reflections in Minkowski space-time and to generate in a second stage all Lorentz transformations. This problem can be overcome in the $4 \times 4$ representation based on the Dirac matrices, where $a_{\mu}\gamma^\mu$ represents the four-vector $(a_{\mu}, a^\dagger)$ and $\gamma^\dagger \neq I$. We have then to postulate $(\sum a_{\mu}\gamma^\mu)^2 = a_{\mu}^2 - a \cdot a$. The simplest representation of the Dirac matrices is the Weyl presentation:
\[ (a_{\mu}, a) \leftrightarrow \left( \begin{array}{c} a_{\mu}I - a \sigma \\ a_{\mu}I + a \sigma \end{array} \right) \]
\[ = a_{\mu}\gamma^\mu + a_{\mu}\gamma^\nu + a_{\nu}\gamma^\mu - a_{\nu}\gamma^\mu = a_{\mu}\gamma^\mu \equiv (a_{\mu}, a)\cdot(\gamma^\dagger, \gamma). \tag{12} \]
This representation is much more easy to manipulate than the traditional text book representation, as due to the block structure of the Weyl representation the formalism reduces to two sets of calculations with $2 \times 2$ matrices. We can write them as $A = a_{\mu}I + a \sigma$ and $A^* = a_{\mu}I - a \sigma$. These matrices occur as blocks on the secondary diagonal. They are both matrices that represent four-vectors in a SL(2,C) representation, but in two different types of SL(2,C) representation. Each of the two vector matrices can be used as starting point to set up a representation SL(2,C) of the Lorentz group [4]. The matrix $A^*$ is obtained from $A$ by the parity transformation $a \rightarrow -a$. The SL(2,C) representations that are working on the vector matrices are tricky. The formalism does no longer permit using a unit four-vector $(a_{\mu}, a)$ to define a general reflection in SL(2,C) as there is no fourth Pauli matrix to represent reflections with respect to $e_1$. Instead of that $a_{\mu}$ is associated with $I$. The SL(2,C) representations do thus not permit a clear distinction between $e_1$ and the identity element $I$ of the Lorentz group, which are both represented by $I$. This difficulty is removed by the introduction of the gamma matrices where clearly $\gamma_1 \neq I$. Nevertheless, if one contents oneself with describing only true Lorentz transformations which are products of an even number of space-time reflections, we can see by following the faith of the matrices within the Weyl representation that the $2 \times 2$ formalism builds a representation, whereby the four-vector $A = a_{\mu}I - a \sigma$ transforms according to $A \rightarrow LA^\dagger$, where $L^\dagger \neq L^{-1}$. In the other SL(2,C) representation, $A^*$ transforms according should however not interpret the apparent consequences of these procedures too literally, which has apparently been the case in the viewpoint voiced by Biedenharn.
to \( A^* \rightarrow L^{-1}A^*L^{-1\dagger} \). We see that in the Weyl formalism the 2 \( \times \) 2 blocks are just sequences where the presence and absence of the symbol * alternates, e.g. \( V_{2n}^* V_{2n-1} \cdots V_1^* V_1 \). The algebra in the other block is just given by inverting the presences and absences of the * symbol. Everything that happens in one 2 \( \times \) 2 block is thus defined by what happens in the other 2 \( \times \) 2 block, such that we can use the 2 \( \times \) 2 blocks as a shorthand for what happens in the 4 \( \times \) 4 formalism. We may note that \( V^* \) has the meaning of \((\nu, -\nu)\) in the representation without stars. This justifies the use we will make of the 2 \( \times \) 2 matrices in the following sections.

It is no longer possible to cram all the information about a general Lorentz transformation that is coded in a one-to-one fashion within a SL(2, \( \mathbb{C} \)) matrix into a single 2 \( \times \) 1 spinor like it was the case in SU(2). Fortunately, will not have to bother about this technicality in this paper. Once again, we refer the reader to reference [2] for more details about the solution of this problem and the group calculus.

We must finally point out that a representation has always its own internal self-consistent logic, such that there can be no ground to question any result correctly derived within a given representation by drawing in considerations from outside the context of that representation.

### 3 Lorentz Symmetry of Electromagnetism

#### 3.1 Some simple algebra in SL(2, \( \mathbb{C} \))

3.1.1 The fields

In view of the facts outlined in Subsection 2.2, in SL(2, \( \mathbb{C} \)) the four-gradient \((\partial/t, \nabla)\) is represented by \( \frac{\partial}{\partial t} I - \nabla \sigma \). Analogously, the four-potential \((V, cA)\) is represented by \( V I - cA \sigma \). We can now check what will happen if we "multiply" these two matrices. Using the identity \([a \cdot \sigma] [b \cdot \sigma] = (a \cdot b) I + i [(a \wedge b) \cdot \sigma]\) we find:

\[
\left[ \frac{\partial}{\partial c} I - \nabla \sigma \right] \left[ \frac{V}{c} I - A \sigma \right] = \\
\left[ \frac{1}{c^2} \frac{\partial V}{\partial t} + \nabla \cdot A \right] I - \frac{1}{c} \left[ (\nabla V + \partial A/\partial t) \cdot \sigma \right] + \frac{i}{c} \left[ (\nabla \wedge A) \cdot \sigma \right]
\]

with the Lorentz gauge condition \( \frac{1}{c^2} \frac{\partial V}{\partial t} + \nabla \cdot A = 0 \), we obtain thus:

\[
\left[ \frac{\partial}{\partial c} I - \nabla \sigma \right] \left[ \frac{V}{c} I - A \sigma \right] = \frac{1}{c} [(E + icB) \cdot \sigma].
\]

We recover thus automatically the expressions for the Lorentz gauge condition, and for the electric and magnetic fields in terms of the potentials. The term \( E + icB \) is the electromagnetic field tensor. The presence of \( i \) in an expression can be seen to signal that it is a pseudo-vector or a pseudo-scalar\(^2\). The vector \( E \) and pseudo-vector \( B \) are the symmetric and anti-symmetric three-component parts of the six-component field tensor. We see thus that symmetry is enough to recover all the definitions. It summarizes in a sense the reason why we need the theory of relativity by showing that Lorentz symmetry is the symmetry that is compatible with the structure of the Maxwell equations. A whole text book development is here elegantly summarized in one line of calculation. With this formalism, one can also write the four Maxwell equations jointly in one, very simple, matrix equation. It seems that this approach was first discovered by Majorana, but most of the time the presentation is less concise than here.

---

\(^2\) This is general and follows from the identity \([a \cdot \sigma] [b \cdot \sigma] = (a \cdot b) I + i [(a \wedge b) \cdot \sigma]\). Let us construct a general expression \([a^{(1)} I + a^{(3)} \cdot \sigma] [a^{(2)} I + a^{(3)} \cdot \sigma] \cdots [a^{(n)} I + a^{(3)} \cdot \sigma]\). First consider the case that \( n \) is even. All quantities in the product are rank \( n \) by construction. Some of them are one-component quantities. One can recognize them by the fact that they are multiples of the unit matrix. When these quantities contain an even number of vector terms \( a^{(3)} \cdot \sigma \), they will be real like \((a \cdot b) I\) and they will not change sign under parity transformation. They are scalars. When they contain an odd number of vector terms, they will be imaginary like \( i (a \wedge b) \cdot \sigma\) and they will not change sign under parity transformation. They are pseudo-scalars. The other quantities are three-component quantities. One can recognize them by the fact that they contain \( \sigma \) in their expression. When they contain an even number of vector terms, they will be imaginary like \( i (a \wedge b) \cdot \sigma\) and they will not change sign under parity transformation. They are pseudo-vectors. When they contain an odd number of vector terms, they are real like \( \sigma \cdot \sigma \cdot \sigma \cdot \sigma \), and they will change sign under a parity transformation. They are vectors.
3.1.2 The interactions

The charge-current four-vector \((\rho, j/c)\) for a moving point charge \(q\) with velocity \(\mathbf{v}\) is (up to the Lorentz factor \(\gamma = (1 - v^2/c^2)^{-1/2}\)) given by \((q, -qv/c)\), which is represented by \(qI - \frac{q}{c} \mathbf{v} \cdot \sigma\). Let us now couple this quantity with the electromagnetic-field tensor and calculate\(^3\):

\[
[qI - \frac{q}{c} \mathbf{v} \cdot \sigma] [(E + \kappa \mathbf{B}) \cdot \sigma].
\]

We obtain then:

\[
- \left[ \frac{q}{c} \mathbf{v} \cdot \mathbf{E} \right] I + q(E + \mathbf{v} \cdot \mathbf{B}) \cdot \sigma - \mathbf{q} \mathbf{v} \cdot \mathbf{B} I + i q \left| \mathbf{B} \cdot \sigma \right| - \frac{q}{c} \left[ (\mathbf{v} \cdot \mathbf{E}) \cdot \sigma \right]
\]

\[\text{(16)}\]

The whole paper is devoted to the meaning and the consequences of this single equation. Again, the presence of \(i\) signals here pseudo-scalars and pseudo-vectors, while the real terms correspond to scalars and vectors, as can be checked from the behaviour of the various terms under a parity transformation. We recognize here the Lorentz force.

All terms in Eq. 16 are referring to phenomena that are occurring to a single particle, not to various different particles. It therefore looks as though one can find a frame wherein such a most general situation is realized. It is therefore as though we get magnetic monopoles out of Eq. 16, while at face value we have not introduced any magnetic monopoles in Eq. 15. All we have introduced is the charge-current four-vector of one single moving point charge within a formalism that automatically accounts for Lorentz symmetry. The terms in Eq. 16 are all forces as Eq. 15 actually generalizes a term \(q\mathbf{I} - \frac{q}{c} \mathbf{v} \cdot \mathbf{E}\) corresponding to the work \(\mathbf{F} \cdot \mathbf{d}\mathbf{r}\) done against the force \(\mathbf{F}\) during an infinitesimal displacement \(\mathbf{d}\mathbf{r}\) over a time interval \(dt\). It is well known that the four-vector generalization of the force three-vector \(\mathbf{F}\) is \((\mathbf{F} \cdot \mathbf{v}/c, \mathbf{F})\), which contains this additional power-related term (up to a constant \(c\)). As the term \(q\mathbf{E} \cdot \mathbf{v}\) is here divided by \(c\), the result has again the dimension of a force. We will call such terms therefore scalar force terms. The other terms in Eq. 16 are all imaginary and they may at first sight look less familiar.

4 Magnetic Monopoles

4.1 Symmetry issue

There is a surprise in Eq. 16 in that it is seen to exhibit a complete symmetry between the electric and magnetic force terms. Each of the imaginary terms in Eq. 16 corresponds to a term that is a relativistic counterpart of a term in \((\mathbf{F} \cdot \mathbf{v}/c, \mathbf{F})\) obtained by using the substitution \(\mathbf{E} \rightarrow c\mathbf{B}\), \(c\mathbf{B} \rightarrow -\mathbf{E}\). The addition of these terms is necessary to obtain full relativistic symmetry for the total result, just like adding \(\kappa\mathbf{E}\) to \(\mathbf{E}\) is necessary to obtain an expression with full relativistic symmetry. Such a perfect symmetry in the forces is something that is believed to occur only if magnetic monopoles were to exist. In such an overall symmetry, the magnetic monopole would be the symmetric counterpart of the electric monopole. It will also not have escaped the attention of the reader that the imaginary three-component quantities in the electromagnetic field on a magnetic monopole \(q_m\):

\[
\mathbf{F}_m = q_m \left[ B \cdot \sigma \right] - \frac{q_m}{c^2} \left( (\mathbf{v} \cdot \mathbf{E}) \cdot \sigma \right),
\]

\[\text{(17)}\]

provided we take \(q_m = eq\).\(^4\) The one-component quantity \(-i \left| q\mathbf{v} \cdot \mathbf{B} \right| I\) is the corresponding power-related term that completes the force four-vector. It looks therefore as though we get magnetic monopoles out of Eq. 16, while at face value we have not introduced any magnetic monopoles in Eq. 15. All we have introduced is the charge-current four-vector of the velocity that automatically accounts for Lorentz symmetry. The terms in Eq. 16 are all forces as Eq. 15 actually generalizes a term \(q \mathbf{I} [E \cdot \sigma]\) in a rest frame to a moving frame by Lorentz covariance. Eq. 16 gives us thus the most general possible expression for an electromagnetic force. It is obvious that one can find a frame wherein such a most general situation is realized. It therefore looks as though invoking magnetic monopoles as a mechanism to obtain the symmetry exhibited by Eq. 16 could be a bit far-fetched. All terms in Eq. 16 are referring to phenomena that are occurring to a single particle, not to various different particles.

\(^3\) Of course, in Eq. 15 we have dropped out \(\gamma\) and \(\left| q\mathbf{I} - \frac{q}{c} \mathbf{v} \cdot \sigma \right|\) is not a four-vector. The true four-vector is actually \(\gamma \left[ q\mathbf{I} - \frac{q}{c} \mathbf{v} \cdot \sigma \right]\) rather than \(\left| q\mathbf{I} - \frac{q}{c} \mathbf{v} \cdot \sigma \right|\). Nevertheless, what we have done is correct, because the terms we are identifying in the Lorentz force equation, like e.g. \(\frac{d}{dt} q\mathbf{E}\) and \(q(E + \mathbf{v} \times \mathbf{B})\), are both not covariant, while the equation \(\frac{d}{dt} q\mathbf{E} = q(E + \mathbf{v} \times \mathbf{B})\) is covariant. In fact, it corresponds to rearranging \(\frac{dE}{dt} = \gamma(E + \mathbf{v} \times \mathbf{B})\) by swapping \(\gamma\) from the right-hand side to the left-hand side, where we can use it to replace \(\gamma d\mathbf{r}\) by \(dt\). Here \(\tau\) is the proper time. The quantities \(d\mathbf{r}, (\gamma dE, dp)\) and \((\gamma v \cdot E/c, \gamma(E + \mathbf{v} \times \mathbf{B}))\) that occur in the equation before the swapping are Lorentz covariant (See e.g. J.D. Jackson, in Classical Electrodynamics, (Wiley, New York, 1963), page 405). Once we know the equation is covariant we can swap sides with any quantity in it. Now that we know that the equation is covariant, further covariant operations on them will yield also covariant equations.

\(^4\) With the identities \(q_m = eq, \varepsilon_{\alpha\beta\mu0} = \frac{1}{\sqrt{2}}\), and the substitution \(E \rightarrow cB\), the equation \(E = \frac{d\mathbf{S}}{dt} + \sigma_0\mathbf{e}_0\) can be transformed into \(B = \frac{d\mathbf{S}}{dt} + \mathbf{e}_0\). Applying the theorem of Gauss to the magnetic monopole yields \(\oint B \cdot d\mathbf{S} = q_m\mu_0\).
4.2 Point-like rotational motion

We will discuss in Subsubsection 6.1.3 how the whole theory of electromagnetism just resumes to the description of interactions of charges and currents with other charges and currents. A magnetic field is produced by moving charges. A constant magnetic field is thus a mathematical expedient to describe moving charges as a static non-moving phenomenon. It will be argued along similar lines that a magnetic monopole is just a mathematical construct to treat a moving electric charge as a static non-moving quantity.

This may look an absurd statement if we think about the charges as performing a uniform rectilinear motion, that can be described by an appropriate Lorentz transformation. However a charge that is in uniform circular motion in the laboratory frame, whereby the radius of the circle is very small such that it would look like a point to the naked eye, could indeed correspond intuitively to a phenomenon at rest with respect to the laboratory frame. As we will argue, this is the seed of the idea behind identifying the force terms that occur in Eq. 17 as forces acting on magnetic monopoles. Pushing the idea of a motion that cannot be detected to the extreme, a monopole can be imagined as the limit of a charge traveling on a circular orbit whose radius tends to zero such that the orbit shrinks to a point. It will then just become a charge that is rotating in a fixed position. In considering such a limit we are defining a new mathematical object, somewhat in the same way as Laurent Schwartz [5] defined the mathematical distribution that would correspond to a point-like dipole. This idea of shrinking the orbit to a point will be further elaborated below in Subsection 4.5 and in Subsection 7.2.

From this point of view, magnetic monopoles and magnetic fields are both just theoretical constructions to describe confined motion as a static phenomenon. Coining such mathematical quantities might look like a stroke of genius, but history shows that this was done unwittingly, as magnetic fields were just introduced as a phenomenological tool to describe the observations in terms of static quantities before their physics were truly understood in terms of motion. They were thus defined based on a visual illusion of rest. Of course, the idea to treat motion as a phenomenon at rest is a kind of tricky and beyond guessing. If not clearly spelled out, it may thus easily lead to confusion.

4.3 Quantization issue

We may note that Dirac’s argument that the existence of monopoles would lead to the quantization of charge amounts to postulating:

\[
\frac{q q_m}{2\pi \epsilon_0 \hbar c^2} \in \mathbb{Z}. \tag{18}
\]

For an electron with charge \( q \) and its associated magnetic monopole charge \( q_m = cq \), the quantity that is required to be an integer becomes then:

\[
\frac{q q_m}{2\pi \epsilon_0 \hbar c^2} = 2\alpha, \quad \text{where: } \alpha = \frac{q^2}{4\pi \epsilon_0 \hbar c} \text{ is the fine-structure constant.} \tag{19}
\]

As \( \alpha \approx 1/137 \), the prediction \( 2\alpha \in \mathbb{Z} \) is way off, but by rewriting Eq. 19 as:

\[
\frac{q q_m}{2\pi \alpha \epsilon_0 \hbar c^2} = \hbar, \tag{20}
\]

and considering \( \alpha \) as a constant of nature, Dirac’s argument could perhaps be saved. In fact, as orbital angular momentum occurs in multiples of \( \hbar \), charge would have to come in multiples of \( q \). However, in particle physics, \( \alpha \) is not considered to be a constant.

4.4 Refining the concepts

There are several attitudes one can adopt with respect to these considerations. The first one would be to reject them all together with contempt, on the grounds that they seem to question the official viewpoint of hard-won “established science”. However, an alternative attitude is possible. Before we can address it we must first further develop our Weltbild for the magnetic monopole.

In the Appendix we show that an electron traveling at a velocity \( v \) can be associated with a singular current (that has not to flow along a closed loop). It will become obvious from this approach that a single moving charge can indeed be considered as a “magnetic charge” with a magnetic moment. This magnetic moment could be called a monopole moment because there is only one charge in motion, in contrast with the macroscopic situation in e.g. a circular current loop where there are many such charges and we talk about a dipole moment. The main difference is that one cannot really claim that there exists a torque pulling on a single charge on a circular orbit described as a loop, while one
can when there is more than one charge traveling around the loop. In this respect, the magnetic moment of a single magnetic charge can be called a “magnetic monopole moment”, where it has to be emphasized that there is no hyphen between “magnetic” and “monopole” because “magnetic” refers to a magnetic moment, not to a magnetic monopole.

A magnetic monopole is an all-together different and unrelated concept. Not all single “magnetic charges” can be considered as magnetic monopoles. Only the magnetic charges that correspond to point-like rotational motion will be considered as true magnetic monopoles. Magnetic monopoles will thus not be new physical particles one would have to search for to confirm Dirac’s predictions. They will appear to be just a mathematical means to deal with moving charges whose motion remains hidden to the eye by remaining confined “inside a point”, with the effect that we think that we are dealing with a purely static situation. There is then a confusion that has to be avoided. The “magnetic monopole moment” of a magnetic monopole could be called a magnetic-monopole moment (with a hyphen), where the latter would be an ellipse for a magnetic-monopole’s magnetic monopole moment.

There is an important reason for making the distinction between “magnetic charges” corresponding to currents $qv$ and true point-like magnetic monopoles. If we associated magnetic monopoles with the visible, not point-like currents $qv$, then the classification of the interaction terms would be wrong. The criterion to classify the terms would then be if some term $j = qv$ occurs in them or otherwise. The part $q(v \land B)$ would then be due to the interaction of the magnetic monopole with the magnetic field, while it is conventionally attributed to the interaction of the moving electric charge with the magnetic field. On the other hand, the part $cqB$ would be due to the interaction of the electric charge with the magnetic field, while it is conventionally attributed to the interaction of the magnetic monopole with the magnetic field.

The classification must rather be based on a distinction between point-like hidden and not point-like visible currents. We will show in Subsection 7.2 that the terms in Eq. 17 are related to precession. Such a precession can be mentally visualized as a kind of point-like hidden motion and it has hidden energy we must account for if we want to get our calculations right. The terms in Eq. 17 correspond thus to hidden rotational motion rather than to translational motion (which is conceptually always visible). These rotational effects can be described also in terms of a vorticity (as will be discussed in Subsubsection 7.2.3). The duality between the force in Eq. 17 and the Lorentz force is thus rather based on a duality between non point-like visible (rotational and translational) and point-like invisible (rotational) motion than on a duality between electric charges and magnetic charges.

4.5 Disentangling the two unrelated issues of symmetry and quantization

On the basis of all this, we can and must now also make a further distinction between two concepts of magnetic monopoles. They should not be confused because they address two completely unrelated issues. The first issue is symmetry in the equations describing electromagnetism. As Eq. 17 shows, we get it for free. We do not need to postulate the existence of a true magnetic monopole to obtain the symmetrical counterpart of the Lorentz force in the form of Eq. 17. These terms are already there and we will show that they have already been experimentally observed in the form of the anomalous Zeeman effect and the spin-orbit coupling. Our first concept of magnetic monopole is thus only a mathematical hype. We rewrite $cq$ as $qm$, just to enhance the symmetry, rendering it more evident. We obtain then a shiny interpretation for the symmetry revealed by the existence of the terms in Eq. 17, but as the derivations show, it is also possible to describe everything in a less glittering way that only calls for electric monopoles. There is no new physical quantity, the only truly existing physical quantity is $q$.

The second issue is the quantization of charge, which we do not get for free at all, which is why Dirac introduced his magnetic-monopole concept. Dirac’s argument does not explain everything, as it hinges on the quantization of angular momentum which is also just an empirical fact. We know very well to describe the quantization of angular momentum by quantum mechanics but our intuition about it is not any better than our intuition about the quantization of charge. In fact, quantization of charge is conceptually a less difficult concept than quantization of angular momentum because charge is a fundamental quantity. Its definition does not rely on the definition of other quantities which are themselves not quantized like $r$ and $p$ in the definition $r \land p$ of angular momentum. Of course, it is eventually just experimental evidence that can tell if Dirac’s construction is justified. If Dirac’s monopole existed then it would lead to a second equation that is completely analogous to Eq. 16, with $q$ replaced by $Qm/c$ for some value of $Qm \neq qm$.

The two different issues lead to two distinct concepts of magnetic monopoles, as we do not need Dirac’s monopole to get the symmetry between the electric and the magnetic force terms, while the monopole $qm$, which accounts for that symmetry does not tally with the prediction based on Dirac’s construction needed to obtain quantization. Traditionally the two issues are mentioned in one breath. If in following this tradition however, we merge the two a priori completely unrelated issues into a single one, then it would appear as though Dirac’s construction misses the point underlying the introduction of $qm = cq$ in the first issue, which is that a magnetic monopole just serves to describe rotational motion confined to a point. In fact, Dirac’s monopole is a current (called a string) that stretches from some point to infinity, which is all but confined (Why this is wrong will be further discussed in Subsubsection 8.6.2). If we bleed the two issues, the existence of two equations, one with $qm$ and one with $Qm$ will also raise the
question why we use only $Q_m$ and not also $q_m$ in Dirac’s argument. If we do not confuse them, then one might perhaps think of an argument why we only use $Q_m$, based on the idea that $q_m$ is not a “true monopole” in Dirac’s sense.

5 Anomalous Zeeman Effect and Spin-Orbit Coupling as Purely Classical Phenomena

5.1 A striking similarity

There is an alluring way to interpret Eq. 16 completely differently as follows:

\[
\begin{align*}
-\frac{i}{c} \left[ \mathbf{v} \cdot \mathbf{E} \right] & \quad \text{power term} \\
-\frac{i}{c} \left[ q \mathbf{v} \cdot \mathbf{B} \right] & \quad \text{dual power term} \\
+q \left( E + \mathbf{v} \cdot \mathbf{B} \cdot \mathbf{\sigma} \right) & \quad \text{Lorentz force } \mathbf{F} \\
+\frac{ie}{c} \left[ \mathbf{B} \cdot \mathbf{\sigma} \right] & \quad \text{looks similar to} \\
-\frac{q}{c} \left[ (\mathbf{v} \times \mathbf{E}) \cdot \mathbf{\sigma} \right] & \quad \text{anomalous Zeeman effect} \\
\end{align*}
\]

\[\text{(21)}\]

We have labeled here the magnetic-monopole terms from Eq. 17 as “looking similar” to the anomalous Zeeman effect and to the spin-orbit coupling. What we want to refer to with the terminology “looks similar”, is that the imaginary terms on the second line of Eq. 21 correspond to force terms which are up to the proportionality factor $-\frac{\Delta}{2mc^2}$ equal to the energy terms derived from the Dirac theory for the anomalous Zeeman effect and for the spin-orbit coupling, but without the correction for the Thomas precession.\(^5\)

As will be explained in Subsection 8, Eq. 21 is obtained from Eq. 15 which just expresses the product of the three terms $\left[ q \mathbf{I} - \frac{2}{c} \mathbf{v} \cdot \mathbf{E} \right] \frac{\partial}{\partial t} \left[ 1 - \nabla \cdot \mathbf{\sigma} \right] \left[ \mathbf{I} - A \cdot \mathbf{\sigma} \right]$, while the correct physics for the anomalous Zeeman effect and the spin-orbit coupling are obtained by considering a variant $-\frac{\hbar}{2mc^2} \left[ \frac{\partial}{\partial t} \mathbf{I} - \nabla \cdot \mathbf{\sigma} \right] \left[ 1 + \frac{2}{c} \mathbf{v} \cdot \mathbf{E} \right] \left[ \mathbf{I} + A \cdot \mathbf{\sigma} \right]$, wherein the three types of terms are occurring in a different order.

For the derivation of the anomalous Zeeman term $-\frac{\hbar q}{2mc^2} \left[ \mathbf{B} \cdot \mathbf{\sigma} \right]$, the change of order has no incidence, such that the algebra that leads to the term $\frac{ie}{c} \left[ \mathbf{B} \cdot \mathbf{\sigma} \right]$ in Eq. 21 is exactly the same as the one that occurs in the derivation of the anomalous Zeeman effect from the Dirac equation. That we obtain the expression for the anomalous Zeeman effect up to the factor $-\frac{\hbar}{2mc^2}$ in Eq. 21 is thus not a coincidence. Also for the calculation of the spin-orbit term the order of the three terms in the product does not matter. For both orders of the three terms the calculation does not account for the correction due to the Thomas precession.

There is a classical rationale for the correct calculation of the spin-orbit coupling that runs as follows. As can be seen e.g. from Eq. 22, the total magnetic field experienced by the electron in its co-moving frame is (to first approximation, whereby one neglects the factors $\gamma$) given by $\mathbf{B} = \mathbf{B} - \frac{1}{c^2} \mathbf{v} \times \mathbf{E} = \mathbf{B} + \mathbf{B}_n$. The electric field of the nucleus gives rise to a magnetic field $\mathbf{B}_n = -\frac{1}{c} \left[ \mathbf{v} \times \mathbf{E} \right] \cdot \mathbf{\sigma}$ in a frame that is co-moving with the traveling electron. The interaction of the electron with the magnetic field $\mathbf{B}_n$ gives rise to an anomalous Zeeman term $-\frac{\hbar q}{2mc^2} \left[ \mathbf{B}_n \cdot \mathbf{\sigma} \right] = \frac{\hbar}{2mc^2} \times \frac{1}{2} \left[ (\mathbf{v} \cdot \mathbf{E}) \cdot \mathbf{\sigma} \right]$. This leads finally to an interaction energy $\frac{\hbar}{2mc^2} \frac{1}{2} \frac{\partial T}{\partial t}$. This has to be corrected for the Thomas precession $\omega_T = \frac{1}{2} \mathbf{v} \times \mathbf{a} = \frac{q}{2mc^2} \mathbf{v} \times \mathbf{E}$ of the electron in its orbit around the nucleus (see Eq. 30). According to Subsection 7.3, the corresponding energy is $\hbar \omega_T/2 = \frac{\hbar}{4mc^2} \frac{1}{2} \frac{\partial T}{\partial t}$ which has to be subtracted from the Zeeman term. The absolute value of the correction for the Thomas precession is half that of the Zeeman term, which is the reason why it is referred to as the Thomas half. Both terms on the second line of Eq. 21 are thus anomalous Zeeman terms due to magnetic fields in the rest frame of the electron, while the Thomas precession is a relativistic correction in the laboratory frame.

As mentioned above in Footnote 5, it is claimed in text books that the exact spin-orbit coupling term including the correction for Thomas precession can be derived from the Dirac theory (see e.g. [6]). We will show that this is a falsehood. The calculations claimed to derive the spin-orbit coupling with its correction for Thomas precession from the Dirac equation are wrong algebra leading to a correct physical result. The traditional Dirac theory is in reality unable to derive any of the two terms containing $\mathbf{v} \times \mathbf{E}$ described above, from the Dirac equation with minimal

\(^5\) Due to this absence of the term that corresponds to the Thomas precession, the reader could be inclined to conclude that the approach that leads to Eq. 21 is inferior to the traditional approach wherein it is possible to derive the correct expression for the spin-orbit coupling from the Dirac equation. However, as will be discussed later on in this subsection, this derivation is not correct and based on a fundamental error in the algebra.
coupling. It only succeeds in deriving the correct result by introducing logical errors. Our approach is thus superior to the traditional approach. By using the Dirac equation with the correct physical coupling, it is able to derive one of the two terms. The term it fails to derive is the correction for Thomas precession.

We may note that even an electron at rest within a magnetic field is subjected to the anomalous Zeeman effect. But in the absence of motion there is no Thomas precession, such that the anomalous Zeeman effect does then not require a correction for Thomas precession. Of course, Thomas precession occurs in any type of motion. There exists thus also a correction for Thomas precession in the case of a particle that is moving in a magnetic rather than in an electric field. However, the non-relativistic correction term for Thomas precession during the motion of an electron in a magnetic field \( \mathbf{B} \) is given by \( \omega = \frac{1}{2m_0} \mathbf{v} \cdot \mathbf{a} = \frac{q}{2m_0 c^2} \mathbf{v} \cdot (\mathbf{v} \wedge \mathbf{B}) \), which remains very small in the non-relativistic limit due to the presence of the factor \( v^2/c^2 \).

As we want to discuss the anomalous Zeeman effect and the spin-orbit coupling in the rest of the paper, the similarities exhibited in Eq. 21 seem to be a good way to make the transition between the two parts of the paper. That we can derive the anomalous Zeeman effect and the spin-orbit effect this way from a variant of the calculation leading to Eq. 16 and Eq. 21 is a major upheaval, because these two phenomena are traditionally considered as purely due to the electron spin. Just as it looked as though we obtained magnetic monopoles from Eq. 16 without having introduced them in Eq. 15, here it looks as though we now obtain spin-related effects without having introduced spin in the variant of Eq. 15.

It is extremely important to note that the matrix calculations one performs in going from Eq. 13 to Eq. 21 or from the variant to the more correct equation are not quantum mechanical. They are independent of any context of wave equations and entirely classical as all we have used is a group-theoretical formalism that automatically accounts for Lorentz symmetry. Very obviously, these derivations also do not contain spin. We have only introduced the charge-current four-vector of a single moving point charge. The algebraic expressions for the anomalous Zeeman effect and the spin-orbit coupling must thus be considered as purely classical and not spin-related.

We might indeed have been convinced that these terms are quantum mechanical rather than classical because they fitted nicely into quantum mechanics after their experimental discovery, while they were not covered by classical mechanics. But they become quantum mechanical only by the way we use them in quantum mechanics, where they fitted nicely into quantum mechanics after their experimental discovery, while they were not covered by classical mechanics. Squaring the Dirac equation will lead to a term \( \mathbf{p}^2 / 2m_0 \), which is used in the Schrödinger equation in the non-relativistic limit. In fact, the transition from relativistic to classical mechanics is obtained by putting \( \mathbf{p} = m_0 c \mathbf{v} \). And from this product we will obtain a term \( -e \mathbf{q} \mathbf{B} \cdot \mathbf{\sigma} \). Squaring the Dirac equation also leads to a term \( \mathbf{\sigma} \cdot \mathbf{E} \), which has to be divided by \( 2m_0 c^2 \) to reduce it to the operator \( \mathbf{p}^2 / 2m_0 \) which is used in the Schrödinger equation in the non-relativistic limit. In fact, the transition from relativistic to classical mechanics is obtained by putting \( E = m_0 c^2 + E_{cl} \). Then \( E^2 - c^2 p^2 = m_0^2 c^4 \) leads to \( E_{cl}^2 + 2m_0 c^2 E_{cl} + m_0^2 c^4 - c^2 p^2 = m_0^2 c^4 \). The terms \( m_0^2 c^4 \) can be dropped on both sides. After dividing both sides by \( 2m_0 c^2 \) and neglecting the term \( E_{cl}^2 / 2m_0 c^2 \) based on the observation that \( E_{cl} \ll m_0 c^2 \), one obtains then \( E_{cl} = 2m_0 c^2 \). The quantum mechanical version of this argument is arguably obtained by introducing \( E = m_0 c^2 + E_{cl} \) in the wave function through \( \psi = e^{-i m_0 c^2 t / \hbar} \psi_{cl} \) in the complete Dirac equation including the minimal substitution (but we will discover later on that this procedure can contain a major pitfall).

This way one can derive the Pauli equation from the Dirac equation\(^6\) and due to the division by \( 2m_0 c^2 \) this equation contains now the anomalous Zeeman term \( -\mathbf{\mu} \cdot \mathbf{B} \cdot \mathbf{\sigma} \). As Eq. 21 contains \( e \mathbf{B} \) this explains the conversion factor \( \frac{e}{2m_0} \). The Pauli equation does not contain a term that looks like the spin-orbit interaction, because it does not use the correct “minimal” substitution. We postpone the discussion of this point to Section 8, because it is rather intricate and raises several subsidiary issues.

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\(^6\) It must be noted that after squaring the Dirac equation, and dividing the result by \( 2m_0 c^2 \), we also recover a term \( -\mathbf{\mu} \cdot \mathbf{B} \cdot \mathbf{\sigma} \), which corresponds to the famous classical term \( -\mathbf{\mu} \cdot \mathbf{B} \cdot \mathbf{I} \) and describes the orbital Zeeman effect.
6 Problems with the Traditional Interpretation of the Anomalous Zeeman Effect

6.1 The anomalous Zeeman effect

6.1.1 An inconvenient truth: The physical imagery violates the symmetry

The anomalous Zeeman effect \( -\frac{he}{2ma} [B \cdot \sigma] \) is traditionally attributed to a coupling between the magnetic field \( B \) and the spin \( \frac{h}{2} \sigma \). In reality \( B \cdot \sigma \) is not a scalar product but just the way the vector \( B \) is written in the group theory. Traditionally one interprets the term \( -\frac{he}{2ma} [B \cdot \sigma] \) indeed as analogous to the orbital Zeeman term \( -\frac{he}{2ma} \hat{L} \cdot \hat{L} \) and \( -\frac{he}{2ma} [B \cdot \sigma] \). This is done by rewriting \(-\frac{he}{2ma} [B \cdot \sigma]\) as \(-\frac{2a}{2ma} [B \cdot \sigma] \), where \( \frac{h}{2} \sigma \) is postulated to be the operator \( \hat{S} \) corresponding to the “spin vector” \( S \), such that the term becomes then \(-B \cdot \frac{2a}{2ma} \hat{S} \) yielding an energy eigenvalue \(-B \cdot \frac{2a}{2ma} \hat{S} = -\mu_e \cdot B \), where \( \mu_e = 2\mu_B \). It is because the eigenvalues of \( S_z \) of \( \hat{S} \) are \( \pm \frac{\hbar}{2} \) that one is then obliged to introduce a factor \( g = 2 \) into the algebra to recover the correct value \(-\frac{2a}{2ma} [B \cdot \sigma] \). This interpretation tries thus to represent the term \(-\frac{he}{2ma} [B \cdot \sigma] \) as corresponding to the potential energy \( \mu_e \cdot B \) of a (spin-induced) magnetic dipole \( \mu_e \) within a magnetic field \( B \). But as we have explained in connection with Eq. 9 in Section 2, the quantity \(-\frac{he}{2ma} [B \cdot \sigma] \) represents a pseudo-vector, not a scalar like \(-\frac{he}{2ma} (B \cdot \hat{L}) \), whose scalar character transpires very clearly from the presence of the unit matrix \( 1 \) in it. The picture of the magnetic dipole \( \mu_e \) that would be proportional to a “spin vector” \( S \) is based on the misleading notation \( a \cdot \sigma \) we warned against above. The quantity \( \sigma \) is in reality the set of the three vectors \( e_x, e_y, e_z \). In the analogy one replaces thus wrongly the scalar quantity \( B \hat{L} \cdot \hat{L} \) \( = -\hbar B_z (x \frac{\partial}{\partial y} - y \frac{\partial}{\partial x}) \) by the vector quantity \( B_z \hat{S}_z = B_z \frac{\hbar}{2} \sigma_z \) (with analogous substitutions for \( B_x \hat{S}_x \) and \( B_y \hat{S}_y \)). The notation \( \frac{\hbar}{2} \sigma_z \) stands for a set of three vectors while the quantity \( \hat{L} \cdot \hat{L} \) stands for a set of three scalars. In fact, the three scalars \( \hat{L} \cdot \hat{L} \) \( = -i\hbar (y \frac{\partial}{\partial z} - z \frac{\partial}{\partial y}) \) stand for a pseudo-vector. Whereas the algebra used to calculate the anomalous \( g \)-factor is exact, such that it correctly reproduces the experimental results, the physical interpretation proposed is thus mathematically unsustainable, even if it might be intuitively appealing.

Of course these considerations clash ignominiously with accepted notions. We must therefore insist that the SL(2,\( \mathbb{C} \)) and Dirac representations are completely self-formalisms and that their algebra is a closed system that contains all it needs to contain, such that it is pointless to attack the conclusion by drawing in considerations that are external to SL(2,\( \mathbb{C} \)). In view of the strong resistance this conclusion might provoke, we give further arguments to back it and make a strong case for it (while we will give the correct interpretation of the term \(-\frac{he}{2ma} [B \cdot \sigma] \) in Section 7).

(1) As we explained above, the quantity \( \frac{he}{2ma} \) in the term \(-\frac{he}{2ma} [B \cdot \sigma] \) does not represent the spin in SU(2). It is thus not the spin operator \( \hat{S} \), and it is not correct to transform the term \(-\frac{he}{2ma} [B \cdot \sigma] \) into \(-\frac{he}{2ma} B \cdot \hat{S} = -\frac{he}{2ma} B \cdot \hat{S} \), with \( g = 2 \). We may note that in the context of the Dirac equation the presence of the term \(-\frac{he}{2ma} [B \cdot \sigma] \) is due to the minimal substitution used to derive the Dirac equation in an electromagnetic field from the free-space Dirac equation. But this is a substitution for a point charge, which is why it is called minimal in the first place (As pointed out in Subsection 8.2, it does not even account for the motion of the electron in the laboratory frame). If we had wanted to account for a potential energy within the magnetic field, of a magnetic dipole \( \mu_e \) associated with the spin, we should have introduced a more complicated substitution, with a term expressing how \( \mu_e \) couples to the electromagnetic field, or how it couples to the charge or magnetic dipole moment of another particle that is also present in its neighbourhood within the magnetic field. As we have not put such spin-related dipole effects into the formalism, they cannot come about by magic.

(2) That there is no spin-related dipole effect in the term \(-\frac{he}{2ma} [B \cdot \sigma] \) can also be appreciated from the fact that Eqs. 14-21 and their variant have been derived without introducing any considerations about spin. As the terms that occur in these calculations are the same ones as those that occur in the Dirac theory, none of the operators in the Dirac theory contains the spin. The physical origin of the terms we recollect from the squared Dirac equation is not the presence of spin. The terms only come about because we treat the problem in full rigor by using relativistic group theory. Within the Dirac equation, the spin occurs only implicitly, viz. inside the spinor wave function (as clearly

\[ \hat{L} = ( -i\hbar (y \frac{\partial}{\partial z} - z \frac{\partial}{\partial y}), -i\hbar (z \frac{\partial}{\partial x} - x \frac{\partial}{\partial z}), -i\hbar (x \frac{\partial}{\partial y} - y \frac{\partial}{\partial x})) \] Hence \( B \cdot \hat{L} \) stands here for the scalar operator \(-i\hbar (B_y \frac{\partial}{\partial z} - z \frac{\partial}{\partial y}) + B_z (x \frac{\partial}{\partial y} - y \frac{\partial}{\partial x}) + B_x (y \frac{\partial}{\partial x} - x \frac{\partial}{\partial y}) \), such that \( -\frac{he}{2ma} (B \cdot \hat{L}) \) is a scalar operator, which for the case \( B \parallel e_z \) reduces to \(-i\hbar B_z (x \frac{\partial}{\partial y} - y \frac{\partial}{\partial x}) \).
explained in [2]). It is the requirement that the spinor wave function must be an eigenstate of a vector operator \( K \cdot \sigma \) or a pseudo-vector operator like \( \frac{-ie}{2m_0} [B \cdot \sigma] \) (or their four-dimensional analogous expressions in terms of gamma matrices), that forces the spin to align itself with the vector \( K \) or pseudo-vector \( B \) in the calculations and is this way responsible for the occurrence of the up and down eigenstates.

(3) We may also appreciate that it would be extremely puzzling if it were true that we can calculate a magnetic dipole moment \( \mu \) produced by the electron spin with fantastic precision in quantum electrodynamics, without having to specify anything in the calculations about the internal charge-current and mass distributions inside the electron. In fact, the presence of such current distributions is intuitively the only mechanism we know to account for the existence of a magnetic dipole \( \mu \). The remark of Lorentz we quoted above shows that using this mechanism to explain the anomalous Zeeman effect could be wrong despite the fact that it is intuitively appealing.

(4) As will be pointed out in the Appendix, the interpretation of the orbital Zeeman term \( -\mu \cdot B \), on which one tries to build here intuition about the anomalous Zeeman term by analogy, is itself flawed, because there does not exist such a thing as a potential energy with respect to a magnetic field. The correct interpretation of \( -\mu \cdot B \) will be given in Subsubsection 7.2.1.

(5) We may further note that a three-component expression for the spin can never be a complete description within a fully relativistic context, due to the theorem about the bilinear covariants mentioned in the Introduction. Only covariants with 1, 4 or 6 components exist. Therefore, either one or three components must be missing in the three-component description. In our work [2], the expression for the relativistic generalization of the spin operator \( \frac{1}{2} [s \cdot \sigma] \) in SU(2), becomes a four-component quantity \( \frac{1}{2} [s \cdot \sigma + s \cdot \sigma] \) within SL(2,C).

(6) As we have not introduced spin in Eq. 15 or its variant, all “dipole” effects we can expect to obtain from Eq. 15 or its variant are orbital effects due to moving point charges. In fact, we introduce magnetic moments into formalism through the term \( -\frac{e}{2m_0} [v \cdot \sigma] \) in Eq. 15 or its variant. This can be understood from our calculation of a magnetic moment produced by a current presented in the Appendix. The anomalous Zeeman term \( \frac{-ie}{2m_0} [B \cdot \sigma] \) can thus not involve any magnetic dipole moment for the electron that we are studying: It cannot contain a spin-induced “intrinsic” magnetic dipole moment as it can be derived in a context without spin. It also cannot contain any orbital magnetic “dipole” moment as it does not contain \( v \). The anomalous Zeeman effect does indeed not depend on the velocity of the electron. In the measurements of the anomalous g-factor within a Penning trap [7], we can reduce the velocity of the electron such that one practically reaches the limit \( v \to 0 \). The anomalous Zeeman effect would still exist.

(7) We may note that the exercise of rewriting \( \frac{-ie}{2m_0} [B \cdot \sigma] \) as \( -\langle \hat{\mu}_z B \rangle \) is not only mathematically flawed but also physically futile. The wrong interpretation of the algebra is used to introduce by brute force an intuitive classical image, while the facts of the non-interpreted algebra will eventually force us to give up on that classical image anyway. The presence of a scalar product \( \mu \cdot B \) conjures up the image of an operator with a continuous spectrum, due to the continuum of possible angles between \( \mu \) and \( B \), while eventually this classical picture of a magnetic dipole in a magnetic field has to be abandoned as only the up and down states are observed experimentally.

(8) To readers who may still feel reluctant to accept the arguments formulated under points (1)-(7) because they fly in the face of the accepted notions, we may perhaps ask to consider how the inherent profound difficulty of quantum mechanics forces us to teach it a little bit like a religion. A nice example of a quasi-religious mystery is the concept of particle-wave duality. With some misplaced irony one could compare it with the Christian dogma of the mystery of the Holy Trinity. Three different persons (the Father, the Son and the Holy Ghost), are claimed to be only one God, and one is invited to accept this puzzling postulate as a factual truth, for which will not be given any further explanation because it is a mystery. In complete analogy, an electron is postulated to be both a particle and a wave. This is also quite puzzling a notion, for which we are told that we should accept it as a quantum mystery [8]. One could claim sarcastically that the two mysteries resemble one another as two peas in a pod. Of course, there is a very essential point that makes all the difference between the quantum mystery and the religious mystery, and justifies why we can postulate that one has to accept the quantum mystery without further asking and just “shut up and calculate”. That point is the agreement of the theory with experimental evidence. Quantum mechanics passes the test of the comparison with experimental data with flying colors by grinding out all the correct answers with impressive precision.

Asking if a physical theory provides the correct answers is indeed a crucial criterion to assess its value. But a formalism that turns out the correct answers is a far cry from a theory if it is mathematically flawed. The points (1)-(7) show that with respect to the criterion of mathematical self-consistency, the traditional interpretation of the anomalous g-factor of the electron in quantum mechanics is not even an option. What is wrong in this respect is not the algebra itself, which is correct, such that it indeed turns out the right answers. It is the interpretation routinely given to that algebra that is absurd as it violates the mathematics, even if the algebraic expression \( -\frac{ie}{2m_0} [B \cdot \sigma] \) correctly accounts for the anomalous Zeeman effect. On this and countless other occasions quantum mechanics has time and again overruled the logically peremptory, true geometrical meaning of the group theory explained in Section 2 in favour of a self-cooked parallel interpretation [2]. The argument that we should accept this because the theory turns out the right answers does not hold sway. What is confirmed by the experiments is just the algebra, not the
interpretation of that algebra. The real anathema resides in interpreting $-\frac{\hbar}{2m_0} |\mathbf{B} \cdot \mathbf{\sigma}|$ as a scalar, not in questioning the traditional orthodoxy.

Of course these observations also raise the question how we must define the spin operator if it is not given by $\hat{S} = \frac{\hbar}{2} \mathbf{s}$. We may mention that it is possible to preserve the physical image of the spin as a vector by using a different definition $\frac{\hbar}{2} [s \cdot \mathbf{\sigma}]$ for the spin operator than $\hat{S} = \frac{\hbar}{2} \mathbf{s}$. This approach can be developed without changing a iota to the results of Pauli’s spin calculus, such that it leads to an identical agreement with experimental data. It even presents less problems of interpretation, but the issue entails a whole domino chain of related questions and answers, whose development and discussion are beyond the scope of any paper of reasonable length [2].

6.1.2 Difficult questions

The problem with this discussion of the anomalous Zeeman effect is that without the traditional interpretation, the physics related to the non-relativistic Zeeman term $-\frac{\hbar}{2m_0} [\mathbf{B} \cdot \mathbf{\sigma}]$ become mysterious. The problem is two-fold. There is a classical enigma, as the interaction exists in principle already within classical electromagnetism, as the calculations given in Eqs. 14-21 or their variant clearly illustrate. The question is then of course what the classical meaning of this classical effect is supposed to be. There is also a quantum mechanical enigma. In fact, the two energy levels observed in the anomalous Zeeman splitting are traditionally interpreted as corresponding to the spin-up and spin-down states. But how can we explain the anomalous Zeeman splitting if the derivation of the term $-\frac{\hbar}{2m_0} [\mathbf{B} \cdot \mathbf{\sigma}]$ does not rely on a notion of spin?

6.1.3 Not a dipole-dipole but a charge-dipole interaction symmetry

The first step in our attempts to make sense of this puzzling situation is as follows. The Lorentz transformation for the electromagnetic field under a boost with velocity $\mathbf{v}$ can be written as:

$$\mathbf{E}'_\parallel = \mathbf{E}_\parallel, \quad \mathbf{B}'_\parallel = \mathbf{B}_\parallel, \quad \mathbf{E}'_\perp = \gamma (\mathbf{E}_\perp + \mathbf{v} \wedge \mathbf{B}_\perp), \quad \mathbf{B}'_\perp = \gamma (\mathbf{B}_\perp - \frac{1}{c} \mathbf{v} \wedge \mathbf{E}_\perp).$$ (22)

Here the indices $\perp$ and $\parallel$ are with respect to $\mathbf{v}$. Starting from a rest frame without magnetic field ($\mathbf{B} = 0$), we obtain a magnetic field ($\mathbf{B}' \neq 0$) in a moving frame, such that a magnetic field can be viewed (schematically) as a relativistic byproduct of the electric field. One could introduce the philosophy that only the Coulomb field really exists and that the magnetic field is just an optical illusion. By adopting this viewpoint, we can see that it is the macroscopic quantity $\mathbf{B}$ that contains a set of terms $q\mathbf{v}_j$, whose sum can be associated with a macroscopic magnetic dipole moment $\mathbf{\mu}$. The whole of Eq. 21 would in essence just be due to a calculation:

$$(q_1 \mathbf{l} + \frac{q_1}{c} [\mathbf{v}_1 \cdot \mathbf{\sigma}]) \sum_j (q_2_j \mathbf{l} - \frac{q_2_j}{c} [\mathbf{v}_2 \cdot \mathbf{\sigma}]) =$$

$$\sum_j (q_1 q_2_j \mathbf{l} + \frac{q_1 q_2_j}{c} [\mathbf{v}_1 \cdot \mathbf{\sigma}] - \frac{q_1 q_2_j}{c} [\mathbf{v}_2 \cdot \mathbf{\sigma}] - \frac{q_1 q_2_j}{c^2} (\mathbf{v}_1 \cdot \mathbf{v}_2_\mathbf{l}) - \frac{q_1 q_2_j}{c^2} [(\mathbf{v}_1 \wedge \mathbf{v}_2) \cdot \mathbf{\sigma}]).$$ (23)

Here $q_1$ and $\mathbf{v}_1$ are the parameters of the electron we are studying, after putting it in the electromagnetic field ($\mathbf{E}, \mathbf{B}$), which is generated by a distribution of charged particles described by the parameters $q_2$ and $\mathbf{v}_2$. The many terms $\mathbf{v}_2_j$ are hidden (with their coupling terms) in the macroscopic quantity $\mathbf{B}$, and the terms $q_2_j$ in $\mathbf{E}$ or $\mathbf{B}$. The macroscopic quantities $\mathbf{E}$ and $\mathbf{B}$ are only tools to express interactions of charges and currents with other charges and currents. We can then “derive” the structure of the backbone of Eq. 23 from the backbone of Eq. 23, by replacing $\sum_j q_2_j \mathbf{v}_2_j \rightarrow c \mathbf{B}$ and for the remaining terms $q_2 \rightarrow \mathbf{E}$. The term $\frac{2q_2_j}{c} [\mathbf{v}_1 \cdot \mathbf{\sigma}]$ will lead to two contributions $\frac{2}{c} (\mathbf{v} \cdot \mathbf{E}) \mathbf{l}$ and $\frac{2}{c} ([\mathbf{v} \wedge \mathbf{E}] \cdot \mathbf{\sigma}$, because $\mathbf{E}$ will be defined by $\mathbf{r}_2 - \mathbf{r}_1$. To make a rigorous derivation one would of course have to weight the various contributions with their coupling terms.

It can be seen this way that it is much more logical to consider the anomalous Zeeman effect as a term that accounts for the interaction of the charge of the electron with the magnetic dipoles produced by the current loop of the moving electron, which is generated by the classical derivation of Eq. 21 wherein the term containing $\mathbf{B} \cdot \mathbf{\sigma}$ does not allow for any “intrinsic” magnetic dipole moment on behalf of an electron of charge $q$ to interact with the magnetic field. That does not make sense to invoke another physical mechanism for the same term in the Dirac equation. The origin of the term containing $\mathbf{q} \mathbf{B}$ is obvious. We might initially just consider the force $q \mathbf{l} \cdot [\mathbf{E} \cdot \mathbf{\sigma}]$ instead of Eq. 15 and then generalize both terms independently by Lorentz symmetry. We recover then the general expression of Eq. 15, wherein $\mathbf{B}$ is no longer zero. We must thus conclude that there is in general an interaction between an electric point charge and a magnetic field. The pre-factor $-\frac{\hbar}{2m_0}$ in the anomalous
Zeeman term $-\frac{\hbar}{2m_0}[B \cdot \sigma]$ has been used to attribute a non-classical magnetic dipole moment $\mu_\sigma$ to the charge $q$, while in reality the term $-\frac{\hbar}{2m_0}[B \cdot \sigma]$ corresponds to a charge-dipole interaction of a point charge $q$ with the macroscopic dipole $\mu$ corresponding to the current loops that produce $B$. We can see that the anomalous value $g = 2$ is just due to the introduction of $\frac{\hbar}{2}$ with the aim to recover $\frac{\hbar}{2} \sigma$ as described above. In reality, if we take the liberty to continue to use the textbook misnomer “magnetic dipoles” for the magnetic moments of single moving electrons, then all “magnetic-dipole” effects in the formalism are orbital, and there is no spin-induced magnetic dipole moment in the algebra. The term in the Dirac equation that gives rise to the anomalous Zeeman term belongs to the symmetric counterpart $cqB \cdot \sigma + \frac{1}{2}(v \wedge E) \cdot \sigma$, of the Lorentz force $q(E + v \times B) \cdot \sigma$ where the roles of the electric and magnetic fields have been exchanged. Within the expression $cqB \cdot \sigma + \frac{1}{2}(v \wedge E) \cdot \sigma$ itself, the term $cqB \cdot \sigma$ is the symmetric counterpart of $\frac{1}{2}(v \wedge E) \cdot \sigma$ wherein the roles of charge and “magnetic dipoles” have been exchanged. Within $cqB \cdot \sigma$ the charge of the electron interacts with the “magnetic dipoles” of the moving electrons that generate the magnetic field. In the term $\frac{1}{2}(v \wedge E) \cdot \sigma$ the “magnetic dipole” generated by the moving electron interacts with the charges of the electric field. The symmetry of the anomalous Zeeman effect is intrinsically different from that of the orbital Zeeman effect, as the latter depends on two velocities, rather than one velocity. In other words, the orbital Zeeman effect corresponds to a “dipole”-dipole interaction, while the anomalous Zeeman effect stems from a charge-dipole interaction and the spin-orbit coupling to a “dipole”-charge interaction. This shows clearly that the anomalous $g$-factor should not be visualized in terms of a “dipole”-dipole interaction, as Dirac has done.

7 The Physical Meaning of the Anomalous Zeeman Effect

7.1 Introduction

Of course, getting a feeling for the anomalous Zeeman effect and explaining why it leads to two Zeeman levels is more complicated, even if it follows very clearly from the algebra. All terms in Eq. 21 we “understood” classically are in reality facts of life that we had to accept after discovering them experimentally and to which we got used. The anomalous Zeeman effect and the spin-orbit coupling should thus be considered on the same footing. However, the spin-orbit term can be understood in terms of precession, as explained in Subsection 8.1. It is a current-charge interaction, and therefore the symmetrical counterpart of the anomalous Zeeman effect, which is of the charge-current type. Let us therefore check if we can also interpret the anomalous Zeeman effect in terms of precession. To do this we need a more detailed understanding of the magnetic potential and its vorticity.

7.2 The magnetic potential

7.2.1 Derivation

How do we deal with the kinetic energy of a moving charge in circular motion when we describe it as a stationary magnetic phenomenon? As we will show, it is done by expressing the kinetic energy in terms of a fake potential energy $U$. Potential energy is a scalar. As the moving charge corresponds to a current $qv$, which is a vector quantity, the only way to create a scalar quantity out of this current is to combine it with another vector quantity $A$ into a scalar product, e.g. $U = -q(v \cdot A)$. For the moment we consider $A$ as a general vector that has not yet been specified any further. In a constant magnetic field $B = Be_z$, with $B > 0$ the moving charge $q < 0$ whose velocity $v = ve_y$, $v > 0$ corresponds to a current $qv$, will perform a uniform circular motion at the cyclotron frequency $\omega_c > 0$, which in the non-relativistic limit is given by $\omega_c = -\frac{eB}{m}$. The velocity is then $v = -\frac{me}{q}B$. Let us rewrite $-qv \cdot A$ as $-qA \cdot v = -\frac{me}{q}B A \cdot v$. To make this correspond to the kinetic energy $p^2/2m_0$ we must thus have $-\frac{me}{q}B A \cdot v = p^2/qm_0 = v^2 = -\frac{q^2}{2m_0}re_y$. From this it follows that $A = \frac{1}{r}Be_y = -\frac{1}{r}v \times B$. This is exactly the magnetic vector potential that corresponds to a constant magnetic field $B$ as can be checked by calculating $B = \nabla \times A$. Using $A = -\frac{1}{r}v \times B$ it is easy to rewrite $U = -q(v \cdot A)$ as $U = \mu \cdot B$, where $\mu = -\frac{1}{2m_0}L$. This derivation does not require any consideration of a current loop. We see thus that the intuitive picture we use for this term in terms of a true potential energy is wrong, as anticipated under point (4) in Subsubsection 6.1.1.

The sign used in the expression $-qv \cdot A$ may surprise, but we must remind here the reason why we introduce the minimal substitution. In the absence of a magnetic field, the correct parameters to write the Lorentz transformation would be $(E - qV(r), cp)$. In a first non-relativistic approximation the quantity $E - qV$ we want to obtain becomes $E - qV \approx m_0c^2 + \frac{p^2}{2m_0} - qV = m_0c^2 + T - U$, just like in Lagrangian dynamics, where one justifies its introduction merely by showing that it makes things work. In the case of a purely magnetic field, there is no electric potential, such that this expression becomes then $m_0c^2 + \frac{p^2}{2m_0}$ in a frame wherein the centre of the orbit is at rest, such that it can be considered as a “static” magnetic situation. The expression we want to obtain requires thus adding the kinetic
energy, and this can be achieved by subtracting the “potential energy” of the current, which is the negative kinetic energy, as defined above.

Of course the preceding lines only explain the case when \( \mathbf{v} \parallel \mathbf{A} \), and not the cosine term in \( U = -q \left( \mathbf{v} \cdot \mathbf{A} \right) \). It is less easy to grasp the meaning of the cosine term in \( U = -q \left( \mathbf{v} \cdot \mathbf{A} \right) \). In principle, in a constant magnetic field \( \mathbf{v} \) must be parallel to \( \mathbf{A} \). Therefore, if \( \mathbf{v} \parallel \mathbf{A} \), the motion of charged particle must be a forced motion with respect to the magnetic field. Let us therefore consider a charge in uniform motion on a circular orbit with a velocity \( \mathbf{v} \) in a constant magnetic field \( \mathbf{B} = B_x \mathbf{e}_x + B_y \mathbf{e}_y \). This motion takes place in a plane perpendicular to \( \mathbf{B} \). We will then have \( \mathbf{A} = \frac{1}{c} \mathbf{v} \times \mathbf{B} = \frac{1}{c} B \times B_x \mathbf{e}_x - \frac{1}{c} B \times B_y \mathbf{e}_y \). Put \( \mathbf{A}_1 = -\frac{1}{c} B_z \mathbf{e}_x \), \( \mathbf{A}_2 = -\frac{1}{c} B_z \mathbf{e}_y \), such that \( \mathbf{A} = \mathbf{A}_1 + \mathbf{A}_2 \). It is then obvious that \( \mathbf{A} \cdot \mathbf{v} = \mathbf{A}_1 \cdot \mathbf{v} + \mathbf{A}_2 \cdot \mathbf{v} \). This explains why the part of the kinetic energy corresponding to the velocity \( \mathbf{v} \) that one can attribute to \( \mathbf{B}_1 = B_z \mathbf{e}_y \) is given by \( \mathbf{A}_1 \cdot \mathbf{v} \). If we consider \( \mathbf{A}_1 \) and we observe the circular orbit with velocity \( \mathbf{v} \), then only the part \( \mathbf{A}_1 \cdot \mathbf{v} \) of the kinetic energy can be attributed to \( \mathbf{B}_1 \). The rest must be attributed to the second force which forces the charged particle to follow an orbit in a plane that is not perpendicular to \( \mathbf{B}_1 \), and which in this analysis is based on \( \mathbf{B}_2 \). But in other situations, the origin of the force could be a different kind of field than a magnetic field \( \mathbf{B}_2 \), e.g. an electric field. It is only in such cases that a negative cosine term (leading to a “negative kinetic energy”) can have physical meaning. The field could also be varying with time rather than a constant field as in our example of \( \mathbf{B}_2 \).

The magnetic vector potential \( \mathbf{A} \) is often presented as a meaningless mathematical quantity that has just been introduced to simplify the calculations and whose use is justified because it makes things work. After the discovery of the Aharonov-Bohm effect, this viewpoint has been challenged by Feynman [12] who stated that \( \mathbf{A} \) is for quantum mechanics more significant than \( \mathbf{B} \). In relativity, \( \mathbf{cA} \) builds a four-vector with \( \mathbf{V} \), such that its meaning should be as physical as the meaning of \( \mathbf{V} \), and conceptually related to it. Despite all this, the vector potential has remained a concept that is not very intuitive. Here we have discovered a clear meaning for it. As often pointed out, \( \mathbf{A} \) is only defined up to a constant, just like \( \mathbf{V} \). Its meaning is thus indeed as clear as the meaning of \( \mathbf{V} \), and both quantities are quite intuitive. Several authors [13]-[14] have tried to make a case for this viewpoint, based e.g. on an experiment by Blondel [15].

7.2.2 Larmor frequency

The kinetic energy \( \frac{p^2}{2m_0} = \frac{1}{2} m_0 \omega_0^2 r^2 \) of the electron on the circular orbit can also be written in terms of the angular momentum \( L = m_0 \omega_0 r^2 \) as \( \frac{p^2}{2m_0} = \frac{1}{2} \omega_0 L \). To express the kinetic energy for a particle with an angular momentum \( L \) in the form \( L \omega_0 \), we must thus not use the true cyclotron frequency \( \omega_0 \) for \( \omega \), but the fictive Larmor frequency \( \omega_L = \omega_0/2 \). This quantity pops up in all quantum mechanical calculations. One may feel tempted to infer from spotting this quantity in the equations that the orbital rotational motion in the physical problem studied is happening at the frequency \( \omega_L \) instead of \( \omega_0 \), which is quite puzzling. One might ask oneself why the electron is turning slower on its orbit than one might expect based on classical mechanics. Is this just one more quantum mystery? The solution to this riddle is thus that \( \omega_L \) is only an auxiliary quantity to simplify the notations.

There is another way to highlight this point that the Larmor frequency is just an auxiliary concept. Larmor’s construction starts from the calculation of the fictitious forces in a rotating frame. In this frame there will be a fictitious centrifugal force and a fictitious Coriolis force. He then expresses that there should be a frequency of rotation that one can attribute to \( \mathbf{B}_1 = B_z \mathbf{e}_y \) as physical as the meaning of \( \omega_0 \). Its meaning is thus indeed as clear as the meaning of \( \omega_0 \), and both quantities are quite intuitive. Several authors [13]-[14] have tried to make a case for this viewpoint, based e.g. on an experiment by Blondel [15].

There is another way to highlight this point that the Larmor frequency is just an auxiliary concept. Larmor’s construction starts from the calculation of the fictitious forces in a rotating frame. In this frame there will be a fictitious centrifugal force and a fictitious Coriolis force. He then expresses that there should be a frequency of rotation where the magnetic field is completely canceled by the fictitious forces. This is the Larmor frequency. The Larmor frame “erases” the magnetic field. In this frame there will be no position and no velocity that make the particle feel the presence of the magnetic field or the fictitious Coriolis and centrifugal forces because all these forces cancel each other exactly. Of course in every point of the inertial frame the particle is not at rest but moving at a velocity \( \mathbf{v} = \mathbf{v} \) rather than being at rest with a velocity \( \mathbf{v} = \mathbf{0} \). The relation between the velocities in the two frames is \( \mathbf{v} = \mathbf{v} + \omega \times \mathbf{r} \).

It looks contradictory that the orbital motion of a particle in a magnetic field takes place at the cyclotron frequency, while we state that the particle does not feel the magnetic force in the frame that is rotating at the Larmor frequency. Following the idea that the particle does not feel the magnetic force it would appear that the particle could be at rest in the Larmor frame, while following from what we know in the inertial frame, the particle should appear to be moving at a residual frequency \( \omega_L = \omega_0 - \omega_L = \omega_0/2 \) in the Larmor frame. If the particle has a residual velocity in the Larmor frame, should it then not have to feel an attractive force that ensures that it stays on the circular orbit at this residual velocity? As Larmor’s construction shows, the answer from Newtonian mechanics, would have been no. This would lead to a contradiction. But relativity shows that there is also an electric field in the Larmor frame, due to the local Lorenz transformation (with boost vector \( \mathbf{v} \)) of the magnetic field in the inertial frame (see Eq. 22). This electric field exerts a local attractive and central force on the particle. It is this electric field that is responsible for the fact that the particle that is in cyclotron motion in the inertial frame is not at rest in the Larmor frame but moving at the residual frequency \( \omega_L = \omega_0 - \omega_L \).

That the Larmor frequency appears in the equations is due to the fact that we must divide all terms that occur in the squared Dirac equation by \( 2m_0 c^2 \) to be able to reduce it to the Pauli equation. We see here the same factor of
2 entering the calculations as the one that occurs in the derivation of the expression of the magnetic potential. It is the factor 2 that occurs in the expression $q^2/2mc$. And in both cases, the true frequency of the motion is $\omega_c$. An analysis of the original meaning of the Larmor frequency shows that drawing the conclusion that the true frequency would be $\omega_L$ rather than $\omega_c$, just because this is the quantity that comes to the fore in the calculation, is wrong because it fails to discern that the rotating frame introduces an electric field.

7.2.3 Larmor precession as the vorticity of the magnetic potential

The traditional minimal substitution is given by:

$$E \rightarrow E - qV, \quad p \rightarrow p - qA. \quad (24)$$

Non-relativistically we can write

$$p - qA = m_0(v - \frac{q}{m_0} A). \quad (25)$$

From this we can appreciate that $-\frac{q}{m_0} A(r)$ behaves as a velocity field $v_s(r)$. For a constant magnetic field $B$ we have

$$A = \frac{1}{2} r \wedge B. \quad (26)$$

On a first contact this looks a bit mysterious, because we can choose the origin at will and calculate $r$ with respect to this origin, the result will always be correct and independent from the choice of origin. But this fact is well known under the name of gauge invariance and expresses the fact that the vector potential is defined up to an arbitrary constant. As the scalar potential and the vector potential are related to each other by Lorentz transformation, it is evident that we cannot choose both arbitrary constants simultaneously at will, which is why they are related through a gauge condition. We will further discuss this arbitrary constant later on in our discussion. For the constant magnetic field, the velocity field $-\frac{1}{m_0} A(r)$ is of the type:

$$v_s(r) = +r \wedge \frac{q}{2m_0} B = -r \wedge \omega_L, \quad (27)$$

where we have introduced:

$$\omega_L = -\frac{q}{2m_0} B. \quad (28)$$

In other words, in the non-relativistic approximation, the velocity field is the same as the one we would observe in a frame that is rotating at an angular velocity $\omega_L$ corresponding to the Larmor frequency. According to the thoughts described above, we can consider this as a fictive auxiliary quantity. The rotating frame we discover here would then be rotating at a fictive frequency, while the true rotational motion would be the cyclotron frequency.

Let us now consider the velocity field $v(r)$ of a liquid and imagine that we would put a paddlewheel inside this liquid. The velocity field has vorticity and it will be make the paddlewheel turn as discussed in [16]. In this reference Auroux considers a circular path of radius $r$ on which he calculates the average velocity $v$ of the paddlewheel. He then takes the limit $r \rightarrow 0$. The paddlewheel becomes then infinitesimal. According to this calculation, the infinitesimal paddlewheel will rotate at a frequency:

$$\omega = \frac{1}{2} \nabla \wedge v(r). \quad (29)$$

When $v = \omega \wedge r$, such that the liquid bodily rotates with the frequency $\omega$, the paddlewheel will locally spin with the same frequency as the rotation of the liquid body. We have a daily-life confirmation of this idea on a merry-go-round. Such a merry-go-round is actually a better realization of the idea of a liquid that bodily rotates at a constant frequency than a real-life liquid where the angular frequencies might be radius-dependent. Let us associate the merry-go-round with a global rotating frame. When you are on a merry-go-round your personal local frame will not describe a circular motion with the directions of its axes $x, y, z$ fixed. The axes of your local frame are also rotating: Your local frame exhibits precession. Most of the time we gloss over of this fact by using a reference frame based on a local basis $e_x, e_y, e_z$ in a symmetry-adapted system of curvilinear coordinates, which already incorporates this effect. But the fact that this is a varying basis must be taken into account in the mathematics, and motivates the introduction of covariant derivatives. The argument of the merry-go-round just expresses Berry’s phase [17] on a circular orbit which is a geodesic.

Imagine now that you have a point-like charged particle without spin that is put into circular orbital motion within a magnetic field. It is then conceivable that the merry-go-round effect could give it spin, whereby the angular frequency of that spin would be the cyclotron frequency. That spinning motion would have a kinetic energy $L\omega_c/2 = L\omega_L$, where again the Larmor frequency is an auxiliary quantity. The spinning motion is intrinsic and if we wanted to describe the spinning point particle geometrically, we would have to describe it as a spinning point. But in geometry, points are not spinning. However, we can describe the spinning motion in terms of vorticity. We can treat the Larmor frequency
geometrically by using the same reasoning on \( \mathbf{v}_* \) as on \( \mathbf{v} \). We see thus that the "vorticity" of the velocity field of a constant magnetic field is such that a charged point-like particle will spin in the field at the cyclotron frequency. Any energy calculations that could be based on the equation \( E = \hbar \omega \) will have to be done using the Larmor frequency and this Larmor frequency can be calculated from:

\[
\omega_L = \frac{1}{2} \nabla \times \mathbf{v}_*(r) = -\frac{1}{2} \nabla \times \frac{q}{m_0} \mathbf{A}(r) = -\frac{q}{2m_0} \mathbf{B}.
\]  

(29)

Of course, in the case of a magnetic field, there is no real liquid that would push on the charge like on a paddlewheel, but we can obtain the results also without assuming the presence of a liquid. The idea of a liquid is only introduced to obtain a velocity field. As we have already a velocity field \(-\frac{q}{m} \mathbf{A}\), we can dispense with the introduction of a liquid to obtain the mathematical result derived. The idea is to describe precession induced by circular motion. Such precession is certainly considered in magnetism, and one even calculates relativistic corrections for it in terms of Thomas precession.

Within a frame in global rotation, the physical effects of rotation that affect the paddlewheel or the charge are independent from the choice of the origin for the local frame. This corresponds to the notion that the magnetic potential is defined up to a constant. Of course, the force that is needed to counterbalance the "centrifugal force" of the rotating frame is provided by the Lorentz force, and this "centrifugal force" will indicate what the true centre of the rotation is. For the calculation of the physical effects of precession however, it does not matter where the centre of the fictive merry-go-round is. We could take any point as the centre of rotation, it would not make any difference. We may also note that in the limit \( r \to 0 \), we reduce the circular current \( q \mathbf{v} \) to a point-like monopole, exactly according to the idea introduced above.

Let us further explore these ideas we used to define monopoles and vector potentials and make the radius of the circular motion of the charged particle in the magnetic field shrink to zero. What will happen then? As the non-relativistic cyclotron frequency is independent from the radius of the circular motion, this will leave us with a point-like spinning motion, even for a spin-less charged particle at rest! A charge in circular motion on an orbit with a diameter that is so small that we cannot see it with the naked eye will be a hidden motion. This hidden motion can be treated by the previous arguments, which lead us to the idea that a magnetic field could make a spin-less particle with charge \( q \) at rest spin at an angular frequency \( \omega_c = -\frac{q}{m} \mathbf{B} \). The point-like hidden motion can be treated as the interaction between the scalar charge \( q \) and the vorticity of magnetic potential \( \mathbf{A} \), such that we end up with an interaction of the electric charge with the magnetic field \( \mathbf{B} \). Energy calculations have to be done however with \( \omega_L = -\frac{q}{2m_0} \mathbf{B} \). However, this energy \( \hbar \omega_L \) will not be the kinetic energy of the orbit shrunk to zero because this is also zero. It is the energy due to the precession. The moment of inertia that intervenes in the precession is not related to a mass distribution that corresponds to a point mass at a distance \( r \) from the centre of the circular orbit. It is the moment of inertia of the mass distribution inside the spinning top that visualizes the spinning electron (see Subsection 7.3). Simultaneously, a point-like monopole is not the magnetic charge of a circular current loop whose radius shrinks to zero. In taking this limit, \( v \to 0 \), such that there is no current or magnetic charge left, and no true magnetic monopole. As also \( L \to 0 \), there is also no magnetic moment left. It is the precession that corresponds to the magnetic monopole. The quantity \( cq \) can be symbolically identified with a monopole \( q_m \), like we have done. The concept of the magnetic monopole is useful to distinguish a geometrical point modeling a point charge from a spinning point charge. We may note that the precession terms which correspond to the anomalous Zeeman effect and the spin-orbit coupling are thus both related to the hidden rotational energy of the hidden rotational motion. This explains thus why the terms in Eq. 17 exist and how they can be associated with magnetic monopoles, such that their classification is correct.

We may finally note why the merry-go-round metaphor fails for the spin-orbit term. In fact, for a circular motion in a central Coulomb field \( \mathbf{F} = \frac{q\mathbf{v}^2}{r^2} \mathbf{r} \), the rotational frequency \( \omega(r) \) is not a constant, but depends on \( r \), such that the image of a liquid that is bodily rotating no longer holds. The consequence hereof is that the paddlewheel will no longer rotate at the same frequency as the liquid.

### 7.3 Precession

Precession changes the rest mass of a particle. In [2] we have shown that the Dirac equation can be derived from the Rodrigues equation (Eq. 8 in Section 9) by putting \( \varphi = \omega r \) in the rest frame of the electron and making the assumption \( \frac{\hbar}{2m_0} = \gamma m_0 c^2 \). The assumption was introduced in a hand-waving way by assuming that the rest mass of the electron would correspond to the kinetic energy stored in its spinning motion.

The derivation proposed in [2] of the Dirac equation is entirely classical, which is a very surprising fact, as the Dirac equation is the core of the whole machinery of quantum mechanics. The Schrödinger equation can be derived from it. How can these equations then possibly be classical? It is explained in [2] that the properties of quantum mechanics that make it different from classical mechanics are only due to the way we use the Dirac and Schrödinger equations in the calculations when we apply them to specific problems. In fact, in following the motto "shut up and
calculate" we introduce unwittingly features into the algebra that are meaningless from the viewpoint of the classical meaning of the algebra. One of those features that are puzzling is the superposition principle. Just as adding rotation matrices does not lead to a new rotation matrix, adding spinors has \textit{a priori} not a clear meaning. Another of the weird things that we cannot understand classically is the quantization of spin and angular momentum. The algebra of the Dirac equation forces the spin and the angular momentum to align with the magnetic field, as discussed above, and it is difficult to understand classically why it could not be otherwise.

Let us here thus just accept the fact that the spin must be aligned with the magnetic field, like quantum mechanics tells us. In fact, when we consider in the calculations the possibility that the spin or the angular momentum are not aligned, we find that the spin or angular momentum must precess, but that this does not lead to a constant energy. To recover a constant energy, one must introduce another force, and the motion becomes then forced with respect to the magnetic field. If we do not draw this force into the calculations, they are incomplete and it is then vain to try to understand them. What it would imply that the spin is not aligned without forcing, such that the energy is not constant is not clear, but this problem is in a sense eluded by the alignment condition. We may thus assume that the treatment ceases to be classical at the moment we are accepting the alignment condition.

We must now point out that we already know the physical meaning of the anomalous Zeeman effect. In fact, it is now time to remember that the Larmor precession term \( \omega L \cdot \sigma = -\frac{e \hbar}{2m_0} B \cdot \sigma \) we obtained from the discussion in Subsection 7.2 is identical to the term obtained from quantum mechanics in the non-relativistic limit of the Dirac equation for an electron in a magnetic field. When the particle is rotating in its rest frame the precession frequency \( \omega_c \) will add up algebraically to the rotation frequency \( \omega_0 \) of the particle, changing the apparent frequency of its rotation, which is why it eventually entails a correction for the energy \( E = \frac{\hbar}{2m_0} \rightarrow E = \hbar (\omega_0 \pm \omega_c)/2 = m_0 c^2 \pm \hbar \omega_c \), where the \( \pm \) sign is due to the existence of spin up and spin down states. This gives a very neat explanation for the anomalous Zeeman splitting for an electron at rest. The very important point that follows from this argument is that the amplitude of the anomalous Zeeman effect in the Dirac theory must be strictly identical to that of the orbital Zeeman effect, because on orbit the precession is in phase with the orbital motion due to the merry-go-round effect, and the precession for the particle at rest is obtained from the precession during the orbital motion by letting the radius of the orbit shrink to zero. Moreover the anomalous Zeeman effect is not due to the spin of the electron, but due to \textit{the additional spin} given to the electron by the interaction of its charge with the magnetic field.

### 7.4 Conclusion about the anomalous Zeeman effect

We see thus that the theory can calculate the energy values of the two equilibrium states \( m_0 c^2 \pm \frac{e \hbar B}{2m_0} \) of the spinning electron in terms of a charge-dipole interaction, without associating a magnetic dipole with the spin. The idea is thus that a magnetic field makes any charge turn due to the electron spin in the calculation of the equilibrium state. Simultaneously this copes with Lorentz's objection that the hypothetical magnetic dipole moment of the electron is too large to allow for an explanation in terms of a current loop inside the electron: There is simply no spin-associated magnetic dipole moment in the formalism.

We may note that the correct theory for ferromagnetism also does not rely on magnetic dipoles, but on Heisenberg's mechanism of an exchange interaction which is based on the Pauli principle and the Coulomb interaction. In the Dirac formalism, charge and spin occur in mere juxtaposition, without blending into a more complex quantity like a dipole. The interaction is just defined by the charge, while the formalism shows that the spin has to line up with the magnetic field \( B \), because the wave function must be an eigenstate of the pseudo-vector operator \( B \cdot \sigma \) or \( B \cdot \gamma \).

The mere juxtaposition of spin and charge in the Dirac equation looks similar to that in the exchange mechanism. The image of a dipole moment is just not present in the algebra of the Dirac equation. One may speculate about defining a magnetic dipole moment \( \mu_c \), because it flatters our intuition of a little magnet, but this idea that \( -\frac{e \hbar}{2m_0} B \cdot \sigma \) must willy-nilly correspond to a magnetic dipole moment is a preconceived notion that is just flawed mathematically.

---

\footnote{The idea that the charge turns means that it is a magnetic monopole according to Section 4. This is in conformity with the fact that the anomalous Zeeman term \( e q \mu B \) in Eq. 21 corresponds to the term that describes the interaction of the electromagnetic field with a monopole at rest in Eqs. 16-17. The whole paper with its apparently meandering narrative is thus constructed around proving this connection and the underlying physical imagery. The rest of the paper is devoted in a similar way to the connection between the interaction of the moving magnetic monopole with the electromagnetic field and the spin-orbit coupling.}
As already pointed out above, the appropriate interpretation is that there are three types of interaction: charge-charge, dipole-dipole, and charge-dipole. The anomalous Zeeman effect is a charge-dipole interaction and one should refrain from identifying this charge-dipole interaction with a dipole-dipole interaction. There is thus always a precession energy associated with the presence of a magnetic field \( B \), which is the reason why we find this term already in Eq. 21, which does not contain spin.

After all this, there is yet another argument in favor of the interpretation of the term \(-\frac{\mu}{2m}B \cdot \sigma\) proposed in this paper. Dirac’s theory works only well for the leptons [18], as the true value of \( q \) for the electron is given by: \( g_e = 2.00231930436153(53) \). It works almost as well for the muon \( g_\mu = 2.003318416(13) \), and much less well for the neutron \( (g_n = 3.82608545(90)) \) and the proton \( (g_p = -5.585694713(56)) \). It is tempting to assume that in the cases of the proton and the neutron, a true dipole moment \( \mu \) due to internal currents might intervene\(^9\) and give rise to some term \(-\mu \cdot B\), while this is just not the case for the electron, which is truly a point particle or is so small that the effect of such currents, if they exist, can be neglected, and only the charge term \(-\frac{\mu}{2m}B \cdot \sigma\) plays a significant rôle.

8 Problems with the traditional derivation of the spin-orbit coupling

8.1 Preliminary remarks

The spin-orbit term \( -\frac{1}{m} \frac{\partial}{\partial t} L \cdot \sigma \) is easily shown to be equal to \( \frac{2}{3} [(v \wedge E) \cdot \sigma] \), where \( U = qV \) is the potential energy.\(^10\) Following the discussion in Subsubsection 6.1.3 the spin-orbit interaction is of the “dipole”-charge type, due to the presence of the term \( qV \wedge E \). Here \( qV \) represents the “dipole” and \( E \) contains the charge. Also the interpretation of \( \sigma \) in \( L \cdot \sigma \) in terms of a “spin operator” \( \Delta = \frac{1}{2} \sigma \) is not appropriate. The term \( L \cdot \sigma \) within \( \frac{1}{m c^2} \frac{\partial}{\partial t} L \cdot \sigma \) just represents the angular momentum. Imaging the spin-orbit coupling as a “dipole”-dipole interaction containing a term \( L \cdot \Delta \) is in conflict with the symmetry. First of all \( \frac{1}{m c^2} \frac{\partial}{\partial t} L \cdot \sigma \) is a vector rather than a scalar. Secondly, its “dipole”-charge interaction symmetry is not compatible with a “dipole”-dipole interaction symmetry. An analogous problem of an over-interpretation of an operator that violates the symmetry occurs in the definition of helicity \( u \cdot \sigma \), with \( u = p/p \) for neutrinos in particle physics.

The frequency \( \omega_T \) of the Thomas precession correction is given [9,10] by:

\[
\omega_T = \gamma^2 \frac{1}{\gamma + 1} (v \wedge a). \tag{30}
\]

In an electric field \( E \), this can be written as \( \gamma \gamma \frac{1}{\gamma + 1} m c^2 \gamma v \wedge qE \). For \( \gamma \approx 1 \) this corresponds to \( \frac{1}{2m c^2} v \wedge qE \). To correct the spin-orbit term for the Thomas precession we must subtract \( h \omega_T \) from its absolute value. This has thus absolutely nothing to do with an electric dipole moment induced by the relativistic motion of a magnetic dipole moment associated with the spin of the electron, as has often been claimed [11]. As we have argued all along, such a hypothetical magnetic dipole moment does not even exist in the context of the Dirac equation. The Thomas precession is a relativistic correction term that contributes to the global spin-orbit precession along the orbit. The global expression is equal to the Thomas precession with the reversed sign.

8.1.1 Simple derivation of the expressions for the Thomas precession

For what will follow it might be useful to explain carefully what Thomas precession is. There are actually two equations for the Thomas precession that one can prove. The first one summarizes the conceptual definition of the effect, which is that the composition of two boosts is a composition of a boost and a rotation: \( B(v_2)B(v_1) = \mathbf{R}(\mathbf{n}, \vartheta_T)B(v) \). The rotation axis \( \mathbf{n} \) is here perpendicular to both \( v_1 \) and \( v_2 \), such that it is just the rotation angle \( \vartheta_T \) and the boost

\[^9\] These currents are the currents produced by the quarks. It is impossible to explain the magnetic moment of the neutron whose overall charge is zero, without falling back on its quark structure.

\[^{10}\] It may be noted that in the classical Coulomb problem, \( v \) can be calculated from \( r \) from both conservation laws (for energy and angular momentum). The kinetic energy \( \frac{1}{2} m v^2 = E - U \) follows just from the total energy \( E \) and the potential energy \( U \). But using the law of conservation of energy ceases to be feasible within the present quantum mechanical context. The problem is that the electron might have radiated to settle down on a given orbit. The total energy \( E \) of the electron on this orbit is now the quantity we want to determine in the first place. If there is no further information available as to the amount of energy that has been radiated away, it is a priori impossible to know \( E \). The lacking information can however be derived from the conservation of angular momentum. Specifying \( v \) through \( L = r \wedge m v \) in the non-relativistic limit permits to calculate \( v \). From this it is possible to calculate \( E = U(r) + \frac{1}{2} m v^2 \). It is for these reasons that we express everything in terms of \( U \) and \( L \). But in a purely relativistic context, the equation \( \hat{L} = r \wedge m v \) introduces also \( m \), which is related to the total energy through \( E = mc^2 \).
parameter \( \mathbf{v} \) that have to be calculated. But specifying the Thomas precession requires only the calculation of \( \vartheta_T \).
The second equation formulates the effect of Thomas precession on an orbit, and states that \( \frac{d\vartheta_T}{dt} = \frac{\gamma^2}{(\gamma+1)c^2} \mathbf{v} \wedge \mathbf{a} \).

The following derivation of the first equation follows the argument in Subsection VIII D of [9]. We note \( \gamma = \cosh w \), \( \beta \gamma = \sinh w \), \( \beta = \tanh w \). We have then \( \cosh \frac{w}{2} = \sqrt{\frac{1 + \gamma}{2}} \), \( \sinh \frac{w}{2} = \sqrt{\frac{\gamma - 1}{2}} \), \( \tanh \frac{w}{2} = \sqrt{\frac{\gamma - 1}{\gamma + 1}} \). In SL(2, \( \mathbb{C} \)), a boost with velocity \( \mathbf{v} = w \mathbf{u} \) is given by \( \mathbf{B}(\mathbf{v}) = \cosh \frac{w}{2} \mathbf{I} - \sinh \frac{w}{2} [\mathbf{u} \cdot \sigma] \). Here \( \mathbf{u} \) is the unit vector parallel to \( \mathbf{v} \). We must now calculate \( \mathbf{B}(\mathbf{v}_2)\mathbf{B}(\mathbf{v}_1) \).

For simplicity, we can take \( \mathbf{u}_1 = \mathbf{e}_x \), \( \mathbf{u}_2 = \cos \alpha \mathbf{e}_x + \sin \alpha \mathbf{e}_y \). We find then:

\[
\mathbf{B}(\mathbf{v}_2)\mathbf{B}(\mathbf{v}_1) = \left( \cosh \frac{w_2}{2} \cosh \frac{w_1}{2} + \sinh \frac{w_2}{2} \sinh \frac{w_1}{2} \cos \alpha \right) \mathbf{I} \\
- i \sinh \frac{w_2}{2} \sinh \frac{w_1}{2} \sin \alpha [\mathbf{e}_z \cdot \sigma] - \cosh \frac{w_2}{2} \sinh \frac{w_1}{2} [\mathbf{e}_y \cdot \sigma] \\
- \sinh \frac{w_2}{2} \cosh \frac{w_1}{2} \left[ (\cos \alpha \mathbf{e}_x + \sin \alpha \mathbf{e}_y) \cdot \sigma \right].
\]

This must be equal to the product \( \mathbf{R}(\vartheta_T, \mathbf{e}_z) \mathbf{B}(\mathbf{v}) = \left( \cos \frac{\vartheta_T}{2} \mathbf{I} - i \cos \frac{\vartheta_T}{2} [\mathbf{e}_y \cdot \sigma] \right) \left( \cosh \frac{w}{2} \mathbf{I} - \sinh \frac{w}{2} [\mathbf{u} \cdot \sigma] \right) \) of a rotation around the \( z \)-axis and a boost:

\[
\mathbf{R}(\vartheta_T, \mathbf{e}_z) \mathbf{B}(\mathbf{v}) = \cos \frac{\vartheta_T}{2} \cosh \frac{w}{2} \mathbf{I} - i \sin \frac{\vartheta_T}{2} \cosh \frac{w}{2} [\mathbf{e}_y \cdot \sigma] \\
- \cos \frac{\vartheta_T}{2} \sinh \frac{w}{2} [\mathbf{u} \cdot \sigma] - \sin \frac{\vartheta_T}{2} \sinh \frac{w}{2} [\mathbf{e}_y \cdot \sigma].
\]

By identifying the parts containing the unit matrix \( \mathbf{I} \) and the parts containing \( i[\mathbf{e}_y \cdot \sigma] \) one obtains:

\[
\tan \frac{\vartheta_T}{2} = \frac{\tanh \frac{w}{2} \sin \alpha}{1 + \tanh \frac{w}{2} \cos \alpha}.
\]

which is Eq. 145 in reference [9].

The second equation can be derived from this equation. But one can calculate the identity \( \frac{d\vartheta_T}{dt} = \frac{\gamma^2}{(\gamma+1)c^2} \mathbf{v} \wedge \mathbf{a} \) also directly, by considering the identity: \( \mathbf{B}(\mathbf{v}_2)\mathbf{B}(\mathbf{v}) = \mathbf{R}(\mathbf{n}, \vartheta_T) \mathbf{B}(\mathbf{v} \oplus \mathbf{v}_2) \), where \( \mathbf{v} \oplus \mathbf{w} \) denotes the boost vector associated with the composition of boosts \( \mathbf{B}(\mathbf{w}) \mathbf{B}(\mathbf{v}) \). We are considering here only \( \mathbf{dv}_\perp \) as collinear boosts do not lead to Thomas precession. By using Taylor expansions one can show that to first order \( \mathbf{B}(\mathbf{v}_\perp) = \mathbf{1} - \frac{1}{2} \mathbf{dv}_\perp \left( [\mathbf{e}_z \cdot \mathbf{u}] \cdot \sigma \right) \).

One obtains then \( \mathbf{B}(\mathbf{dv}_\perp)\mathbf{B}(\mathbf{v}) = \sqrt{\frac{2+\gamma}{2}} \mathbf{I} - \frac{1}{2} \sqrt{\frac{2+\gamma}{2}} \mathbf{dv}_\perp \left( [\mathbf{e}_z \cdot \mathbf{u}] \cdot \sigma \right) - \sqrt{\frac{2-\gamma}{2}} \mathbf{dv}_\perp \left( [\mathbf{e}_y \cdot \sigma] \right) \). Here \( \gamma \) is the Lorentz factor that corresponds to \( \mathbf{v} \). The identification of this result with Eq. 32 yields then:

\[
\frac{d\vartheta_T}{dt} = \frac{\gamma^2}{(\gamma+1)c^2} \mathbf{v} \wedge \mathbf{a}.
\]

(wherby we have replaced of course \( \vartheta_T \) by \( d\vartheta_T \)). From this one obtains \( \frac{d\vartheta_T}{dt} = \frac{\gamma^2}{(\gamma+1)c^2} \mathbf{v} \wedge \mathbf{a} \) in the co-moving frame, i.e. the rest frame of the electron. Here \( \tau \) is the proper time of the electron, i.e. the time in the co-moving frame. Taking into account the time dilatation this yields \( \frac{d\vartheta_T}{dt} = \frac{\gamma^2}{(\gamma+1)c^2} \mathbf{v} \wedge \mathbf{a} \) in the laboratory frame. Here \( \mathbf{a} \) takes the same value in the laboratory frame and in the co-moving frame because it is perpendicular to \( \mathbf{v} \). This is presumably the simplest derivation possible of the expression for the Thomas precession. It avoids using hyperbolic geometry to derive this result, as was done in reference [9]. We can also see from this derivation why the Dirac equation with the minimal substitution cannot be used to derive the Thomas precession. As discussed in Subsection 8.2 (and explained in reference [2]), the minimal substitution accounts for the instantaneous value of \( \mathbf{B}(\mathbf{v}) \) in (\( \mathbf{r}, t \)), but does not take care of \( \mathbf{a}_\perp \) or \( \mathbf{R}(\vartheta_T, \mathbf{e}_z) \).

8.2 Generalization of Dirac’s minimal substitution for the case of a moving charge

As already stated in Subsection 5.2, Dirac’s minimal substitution is not general enough as it does not account for the interactions of the current \( \mathbf{q} \mathbf{v} \) of the moving electron with the electromagnetic potential. It is explained in [2] that in the context of the Dirac equation, the minimal substitution must not be seen as an abstract rule that one copies mechanically from Lagrangian dynamics and that one can justify with hindsight by the fact that it works. The goal of the minimal substitution is to introduce the true instantaneous kinetic parameters \( (\gamma, \gamma \mathbf{v} / c) \), that will permit us to write the true instantaneous Lorentz transformation for the time. As explained in [2], the primary idea is that the the transformation of the time \( ct' = \gamma(ct - \mathbf{v} \cdot \mathbf{x} / c) \) under a free-space Lorentz boost can also be written using the parameters \( (E, cp) = m_0 c^2 (\gamma, \gamma \mathbf{v} / c) \) and \( m_0 c^2 \) instead of \( (\gamma, \gamma \mathbf{v} / c) \). In fact, the Dirac equation is derived
from the Rodrigues equation by transforming \((\frac{d}{dt}, 0, 0, 0) \rightarrow \frac{d}{dt}, \frac{q}{c}, \frac{d}{dt}, \frac{q}{c})\) according to the boost that transforms \((m_0 c^2, 0, 0, 0) \rightarrow (E, cp)\).\(^{11}\) But in a potential \(V\), the rest energy of a particle is no longer \(m_0 c^2\) but \(m_0 c^2 + qV\), such that the parameters \((E, cp)\) of a moving particle are no longer the correct set of kinetic parameters to describe the instantaneous Lorentz transformation, because \(E\) is no longer purely kinetic. To be able to write the correct Lorentz transformation, one must know the “true kinetic energy”, i.e. that part of the total energy that is not potential energy. This leads to the substitution \(E \rightarrow E - qV, p \rightarrow p - qA\). The idea is that for an electron at rest in an electric potential, the equation is \(-\frac{d}{d\tau}\gamma \psi = (m_0 c^2 + qV)\psi\), where \(\tau\) is the time in the rest frame. By generalizing this to a general frame by Lorentz covariance we obtain then arguably the general Dirac equation with the minimal substitution \(E \rightarrow E - qV, p \rightarrow p - qA\).

The problem is here that \((qV, cqA)\) is not exactly what one would obtain from \((qV, 0)\) by transforming it to a general frame. Whereas the four-potential \((V, cA)\) is Lorentz covariant, the quantity \(q\) is not, because \(q\) is part of a charge-current four-vector \((\gamma q, \gamma qV/c)\). Therefore \((qV, cqA)\) is not the correct Lorentz covariant generalization for the expression that must be used in the most general substitution. In the most general substitution we would need in stead of \(qV\) and \(qA\) the terms that we can obtain by considering: \([\gamma q \mathbb{I} + \gamma q \frac{V}{c} \mathbf{\sigma}] [V \mathbb{I} + cA \mathbf{\sigma}]\). The result will lead to two terms \(qV\) and \(qcA\) with the same sign. This is needed because the terms \(qV\) and \(qcA\) are used with the same signs in the traditional minimal substitution. Furthermore, we obtain a term \(\gamma q (v \cdot A) \mathbb{I}\), which after the substitutions \(E \rightarrow E - qV, p \rightarrow p - qA\), will lead to a term \(-\gamma q (v \cdot A) \mathbb{I}\), whose sign is in conformity with the discussion in Subsubsection 7.2.1. It may look confusing that in the derivation of Eq. 13 and Eq. 34 we have not used the strategy of alternating signs as would follow from the rules that prevail for SL(2,\(\mathbb{C}\)). For Eq. 13 this may be due the way the magnetic potential has been historically defined. For Eq. 34, we note that we can make juxtapostions like \([\gamma q \mathbb{I} + \gamma q \frac{V}{c} \mathbf{\sigma}] [V \mathbb{I} + cA \mathbf{\sigma}]\) with any number of terms and any choice of signs. The goal is to check to what kind of multi-vectors this leads. This is more the case here and the choice made here leads to the correct results for the further use. These considerations lead to:

\[
[\gamma q \mathbb{I} + \gamma q \frac{V}{c} \mathbf{\sigma}] [V \mathbb{I} + cA \mathbf{\sigma}] = 
\gamma q V \mathbb{I} + \gamma q c A \mathbf{\sigma} + \gamma q \frac{V}{c} V \mathbf{\sigma} + \gamma q (v \cdot A) \mathbb{I} + \gamma q (v \wedge A) \mathbf{\sigma}.
\] (34)

These are the quantities that one would have to use in the substitution that generalizes the minimal substitution.

8.3 How we obtain the spin-orbit-coupling term from the Dirac equation with the generalized substitution

The generalized substitution was given in Eq. 34. The term \(\gamma q \frac{V}{c} V \mathbf{\sigma}\) in Eq. 34 is a vector and cannot contribute to the potential energy. However, after squaring the Dirac equation, which will produce the announced variant \([\frac{d}{d\tau} \mathbb{I} - \nabla \mathbf{\sigma}] [\gamma q \mathbb{I} + \gamma q \frac{V}{c} \mathbf{\sigma}] [V \mathbb{I} + cA \mathbf{\sigma}]\), it will lead to a term containing \([\nabla \mathbf{\sigma}] [\gamma q \frac{V}{c} V \mathbf{\sigma}]\). For \(A = 0\), this will lead in the non-relativistic limit to a term \(\frac{1}{2} (\mathbb{I} \nabla (\mathbb{I} V)) \mathbb{I}\) and a term \(\frac{1}{2} (\mathbb{I} \nabla (V \wedge A) \cdot \mathbf{\sigma})\). The first term leads to \(-\frac{1}{2} (v \cdot E) \mathbb{I} + \frac{v}{c} \nabla \cdot v \mathbb{I}\). In the former term we can recognize the power term in Eq. 21. The calculation of the term \(\frac{1}{2} (\mathbb{I} \nabla (V \wedge A) \cdot \mathbf{\sigma})\) yields \(\frac{1}{2} [V (\nabla \wedge v) + v \wedge E]\). For uniform circular motion \(\nabla \wedge v = 2i\omega\). The term \(V (\nabla \wedge v)\) becomes then \(2i\omega E = -2i v \wedge E\). In total we have thus \(V (\nabla \wedge v) + v \wedge E = -v \wedge E\), such that \(\frac{1}{2} (\mathbb{I} \nabla (\mathbb{I} V)) \mathbb{I} - \frac{1}{2} (\mathbb{I} \nabla (V \wedge A) \cdot \mathbf{\sigma})\). After multiplication by \(-\frac{\hbar}{2m_0 c}\) the term that goes with \(v\) becomes \(-\frac{\hbar}{2m_0 c} \mathbb{I} (\frac{1}{2} v \wedge E)\), which is the spin-orbit precession in an electric field, but without the correction for Thomas precession.

8.4 Apparent failure with respect to the traditional derivation

If we derive the Pauli equation from the Dirac equation with the generalized substitution, it will thus also contain the spin-orbit term, with the correct sign, but without the correction for Thomas precession. Traditionally, the spin-orbit term is derived from the Dirac equation with the minimal substitution \(E \rightarrow E - qV, p \rightarrow p - qA\). This seems to entirely discredit our approach as it makes it look as though it contradicts our criticism that this substitution does not take into account the velocity of the electron. By combining the traditional approach with the present approach it may even seem that we get the result for the spin-orbit coupling twice. But as already stated previously, all this is not true because the traditional approach is wrong and it is not possible at all to derive the spin-orbit term from the Dirac equation based on the traditional minimal substitution. Let us now explain why.

\(^{11}\) This shows that the Dirac equation does thus not account for the Thomas precession by construction. This may look as a startling claim to the reader, but it will be part of a discussion that we will unfold from Subsection 8.4 onwards.
8.5 Errors in the traditional derivation

8.5.1 How a change of basis offers a first hint about the existence of an error

Let us point out in which way the traditional derivation is based on flawed mathematics. The way we are writing the Dirac equation in this paper is different from the traditional one. In the Weyl representation we have:

\[
e_t \leftrightarrow \gamma^t = \begin{pmatrix} 1 & \mathbb{1} \\ \mathbb{1} & -1 \end{pmatrix}, \quad e_5 \leftrightarrow \gamma^5 = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix},
\]

while in the traditional representation we have:

\[
e_5 \leftrightarrow \gamma^5 = \begin{pmatrix} 1 & \mathbb{1} \\ \mathbb{1} & -1 \end{pmatrix}, \quad e_t \leftrightarrow \gamma^t = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}.
\]

The difference just corresponds to a change of basis in \( \mathbb{R}^5 \) provided with a metric \( x_1^2 + x_2^2 - x^2 - y^2 - z^2 \), whereby we change the orientation of the ct-axis between the axes \( OX_4 \) and \( OX_5 \) in the \( OX_5X_4 \) plane, e.g. by a rotation over \( \frac{\pi}{4} \). We can consider space-time as embedded into the five-dimensional space, and its mathematical properties do not depend on the way it is embedded: Meaningful mathematical properties do not depend on a choice of basis. The two representations are therefore equivalent by a similarity transformation. This argument actually indicates how one can try to prove Pauli’s theorem that all choices of gamma matrices are equivalent. Equivalent choices of gamma matrices correspond just to different choices of a basis. Now, in the Weyl representation, the Dirac equation for an electron in a Coulomb field will be:

\[
\left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} - \frac{\hbar}{i} \nabla \cdot \sigma = \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} + \frac{\hbar}{i} \nabla \cdot \sigma \right) \begin{pmatrix} \psi_1 \\ \psi_2 \end{pmatrix}
= m_0 c \begin{pmatrix} \psi_1 \\ \psi_2 \end{pmatrix},
\]

We have thus:

\[
\left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} + \frac{\hbar}{i} \nabla \cdot \sigma \right] \psi_2 = m_0 c \psi_1,
\]

\[
\left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} - \frac{\hbar}{i} \nabla \cdot \sigma \right] \psi_1 = m_0 c \psi_2,
\]

which leads to:

\[
\left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} + \frac{\hbar}{i} \nabla \cdot \sigma \right] \left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} - \frac{\hbar}{i} \nabla \cdot \sigma \right] \psi_1 = m_0^2 c^2 \psi_1,
\]

\[
\left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} - \frac{\hbar}{i} \nabla \cdot \sigma \right] \left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} - \frac{qV}{c} \right) \mathbb{1} + \frac{\hbar}{i} \nabla \cdot \sigma \right] \psi_2 = m_0^2 c^2 \psi_2.
\]

The point is now that these equations are completely decoupled. The terms that do not contain \( V \) are e.g. \( \left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} \right) \mathbb{1} + \frac{\hbar}{i} \nabla \cdot \sigma \right] \left[ \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} \right) \mathbb{1} - \frac{\hbar}{i} \nabla \cdot \sigma \right] \psi_1 = \left[ \hbar^2 (\Delta - \frac{1}{c^2} \frac{q^2}{\sigma^3}) \right] \psi_1 \), for the first equation. The terms that combine \( \left( -\frac{\hbar}{i} \frac{\partial}{\partial ct} \right) \mathbb{1} \) and \( \frac{\hbar}{i} \nabla \cdot \sigma \) give rise to: \( \frac{\hbar}{c} \frac{qV}{c} \cdot \frac{qV}{c} \), where we have assumed that \( V \) does not vary with time. The interesting terms with vector symmetry are those that combine \( \frac{\hbar}{i} \nabla \cdot \sigma \) and \( \frac{\hbar}{i} \nabla \cdot \sigma \). They give rise to \( \frac{\hbar}{c} \left[ E \cdot \sigma \right] \psi_1 \). There is thus a term \( \left[ E \cdot \sigma \right] \psi_1 \), but no term containing \( \left[ (E \cdot p) \cdot \sigma \right] \psi_1 \). There is even not a term containing \( \left[ p \cdot \sigma \right] \psi_1 \), because the two terms containing \( \frac{\hbar}{c} \left[ \nabla \cdot \sigma \right] \) have opposite signs and cancel in the algebra. This is an exact result. There is here no fuss about approximations or series expansions. The vector operators working on \( \psi_1 \) are all based on vector quantities in the plane of motion, while \( (E \cdot p) \cdot \sigma \) is perpendicular to it. The term \( (E \cdot p) \cdot \sigma \) can thus never occur in the algebra if it is carried out correctly.

The operator \( \frac{\hbar}{c} \frac{qV}{c} \cdot \frac{qV}{c} \) will admittedly lead to a vector term that is perpendicular to the plane of motion, that becomes equal to \( -i \left[ \omega \cdot \sigma \right] = -i \omega [s \cdot \sigma] \) for a particle at rest. Here \( s \) is the spin axis, which is identified with \( e_2 \), while the plane of motion is identified with the \( Oxy \) plane. After the necessary algebra this term will lead to a change \( m_0 \to m_0 + qV/c^2 \) for the rest mass or \( h \omega_0 \to h \omega = h \omega_0 + qV \) for the rest energy of the particle in a potential. As explained in Subsection 8.2, the purpose of the minimal substitution was exactly to reproduce this result. Also the term \( \frac{\hbar}{i} \frac{qV}{c} \frac{\partial}{\partial ct} \) does thus not incorporate the spin-orbit coupling.
As the traditional representation is equivalent to the Weyl representation up to a similarity transformation, this conclusion must also be valid in the traditional representation. Note that the difference between the Weyl representation and the traditional interpretation does not affect the coordinates \((x, y, z)\), as it corresponds only to a change of the orientation of the time axis in the \((x_4, x_5)\)-plane. The two representations must thus lead to the same results and to the same expressions for the results. This suggests that something with the traditional derivation of the spin-orbit term must be wrong.

We can further argue that the only way to derive the terms for the spin-orbit coupling correctly from a Dirac equation is using the extended substitution we introduced in Eq. 34. In fact, to obtain a term \(|(E \wedge p) \cdot \sigma \psi_j|\) one needs a succession \(|\nabla \cdot \sigma V[p \cdot \sigma] \psi_j|\). This can never be obtained from a succession of two \(2 \times 2\) SL(2, \(\mathbb{C}\)) operators like the one that occurs in Eq. 39. A succession of three such operators is needed. With the extended substitution one obtains the succession \(|\nabla \cdot \sigma V[v \cdot \sigma] \psi_j|\) after squaring the Dirac equation, and this will lead to the required spin-orbit term. The traditional approach is unable to accomplish this because it misses the presence of \(\gamma \left[ \frac{\partial L}{e} \cdot v \cdot \sigma \right]\) in the minimal substitution.

8.5.2 The origin of the error: the solutions are mixed states

Nevertheless, the traditional approach presents a derivation of the spin-orbit term whereby it even seems to account correctly for the Thomas precession. This traditional derivation of the spin-orbit coupling is a piece of wrong algebra that by good fortune produces the correct physical result desired. As explained in the preceding lines, it starts from the wrong minimal substitution which can never yield the desired result because it does not contain the interactions with the current. It then introduces a second error to obtain the result that agrees with the experimental observations by brute force. What the second error is can be discovered by considering the Dirac equation for an electron at rest in the absence of any electromagnetic field. In the Weyl representation this leads to:

\[
\left[ -\frac{i}{\hbar} \frac{\partial}{\partial ct} \right] \psi_1 = m_0 c \psi_1, \\
\left[ -\frac{i}{\hbar} \frac{\partial}{\partial ct} \right] \psi_2 = m_0 c \psi_2,
\]

and after “squaring” to:

\[
\left[ -\hbar^2 \frac{\partial^2}{\partial c^2 t^2} \right] \psi_1 = m_0^2 c^2 \psi_1, \\
\left[ -\hbar^2 \frac{\partial^2}{\partial c^2 t^2} \right] \psi_2 = m_0^2 c^2 \psi_2,
\]

A viable simultaneous solution of Eqs. 40 and 41 is:

\[
\psi_1 = \psi_2 = e^{-i m_0 c^2 t / \hbar} \begin{pmatrix} 1 \\ 0 \end{pmatrix}.
\]

In the traditional representation, the Dirac equation becomes:

\[
\left[ -\frac{i}{\hbar} \frac{\partial}{\partial ct} \right] \psi_1 = m_0 c \psi_1, \\
\left[ -\frac{i}{\hbar} \frac{\partial}{\partial ct} \right] \psi_2 = -m_0 c \psi_2.
\]

This needs no further squaring as the equations are already decoupled. One possible solution is:

\[
\psi_1 = e^{-i m_0 c^2 t / \hbar} \begin{pmatrix} 1 \\ 0 \end{pmatrix}, \quad \psi_2 = e^{+i m_0 c^2 t / \hbar} \begin{pmatrix} 1 \\ 0 \end{pmatrix}.
\]

What transpires from the solution in the traditional presentation is that the spinor is a mixed state. It contains two pure states that lead to the same rest energy \(m_0 c^2\) for the electron, one with the electron spinning counterclockwise and one with the electron spinning clockwise around the \(z\)-axis:
The mixed state must of course still be normalized. In the mixed state each pure state has thus the same probability \( \frac{1}{2} \). In fact, the rotation angle \( \omega_0 t \) is an algebraic quantity. Feynman found out also that one has to use such mixed states composed of pure states with opposite sign and equal probabilities in quantum electrodynamics. In following the tradition to associate negative-energy states with anti-particles, the mixed state contains particles and anti-particles with equal probabilities.

Feynman wondered what the meaning of this would be. The solution of that riddle is that the mixed states describe statistical ensembles of electrons. Negative frequencies already occur in SU(2), which is just Euclidean geometry and does not contain anti-particles. There is therefore absolutely no necessity to associate negative frequencies with anti-particles. It is more appropriate to associate negative frequencies with particles that spin the other way around than those that give rise to positive frequencies. Both signs of the frequency will however yield the same kinetic rotational energy, and therefore the same rest mass and energy for the electron. Considering such a mixed state makes thus perfect physical sense, while the mixed state based on anti-particles is puzzling.

Of course, the interpretation of the negative frequencies we are proposing here clashes once more acrimoniously with currently accepted notions. But in reference [2] we give a derivation of the Dirac equation, that does not rely at all on the existence of anti-particles. Charge conjugation symmetry is just not part of the picture. As we explain in reference [2] we can consider anti-particles in a later stage, and then the traditional arguments can be used to show that we should associate them with negative frequencies. But this is then a different representation than the original one. The situation is a little bit like that with the two different versions of SL(2,C). Just as the two versions of SL(2,C) are related one to another by parity transformation and should not be merged into a single SL(2,C) formalism, the two particle and anti-particle representations are related by charge-conjugation and should not be merged into a single formalism, because a negative frequency would then acquire two different, mutually exclusive interpretations. Moreover in the merged formalism, the rest energies of a positron and an electron are truly considered as opposite, such that they would add up to zero, while they should add up to twice 511 keV. Of course, the alternative interpretation of the negative frequencies puts the whole discussion about Majorana and Dirac neutrinos in a different context.\(^\text{12}\)

Also in the case of the Weyl representation the states are mixed, even if it is less obvious here because the two pure states carry the same sign for \( \omega_0 t \). But in the Weyl representation, the equation for the pure state of a single electron in its rest frame is:

\[
\psi = e^{-im_0c^2t/\hbar} \begin{pmatrix} 1 \\ 0 \\ 0 \\ 0 \end{pmatrix} + e^{im_0c^2t/\hbar} \begin{pmatrix} 0 \\ 0 \\ 1 \\ 1 \end{pmatrix}.
\]

(45)

The mixed state must of course still be normalized. In the mixed state each pure state has thus the same probability \( \frac{1}{2} \). In fact, the rotation angle \( \omega_0 t \) is an algebraic quantity. Feynman found out also that one has to use such mixed states composed of pure states with opposite sign and equal probabilities in quantum electrodynamics. In following the tradition to associate negative-energy states with anti-particles, the mixed state contains particles and anti-particles with equal probabilities.

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Also in the case of the Weyl representation the states are mixed, even if it is less obvious here because the two pure states carry the same sign for \( \omega_0 t \). But in the Weyl representation, the equation for the pure state of a single electron in its rest frame is:

\[
\begin{pmatrix} \frac{-\hbar}{c} \frac{\partial}{\partial \tau} \mathbb{I} \\ \frac{\hbar}{c} \frac{\partial}{\partial \tau} \mathbb{I} \end{pmatrix} \begin{pmatrix} \psi \\ \psi^{-1\dagger} \end{pmatrix} = m_0c \begin{pmatrix} s \sigma & s \sigma \end{pmatrix} \begin{pmatrix} \psi \\ \psi^{-1\dagger} \end{pmatrix}.
\]

(46)

Here the \( \psi \) is a two-column SL(2,C) matrix representing the spinning electron, while \( \psi^{-1\dagger} \) represents the spinning electron in the SL(2,C) representation of opposite handedness; \( \tau \) is the proper time. The block-diagonal \( 4 \times 4 \) matrix \( D(g) \) with the blocks \( \psi \) and \( \psi^{-1\dagger} \) on the diagonal is the Weyl representation of a group element \( g \) obtained from an even number of reflections and represents the spinning electron. This equation can never lead to the Dirac equation, because the right-hand side can never be simplified according to:

\[
\begin{pmatrix} s \sigma & s \sigma \end{pmatrix} \begin{pmatrix} \psi \\ \psi^{-1\dagger} \end{pmatrix} = m_0c \begin{pmatrix} \psi \\ \psi^{-1\dagger} \end{pmatrix}.
\]

(47)

In fact, let us divide out \( m_0c \) on both sides. Then on the left-hand side the non-zero blocks are off-diagonal, while on the right-hand side they are diagonal. The two sides have different symmetries and they can therefore never be identical. In the traditional representation this kind of observation becomes hidden by the fact that there are no longer vanishing blocks. Due to the fact that the simplification expressed in Eq. 47 is not possible, one is forced to adopt a linear combination of single-electron states to obtain the Dirac equation. This linear combination is (see reference [2]):

\[
M = D(g) + \begin{pmatrix} s \sigma & s \sigma \end{pmatrix} D(g).
\]

(48)

Furthermore, it is impossible to derive the Dirac equation for a particle with zero rest mass, because in the derivation given in reference [2] it is necessary to consider a particle in its rest frame where \( 2m_0c^2 = \hbar \omega_0 \). In reality neutrinos do have mass, which further wrong-foots the discussion about Majorana and Dirac neutrinos, because the wave functions of the traditional Dirac neutrinos are considered to be just two-component Weyl spinors, while neutrino wave functions that are solutions of a Dirac equation with non-zero rest mass must be four-component mixed states, just like the so-called bi-spinor electron wave functions.
and we will have then:

\[
\begin{pmatrix}
  s \cdot \sigma \\
  s \cdot \sigma
\end{pmatrix} M = M.
\]

This shows that also in the Weyl representation the solutions of the Dirac equation are mixed states.

Mixed states do not have a direct obvious meaning in the pure group theory. E.g. the sum of two rotation matrices is not a rotation matrix, and therefore the sum of two spinors is \textit{a priori} not a new spinor. Group elements cannot be added, they can only be multiplied. Linear combinations of representations of group elements do not represent new group elements. They belong to the so-called group ring. One can give such a mixed state however a meaning by considering it as defining a statistical ensemble, just as is done in the traditional interpretation of such mixed states in quantum mechanics. We can thus consider the free-space solution as degenerate, and the presence of electromagnetic fields can lift this degeneracy.

We may note that in the simpler formalism of SU(2), the solution of the eigenvalue equation \( \frac{d}{dt} [s \cdot \sigma] \psi = \pm \frac{\hbar}{2} \psi \) for a spin aligned with the unit vector \( s \) is also a mixed state. The not normalized “spinor” \( \psi \) is the first column of the sum \( S = I + s \cdot \sigma \) of the two group elements \( I \) and \( s \cdot \sigma \). For this sum, we will have \([s \cdot \sigma] S = S\). In other words: \( \psi = (I + s \cdot \sigma) \psi_1 \), where \( \psi_1 \) is the first column of \( s \cdot \sigma \). This leads indeed to \( \psi = [s \cdot \sigma] \psi \). When \( s = e_z \), the mixed nature of \( \psi \) becomes concealed after normalization by the fact that \( \psi_1 \) and \( [s \cdot \sigma] \psi_1 \) accidentally take the same numerical values. This is also further discussed in reference [2].

8.5.3 Problems in taking the Schrödinger limit of the mixed states

Now, the non-relativistic Schrödinger limit is obtained by removing the rest mass from the equation. If we want to get rid of the rest mass in the Weyl representation, we can define the classical wave function:

\[
\begin{pmatrix}
  \psi_1^c \\
  \psi_2^c
\end{pmatrix} = e^{im_0c^2t/\hbar} \begin{pmatrix}
  \psi_1 \\
  \psi_2
\end{pmatrix}.
\]

After such a substitution, the rest mass will be removed from the calculations. But using the same trick in the traditional representation (see e.g. [6], page 65) will lead to a meaningless statistical ensemble containing states with rest masses 0 and 2\(m_0\). The correct substitution to be used in the traditional representation is thus:

\[
\begin{pmatrix}
  \psi_1^c \\
  \psi_2^c
\end{pmatrix} = \begin{pmatrix}
  e^{im_0c^2t/\hbar} \\
  e^{-im_0c^2t/\hbar}
\end{pmatrix} \begin{pmatrix}
  \psi_1 \\
  \psi_2
\end{pmatrix},
\]

rather than Eq. 50, because this is the substitution that leads to a meaningful statistical ensemble. This substitution would look absurd if we thought that the state given by Eq. 45 is a pure state. But for the mixed state, this substitution does make sense. This difference between the substitutions needed in Eqs. 50 and 51 just reflects the difference between the choices \(\gamma_4\) and \(\gamma_5\) for the gamma matrix that has to be associated \(\frac{1}{\hbar}\) \(\partial/\partial t\) in the algebra. It is actually much more logical to write the Dirac equation in the traditional representation as:

\[
\begin{bmatrix}
-\frac{\hbar}{i} \frac{\partial}{\partial t} & I \\
-\frac{\hbar}{i} \nabla \cdot \sigma & \frac{\hbar}{4} \nabla \cdot \sigma
\end{bmatrix} \psi = m_0 c \psi,
\]

thereby considering \(-\frac{\hbar}{i} \frac{\partial}{\partial t} \gamma_0\) rather than \(-\frac{\hbar}{i} \frac{\partial}{\partial t}\) as the energy operator that generalizes the prescriptions that are valid in the context of the Schrödinger equation. All this is of course a major watershed because it invalidates the whole philosophy of small and large 2 \times 2 representations in the Dirac equation. However, there are just two \(\operatorname{SL}(2, \mathbb{C})\) representations in a 4 \times 4 representation based on gamma matrices, and these two \(\operatorname{SL}(2, \mathbb{C})\) representations are mathematically of “the same size”. As can be seen from Subsection 2.2, they are defined in a completely symmetrical way. They are just of different handedness. If we refuse to accept this, then the Weyl representation would have two representations that are equally large, while the traditional representation would have a small and a large one. Why should such differences all at once pop up in the application of the mathematics if the physics in the two choices of gamma matrices must be the same?

It is further very important to realize that it is absolutely not necessary to make the substitution of Eqs. 50 or 51 in order to obtain the non-relativistic limit, as we showed above. One can do the calculations fully relativistically. It suffices then to consider afterwards the limit whereby \(v\) becomes small and to subtract \(m_0 c^2\) from the final result. But when we do the calculations that way, we will never get a term \(2m_0\) into the algebra. It is the division by the term \(2m_0\) in the equations that one obtains from the wrong substitution which leads to the illusion that the algebra can describe the Thomas half correctly. This shows that it is not possible to obtain the Thomas half from the Dirac formalism, and that this is true both in the Weyl and in the traditional representation!
Of course it could be argued that we could solve Eq. 43 by taking another solution than Eq. 44, whereby we keep \( \psi_1 \) and put \( \psi_2 = 0 \). We would then obtain a pure state. But this solution will also no longer lead to a term \( 2mc \) in the algebra, and it could only be used for an electron at rest.

Finally, the traditional approach suggests that up to a certain order the expression for the Thomas precession in terms of the quantity \( v \wedge E \) would be generally valid and not depend on the geometry of the orbit, while the further developments will show that it is in many instances necessary to assume that the motion is uniform and circular.

8.5.4 Methodological remarks

The conclusion that the traditional calculation of the spin-orbit coupling is wrong is very unsettling. The reason why it has not been noticed that this error crept in, is that in the traditional approach the Dirac equation has been guessed. This means that we use it without knowing on what kind of precise assumptions it is built. In the approach described in reference [2], the equation has been derived. This enables discerning a number of issues that in the traditional approach are not even suspected. It is very hard to suspect that the solutions of the Dirac equation have to be mixed states. And it remains difficult to spot that this leads to errors creating the illusion that the spin-orbit coupling would be treated correctly by the Dirac equation, even when one knows the underlying assumptions. All this reveals the limits of the nonchalant advice that one should just “shut up and calculate” because it would “work”. But contrary to the claims, it does not work. As seen here, following the advice in a blindfolded way can lead to errors of appreciation. Spotting such errors and teasing out their consequences can prove a very difficult task.

Of course it is not easy to decide which approach is correct and which one is wrong when we pit two different approaches one against another. It is in this respect always better to give several arguments in support of a conclusion, in order to prove the internal consistency of the logic. Here we are giving four arguments.

(1) The first one is based on the fact that the solutions of the Dirac equation are mixed states. This may be less convincing to the reader because he does not know the results derived in reference [2], but again these results have an internal consistency.

(2) The second one is based on the fact that the approach based on the Weyl representation should yield the same result as the approach based on the traditional representation. This is the argument that should be really clear to all readers.

(3) A third one is based on the fact that the derivation of the Dirac equation ignores instantaneous accelerations. One should indeed not be surprised that the calculation does not take into account the Thomas precession. As mentioned above and in Subsubsection 7.2.1, the Dirac equation is derived by using only boost parameters, like e.g. \((E, c\mathbf{p})\) in the case of the free-space equation. It considers only instantaneous boosts, not instantaneous accelerations. The equation makes sure that we get the instantaneous local boost correct in every point \((r, t)\). Thomas precession can only occur in a sequence of non-collinear boosts. It is thus the result of an instantaneous acceleration perpendicular to the velocity. An instantaneous boost as used to derive the Dirac equation can thus not account for Thomas precession. This is also discussed in further detail in [2].

(4) One can appreciate the point that the Dirac equation with a four-potential only treats boosts and not Thomas precession also as follows. Defining a general Lorentz transformation requires six independent parameters. The electromagnetic four-potential is defined by four parameters, but they are linked by the Lorentz gauge condition, such that the four-potential contains three independent parameters. These three parameters define the boost part of the Lorentz transformation, while the three remaining independent parameters of the general Lorentz transformation correspond to the rotational part of it. These can thus not be defined by the Dirac equation with a four-potential, because there are just no parameters in the equation that would permit defining them.

A weak point of the Dirac equation is thus that it is covariant with respect to boosts, but not with respect to instantaneous accelerations that are perpendicular to the instantaneous velocity.

8.6 A more physical approach to the spin-orbit coupling

8.6.1 Road map

Thomas precession is by definition an effect in the co-moving frame. It is a mis-appreciation of the correct clock rate of the electron due to the fact that the co-moving frame is rotating. According to Purcell’s calculation of the Thomas precession [10] the Thomas precession corresponds exactly to the difference of the merry-go-round effects in the laboratory frame and in the co-moving frame. The two merry-go-round effects are different due to Lorentz contraction (or time dilatation). Of course effects of the motion on the rest mass must be made in the co-moving frame, as it is in this frame that the electron is at rest. These effects must then be transformed back to the laboratory frame. But whereas Purcell’s calculation gives the correct algebraic result, it does not tell us why we can make the calculation the way he does. Why do we need to consider the difference between two merry-go-round effects?
A similar problem exists for the other part of the spin-orbit effect, the part that occurs (up to a proportionality factor) in Eq. 21 and is not corrected for Thomas precession. In our derivation of Eq. 21 it is mere algebra. As this is not satisfactory, one would like a physical explanation for it. Such a classical derivation for it has been proposed in Subsection 5.1. But also here the argument developed leads to the correct algebraic result, while it is not completely explained what the idea is behind the calculation. We may wonder e.g. why one should consider the magnetic field experienced by the electron in the co-moving frame in the first place, while we want to calculate results in the laboratory frame. Is it not possible to make a calculation in the laboratory frame right ahead? In the calculation only the magnetic part of the electromagnetic field in the co-moving frame intervenes. Due to our previous discussion of the anomalous Zeeman effect, we understand very well the effects of the magnetic field on the electron in its rest frame. In this calculation we only consider the instantaneous boost. This result must of course be back-transformed to the laboratory frame. What remains to calculate, is the effect of the accelerations perpendicular to the instantaneous velocity, and this is the Thomas precession. The Thomas precession due to the magnetic field can to a first approximation be neglected. What remains is thus the effect of the component of the electric field that is perpendicular to the instantaneous velocity, and that will be the Thomas precession.

In the following we will talk a lot about phases corresponding to the merry-go-round effect as opposed to Berry phases. We will however not consider such phases accumulated over a whole loop, but rather the instantaneous rates of change of them, because these correspond to the idea of precession. What we want to explain are the following points:

1. One can consider two types of precession, a merry-go-round effect in position space \( \mathbb{R}^3 \) (which is not curved) and a precession in velocity space. The velocity space is curved and helps actually to visualize the group manifold. The precession on this curved space yields the Berry phase after an integration over a closed loop.
2. The instantaneous rate of change in phase due to the merry-go-round effect in position space corresponds to the part of the spin-orbit effect without Thomas precession. It is the part that occurs in Eq. 21.
3. This instantaneous rate of change in phase due to the merry-go-round effect in position space does not correspond to the exact rate of change of the geometrical phase due to parallel transport. The exact calculation of the latter can be made by first considering the merry-go-round effect in position space and then making a correction to it.
4. The correction we have to carry out to obtain the correct rate of change of geometrical phase is the Thomas precession. According to Purcell’s calculation, it is the difference between the merry-go-round effects in velocity space in the co-moving frame and in the laboratory frame. Perhaps it is worth pointing out that there is no real merry-go-round effect on the group. When we make a closed orbit on the group, we are getting back to the Lorentz transformation that is identical to the one we started from, such that the phase difference can only be a multiple of \( 2\pi \). That we have to worry about the Thomas precession is due to the fact that we are dealing with a closed orbit in \( \mathbb{R}^3 \) rather than on the group and that we have constructed the Dirac equation by only taking into account the boost part of the Lorentz transformations (see Footnote 11).

In a merry-go-round calculation one basis vector of the co-moving frame remains always aligned with the instantaneous velocity. The problem is thus that such a continuous alignment with the tangent to the orbit in position space does in general not correspond to parallel transport. It is possible to get a feeling for this by an analogy with the motion of a triad of basis vectors along a closed loop on the surface of a sphere. The closed loop could e.g. be a small circle. Let us assume we have defined spherical coordinates \((r, \theta, \phi)\) and local frames with basis vectors \(e_r, e_\theta, e_\phi\) as usual, such that the \(z\)-axis corresponds to \( \theta = 0 \). In a trip with uniform velocity \( v \) along a small circle defined by \( \theta = \theta_0 \neq 0 \), the geometrical phase will not be given by \( 2\pi \) as one could be tempted to conclude from the permanent alignment of \( e_\phi \) with the instantaneous velocity \( v \). This permanent alignment corresponds to the merry-go-round effect, and does not account correctly for the value of the geometrical phase. As shown by the Gauss-Bonnet theorem, the geometrical phase will be given by \( \Omega = 2\pi(1 - \cos \theta_0) \), where \( \Omega \) is the solid angle subtended by the small circle. Only when \( \theta_0 = \frac{\pi}{2} \) will the procedure of aligning \( e_\phi \) with \( v \) yield the correct result, because then the path is a geodesic. The calculation \( \Omega = 2\pi(1 - \cos \theta_0) \) reproduces exactly what happens e.g. in Foucault’s pendulum. This will be further developed in Subsubsection 8.6.2.

We may note that the calculation of the global Wigner rotation over a closed path in reference [9] is based on a theorem of hyperbolic geometry that just rephrases the Gauss-Bonnet theorem. This shows very clearly that the angle built up by Thomas precession along a path in velocity space is the Berry phase. As the hyperbolic geometry is the geometry of velocity space, we are getting here a first clue for the existence of two precession effects. One in the hyperbolic velocity space, and one in position space. Instead of a path in a curved space of positions, Thomas precession occurs on a path in a curved space of velocities. This curved space represents actually the parameters that define an element of the Lorentz group. The velocities are the boost parameters and correspond to the boosts, the Berry angles correspond to the rotations.

In summary, we can state that the Dirac equation is wrong because it does not account properly for the parallel transport, such that it gets the Berry phase wrong. The reason for this is that the Dirac equation just calculates the merry-go-round effect in position space. This idea is already present in reference [2], but I was just unable to find the appropriate words to verbalize it. In reference [2], it was also discussed how strange it is to postulate that the wave
function is a function. It means that the Berry angle over a given path must be multiple of 2π and it is this feature that leads to quantization. But phases differences that are a multiple of 2π occur only on geodesics. Hence, quantum mechanics postulates that the orbits must be geodesics with respect to the spinning motion on the group manifold, and are therefore quantized. General relativity postulates that orbits must be geodesics with respect to displacement motion. This also implies rotations to a certain extend, but it does not consider the rotation due to the spin. The fully correct picture would thus imply that the orbits must be geodesics with respect to the combined effect of rotation and translation.

### 8.6.2 The Berry phase on the surface of a sphere

The merry-go-round effect on a circle with radius $r$ in the $Oxy$ plane can be obtained by considering the infinitesimal angle $\Delta \theta = \Delta r / r$. In vector form this can be rewritten as $\Delta \theta \mathbf{e}_z = r \Delta r / r^2$. This leads to $\omega = \frac{d}{dr} \mathbf{e}_z = \frac{r}{r} \mathbf{v} \wedge \mathbf{v}$. But such a calculation corresponds only to the correct value for the geometrical phase in flat space (which is here the $Oxy$ plane). It is no longer true if we consider the geometrical phase along a small circle with radius $r$ on the surface of a sphere with radius $R$, as it does not account for the curvature of the surface of the sphere.

How the correct calculation of the geometrical phase runs in curved space can be illustrated on this example of a motion along a small circle with radius $r$ on a sphere with radius $R$. This case study corresponds exactly to description of Foucault's pendulum. Let us take the rotation axis of the Earth as the $z$-axis. The locally vertical direction on the surface of the sphere is given by the vector $\mathbf{e}_R$. The precession of Foucault's pendulum can be trivially calculated from $(\omega \mathbf{e}_\theta) \mathbf{e}_R$, which is the component of $\omega = \omega \mathbf{e}_z$ along $\mathbf{e}_R$. This shows that the calculation in curved space cannot be made by using $\frac{1}{r} \mathbf{v} \wedge \mathbf{v}$, whereby $\mathbf{v}$ would be the position vector of the pendulum with respect to the centre of the small circle.

In the calculation one must take $dr = dr \mathbf{e}_\phi$, where $dr = R \cos \theta$, and $d\phi = r d\phi$. The term $dr / r$ must then be multiplied by $\mathbf{e}_\theta \mathbf{R} / R = \mathbf{e}_\phi$, rather than $r / r$, because what one wants to calculate is the rotation in the local tangent plane spanned by $\mathbf{e}_\phi$, $\mathbf{e}_\theta$. This is a rotation around the local vertical defined by the vector $\mathbf{e}_R$. This calculation leads to $\mathbf{e}_\theta \mathbf{R} \wedge \mathbf{e}_R = d\phi \mathbf{e}_R$. The integral over the circle of this quantity yields $2\pi \cos \theta \mathbf{e}_z$. The de-phasing is then $\Delta \phi = 2\pi - 2\pi \cos \theta$, or $2\pi (1 - \cos \theta) \mathbf{e}_z$ in vector form. This is equal to $\int \mathbf{E} \cdot d\mathbf{S}$, where $d\mathbf{S} = R^2 \sin \theta d\theta d\phi \mathbf{e}_R$, in agreement with the Gauss-Bonnet theorem. The quantity $\Delta \phi$ is the Berry phase, which can be expressed as $\oint A \cdot d\mathbf{r}$, where the vector quantity $\mathbf{A}$ is analogous to a vector potential. Putting $\int \mathbf{E} \cdot d\mathbf{S}$ along $\mathbf{e}_\theta R$ permits to rewrite $\Delta \phi = \oint (\mathbf{A} \wedge \mathbf{A}) \cdot d\mathbf{S} = \oint A \cdot d\mathbf{r}$. Here $dr = R \sin \theta d\phi \mathbf{e}_\phi$, such that one must thus take $\mathbf{A} = \frac{\varepsilon_0}{2} \frac{q}{R} \mathbf{e}_\phi$ to obtain the correct result. Using the general expression for $\mathbf{A} \wedge \mathbf{A}$ in spherical coordinates one can verify that this value of $\mathbf{A}$ leads indeed to $\mathbf{V} \wedge \mathbf{A} = \frac{q}{2\pi R^2} \mathbf{e}_R$. As $\mathbf{A}$ is a vector, one is tempted to interpret its as a vector potential.

The vector $\mathbf{A}$ shows many similarities with a magnetic vector potential. But of course it is not a magnetic vector potential, because there are no magnetic fields in our problem. The vector $\mathbf{A}$ does thus not define a magnetic monopole. This can be seen from the theorem of Gauss for an electric field: $\int \mathbf{E} \cdot d\mathbf{S} = \frac{4\pi}{c^2} \rho (\mathbf{r})$. This is the sum of two surface integrals $\int \mathbf{E} \cdot d\mathbf{S} = \frac{4\pi}{c^2} \rho (\mathbf{r})$, and $\int \mathbf{E} \cdot d\mathbf{S} = \frac{4\pi}{c^2} \rho (\mathbf{r})$ over the two areas $S_1$ and $S_2$ on both sides of the small circle, and which together make up the full sphere. The result for one of the two areas is then perfectly analogous with the calculation we have performed above. As there is by definition only an electric field $\mathbf{E} = \frac{q}{4\pi \varepsilon_0} \mathbf{r}$ within this problem, the quantity $\mathbf{A} = \frac{q}{2\pi} \mathbf{e}_\phi$, which leads to $\mathbf{V} \wedge \mathbf{A} = \mathbf{E}$, is not a magnetic potential. We have always $\nabla \cdot \mathbf{E} = \frac{q}{4\pi \varepsilon_0}$, with $\rho (\mathbf{r}) = q \delta (\mathbf{r})$. For all points different from the origin, the fact that $\mathbf{E} = \nabla \wedge \mathbf{A}$ leads to $\nabla \cdot (\mathbf{V} \wedge \mathbf{A}) = 0$. It can be checked indeed that $\nabla \cdot \mathbf{E} = 0$ for $\mathbf{r} \neq 0$ by using the general expression for the divergence in spherical coordinates. The pitfall is now to conclude from this that $\oint (\mathbf{V} \wedge \mathbf{A}) \cdot d\mathbf{S} = \int \mathbf{V} \cdot (\mathbf{V} \wedge \mathbf{A}) \cdot d\mathbf{r} = 0$. In fact, one expects $\nabla \cdot (\mathbf{V} \wedge \mathbf{A}) = 0$ by analogy with $\mathbf{a} \cdot (\mathbf{a} \wedge \mathbf{b}) = 0$, but this is not true at the origin where $\mathbf{A}$ has a singularity. In putting $\nabla \cdot (\mathbf{V} \wedge \mathbf{A}) = 0$, we forget the singularity in $\nabla \cdot \mathbf{E}$ at the origin. As $\oint (\mathbf{V} \wedge \mathbf{A}) \cdot d\mathbf{S} = \oint \nabla \cdot \mathbf{E} \cdot d\mathbf{r}$, we obtain the correct result when we use $\nabla \cdot \mathbf{E} = \frac{q \delta (\mathbf{r})}{2\pi \varepsilon_0}$. The same reasoning can be applied to the magnetic charge $q_m \delta (\mathbf{r}) = cq \delta (\mathbf{r})$ following the analogy developed in Footnote 4.

The mathematical difficulties that arise here are due to the abstraction of representing the charge distribution by $q \delta (\mathbf{r})$. The delta “function” can be described as a limit of test charge distributions $\lim_{\mathbf{r} \to \mathbf{r}_0} \rho_m (\mathbf{r})$. The problem is that for integration and differentiation we are not allowed to assume that the derivative of the limit will be the limit of the derivative, or the integral of the limit will be the limit of the integral: $\lim_{\mathbf{r} \to \mathbf{r}_0} \int f (\mathbf{r}) \rho_m (\mathbf{r}) d\mathbf{r} \neq \int \left[ \lim_{\mathbf{r} \to \mathbf{r}_0} f (\mathbf{r}) \rho_m (\mathbf{r}) \right] d\mathbf{r}$, and $\lim_{\mathbf{r} \to \mathbf{r}_0} D [f (\mathbf{r}) \rho_m (\mathbf{r})] \neq D [f (\mathbf{r})] \lim_{\mathbf{r} \to \mathbf{r}_0} \rho_m (\mathbf{r})$. Changing the order of the operations is in general not a valid procedure. This error occurs in Dirac’s definition of the “delta function”, and now also here in the differentiation procedure, where it gives rise to the confusion that $\nabla \cdot (\mathbf{V} \wedge \mathbf{A})$ would be zero at $\mathbf{r} = 0$. The limits must be defined in the sense of distributions, and then these difficulties will disappear. In fact, $\nabla \cdot (\mathbf{V} \wedge \mathbf{A})$ is not zero at the origin, even though $\lim_{\mathbf{r} \to 0} \nabla \cdot (\mathbf{V} \wedge \mathbf{A} (\mathbf{r})) = 0$. The correct result for the example of the electric monopole is $\nabla \cdot (\mathbf{V} \wedge \mathbf{A}) = \nabla \cdot \mathbf{E} = \frac{q}{2\pi} \delta (\mathbf{r})$.
\[ \mathbf{v} \cdot \mathbf{B} = 0 \] when there is no monopole. This tallies then exactly with the meaning of the equation \( \nabla \cdot \mathbf{B} = 0 \), which expresses that there are no magnetic monopoles.

The introduction of the Dirac string is illogical and runs contrary to the principle of Occam’s razor. Based on symmetry reasons one wonders if magnetic monopoles could exist. As we have shown in Section 4, the equations contain already for each term its symmetric counterpart. The fields and potentials of the electric and the magnetic monopoles have exactly the same mathematical structure. In the further development one stumbles then on a result \( \nabla \cdot \mathbf{E} = q_\delta \mathbf{r} \mathbf{r} \), where it had been \( \nabla \cdot \mathbf{B} = 0 \) without the monopoles. This result presents a golden opportunity to enhance the symmetry even further by rendering this completely analogous to \( \nabla \cdot \mathbf{E} = \frac{1}{\gamma} \delta \mathbf{r} \mathbf{r} \). But instead of accepting this tried and proved solution one rejects it and postulates that \( \nabla \cdot \mathbf{B} = 0 \) should remain universally valid. This choice is mathematically wrong, it breaks the symmetry, and to cover up for the error, one is then forced to introduce the stunning concept of a Dirac string. It is all together an egregious procedure and boils down to inventing “new physics”, just to explain away a trivial mathematical paradox inherent to the use of singular distributions.

8.6.3 Spin-orbit coupling without Thomas precession

There are several steps in the following demonstration that the merry-go-round effect is equivalent to the part of the spin-orbit coupling that occurs in Eq. 21. We will neglect here often the relativistic \( \gamma \)-factors, relativistic effects on the mass and consider only uniform circular motion. From the treatment it will transpire that for non-circular motion the calculations could be off the mark.

1. According to Subsubsection 7.2.1, \( \nabla \wedge \frac{p}{2} = \nabla \wedge q \mathbf{a} = q \mathbf{B} \). Furthermore Eq. 28 shows that in continuing to neglect relativistic modifications in the mass, we have \( \nabla \wedge \frac{p}{2} = m_0 \omega \). Combining the two we obtain \( \omega = \frac{qB}{m_0} \).

2. For circular motion in cylinder coordinates we have \( \nabla \wedge \frac{p}{2} = \frac{1}{r} \frac{d}{dr} e_z \).

3. For uniform circular motion we can put \( \frac{d\phi}{dt} = \omega \). We have then \( L = m_0 \omega^2 \). From this it follows that \( \frac{1}{2r} \frac{dl}{dt} = m_0 \omega = \frac{1}{2r} \).

4. The calculation of the merry-go-round effect yields \( \omega = \frac{1}{2r} \mathbf{r} \wedge \mathbf{v} \), such that \( m_0 \omega = \frac{1}{2r} \mathbf{r} \wedge \mathbf{p}_0 \).

5. From (1)-(4) it follows that the merry-go-round effect leads to \( m_0 \omega = q \mathbf{B} = \nabla \wedge q \mathbf{a} \). In deriving the Pauli equation we divide this by \( 2 m_0 \), such that the frequency that intervenes in the energy calculations is the Larmor frequency \( \frac{qB}{2m_0} \).

6. When there is no magnetic field in the laboratory frame, the electric field will yield a magnetic field in the co-moving frame which is given by \( \mathbf{B}' = \gamma \frac{1}{2} \mathbf{v} \wedge \mathbf{E} \). We neglect here the factor \( \gamma \). The calculation of the merry-go-round effect in the co-moving frame for an electron that has a relative velocity \( \mathbf{v} = \gamma v \gamma v \mathbf{c} / c \), such that \( \gamma^2 - (\gamma v \gamma v / c)^2 = 1 \). This the equation of a hyperboloid in \( \mathbb{R}^3 \). In formal analogy with what has been done for the calculation of the Berry phase on a sphere, we will introduce here hyperbolic coordinates \( (R, u, \phi) \) for points \( (X, Y, Z) \) in \( \mathbb{R}^3 \). The transformation between the Cartesian coordinates \( (X, Y, Z) \) and their corresponding hyperbolic coordinates is given by: \( Z = R \cosh u \), \( X = R \sinh u \cos \phi \), \( Y = R \sinh u \sin \phi \). This can be considered as an abstract change of coordinates, whereby the real meaning of \( (X, Y, Z) \) is irrelevant. This change of coordinates is not a 1-1-mapping between \( \mathbb{R}^3 \) and \( \mathbb{R}^3 \) as for spherical coordinates, as it implies that \( Z^2 - X^2 = Y^2 > R^2 > 0 \). The coordinate transformation is thus only defined for points inside the cone \( Z^2 - X^2 = Y^2 > 0 \). For the special choice \( (X_0, Y_0, Z_0) = (\gamma v_z / c, \gamma v_y / c, \gamma) \), we have then: \( \gamma = \cosh u \), \( \gamma v_z / c = \sinh u \cos \phi \), \( \gamma v_y / c = \sinh u \sin \phi \) such that the velocity space is just the surface \( Z^2 - X^2 = Y^2 = 1 \) (corresponding to \( R = 1 \)) in the vector space \( \mathbb{R}^3 \) with coordinates \( (X, Y, Z) \). We can define \( e_R, e_u, e_\phi \) as usual. Note that \( e_R \) is here no longer orthogonal to \( e_u \) and \( e_\phi \) in the Euclidean sense as the metric of the velocity space is not Euclidean. The basis vectors are rather mutually orthogonal with respect to the metric \( Z^2 - X^2 - Y^2 \). Calculation of the Jacobian matrix shows that the volume element is given by \( R^2 \sinh u \, du \, d\phi \, dR \). The oriented surface element on the revolution hyperboloid \( R = 1 \) is given by:
$dS = \sinh u \, du \, d\phi \, e_z$.\textsuperscript{13} The surface surrounded by the closed loop defined by $\gamma = \cosh u_0 = \text{thus } 2\pi (\cosh u_0 - 1)$. From $v_y/v_z = \tan \phi$, one can calculate $(1 + \tan^2 \phi) \, d\phi = (v_x v_y - v_y v_x)/v_z^2$, which leads to $d\phi = \gamma^2/(\gamma^2 - 1) \, v \wedge a$. As the Thomas precession along the closed loop $u = u_0$ is given by $(\cosh u_0 - 1) \, \gamma^2/(\gamma^2 - 1) \, 2\pi$, along a part $d\phi$ of this closed loop it is therefore given by $d\theta T = (\cosh u_0 - 1) \, \gamma^2/2 \, v \wedge a$. This is completely analogous to what we did for the small circle on a sphere. The coordinate lines $\phi = \phi_0$ correspond to accelerations $a \parallel v$. The contributions $du$ do therefore not contribute to the Thomas precession. The integration of the Thomas precession over a closed loop of any shape will give the total Berry phase according to the Gauss-Bonnet theorem. No taking into account the contribution $a \parallel v$ is here just analogous to the way we define an integral as the limit of a procedure whereby we use the Simpson rule with ever smaller meshes.\textsuperscript{14}

\subsection*{8.6.5 A critical remark}

After all this a critical question remains. In the fully relativistic solution of the hydrogen problem, the spin-orbit coupling terms do not show up in the calculations. Where are they? And why do we get the correct solutions from the minimal substitution if it is incomplete? The answer is that by solving the problem in cylindrical or spherical coordinates, without introducing covariant derivatives\textsuperscript{15}, we actually introduce the spin-orbit effect (without the correction for Thomas precession) for circular orbits.

As a matter of fact, the transition between Cartesian coordinates and cylindrical coordinates is given by the entirely geometric transformation:

\[ \frac{\partial f}{\partial r} e_r + \frac{1}{r} \frac{\partial f}{\partial \phi} e_\phi + \frac{\partial f}{\partial z} e_z = \frac{\partial f}{\partial x} e_x + \frac{\partial f}{\partial y} e_y + \frac{\partial f}{\partial z} e_z, \]

which shows that $\nabla \equiv \frac{\partial}{\partial r} e_r + \frac{1}{r} \frac{\partial}{\partial \phi} e_\phi + \frac{\partial}{\partial z} e_z$. But when we perform dynamical calculations and calculate temporal derivatives, we must take into account the fact that $(e_z(t), e_\phi(t), e_x(t))$ are functions of time. In the dynamical calculations we must correct for this temporal dependence by introducing covariant derivatives. When we use $(e_z(t), e_\phi(t), e_x(t))$ in dynamical calculations of the position $r(t)$ of a particle, $(e_z(t), e_\phi(t), e_x(t))$ will be a precessing frame. Let us now consider a co-moving copy of this frame at the position $P(r(t))$ of the particle rather than at the origin $O$. By making the calculations in cylindrical coordinates without introducing the covariant derivatives, this co-moving frame will for uniform circular motion carry out exactly the merry-go-round motion in position space. We have seen that this corresponds to the part of the spin-orbit coupling that occurs in Eq. 21. This is fortunate, as the Dirac equation based on the minimal substitution does not correct for the merry-go-round effect. The Dirac equation

\textsuperscript{13} The presence of $e_z$ may look here surprising, but we are dealing with a problem in four-dimensional space with Minkowski metric, where the coordinates of a point are $(\gamma, \gamma v_x/c, \gamma v_y/c, \gamma v_z/c)$. We have dropped the coordinate $\gamma v_z/c = 0$ in our approach in order to be able to represent the hyperboloid with a three-dimensional model. But as the motion takes place in the $Oxy$-plane, the Thomas precession must be around the $z$-axis, which we dropped out of the description. The vector $e_z$ can no longer be written as $e_z \wedge e_x$ because we are dealing with calculations in a four-dimensional space. The vector $e_\phi$ is also orthogonal to $e_z$ and $e_x$ in terms of the Minkowski metric but it is the unit vector of the tetrad that is analogous to $e_t$ in the space-time tetrad. In our three-dimensional subspace $v_z = 0$, we have $e_\phi = e_x \wedge e_z$, where $\wedge$ is the hyperbolic wedge product.

\textsuperscript{14} When this work was finished, A. Gontijo Campos sent me a copy of a paper by N. Alamo and C. Criado, \textit{American Mathematical Monthly}, 116 (2009), pp. 439-446, which takes essentially the same route to calculate the Thomas precession.

\textsuperscript{15} The covariant derivative for a tensor is different from the covariant derivative for a vector. For a scalar we do not need to consider covariant derivatives. For each type of symmetry, their will thus be a different type of covariant derivative. We can thus expect that the covariant derivative for a spinor will be different from that of a vector. The coordinate transformations we use for vectors are not compatible with a coordinate transformation for the spinors, since in a coordinate transformation for a spinor the phase will be unambiguously defined on the whole group. We are thus taking a shortcut to a lot of algebra and conceptual problems in neglecting covariant derivatives.
Lorentz transforms \((\frac{d}{dt}, 0)\) to \((\frac{dr}{dt}, \nabla)\) in \((r, t)\) by using the local instantaneous boost with parameter \(v(r, t)\) and without considering the Thomas precession part of the Lorentz transformation. In these instantaneous boosts, the triads remain just all the time parallel to the \((e_x, e_y, e_z)\), such that any kind of spin-orbit effect remains ignored. By introducing cylindrical coordinates without introducing covariant derivatives, we are thus making up for this shortcoming of the Dirac equation in the case of uniform circular motion, as we are including the merry-go-round effect in the laboratory frame. But as we already pointed out, this fails to account for the Thomas precession and it is thus not completely exact, even for uniform circular motion.

We see thus that the solution of the problem of the hydrogen atom is based on a completely different approach than the one that tries to calculate the various contributions to the spin-orbit coupling explicitly, as attempted in our approach. By not introducing the covariant derivatives we take a short cut to a part of the calculation. We are making an error by using the minimal substitution in the Dirac equation, with the result that we fail to take into account the spin-orbit effect (without the correction for the Thomas precession). But by making a second error when we forget to use covariant derivatives we finally get a result that includes all effects, except the Thomas precession. We may finally note that aligning the reference frame with \(v\) we forget to use covariant derivatives we finally get a result that includes all effects, except the Thomas precession. By not introducing the covariant derivatives we take a short cut to a part of the calculation. We are making an error by using the minimal substitution in the Dirac equation, with the result that we fail to take into account the spin-orbit effect (without the correction for the Thomas precession). But by making a second error when we forget to use covariant derivatives we finally get a result that includes all effects, except the Thomas precession. We would like to solve this equation for \(\omega_T\). To do so, we must first eliminate all reference to \(v\) and \(v\) from the equation. As explained in Footnote 10, \(v\) cannot be calculated from a conservation law between kinetic and potential energy like in classical mechanics, because the electron might have emitted radiation. However, quantized emission of radiation can be taken into account in the value of \(L\), if we admit that the emission of radiation conserves total angular momentum. One can thus use \(v \times qE = \frac{1}{m} \frac{d}{dt} \gamma m c^2 L\), where \(mc^2 = \gamma m_0 c^2 + qV \pm \frac{h\omega_T}{2}\). When \(r \perp v\) we can also use \(L = r \times m v\) to put \(v \times qE = Lc^2 / (r(\gamma m_0 c^2 + qV \pm \frac{h\omega_T}{2}))\) and solve this as an equation of \(v\) in terms of \(\omega_T\) and \(r\). This expression for \(v\) must then consistently be used to replace all occurrences of \(v\) in the equation that results from Eq. 54 after the replacement \(v \times qE = \frac{1}{m} \frac{d}{dt} \gamma m c^2 L\). This way we obtain a complicated equation for \(\omega_T\), with \(r\) as a parameter. If the degree of this equation is larger than 4 it will only be possible to solve it by numerical methods.

It is not obvious that it would be a correct procedure to add this term to the Dirac equation, because after squaring it will lead to new terms. The other contributions to the spin-orbit coupling only enter the scene after squaring. This raises the question if \(m = \gamma m_0 + (qV \pm \frac{h\omega_T}{2}) / c^2\) is actually the correct ansatz. Should we perhaps not just use \(E = mc^2\), where \(E\) is the total energy, to calculate the mass? At first sight, it seems that the equation \(m = \gamma m_0 + (qV \pm \frac{h\omega_T}{2}) / c^2\) seems to account for all corrections on the rest mass in the co-moving frame after a subsequent transformation to the laboratory frame, provided we introduce spherical or cylindrical coordinates in the solution of the Dirac equation obtained by a minimal substitution. When we solve the equation rather in Cartesian coordinates, then we must use the generalized substitution in order to get the spin-orbit coupling correct.

\[\omega_T = \gamma^2 \frac{v \times qE}{\gamma m_0 c^2 + qV \pm \frac{h\omega_T}{2}}\]  

\[(54)\]

We would like to solve this equation for \(\omega_T\). To do so, we must first eliminate all reference to \(v\) and \(v\) from the equation. As explained in Footnote 10, \(v\) cannot be calculated from a conservation law between kinetic and potential energy like in classical mechanics, because the electron might have emitted radiation. However, quantized emission of radiation can be taken into account in the value of \(L\), if we admit that the emission of radiation conserves total angular momentum. One can thus use \(v \times qE = \frac{1}{m} \frac{d}{dt} \gamma m c^2 L\), where \(mc^2 = \gamma m_0 c^2 + qV \pm \frac{h\omega_T}{2}\). When \(r \perp v\) we can also use \(L = r \times m v\) to put \(v \times qE = Lc^2 / (r(\gamma m_0 c^2 + qV \pm \frac{h\omega_T}{2}))\) and solve this as an equation of \(v\) in terms of \(\omega_T\) and \(r\). This expression for \(v\) must then consistently be used to replace all occurrences of \(v\) in the equation that results from Eq. 54 after the replacement \(v \times qE = \frac{1}{m} \frac{d}{dt} \gamma m c^2 L\). This way we obtain a complicated equation for \(\omega_T\), with \(r\) as a parameter. If the degree of this equation is larger than 4 it will only be possible to solve it by numerical methods.

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with time, because the quantities \( v(r,t) \) and \( \gamma(r,t) \) will in general still depend on space and time. There will in any case also be a term with \( \mathbf{a} \cdot \mathbf{\sigma} \) due to the combination \( \left[ \frac{\mathbf{a}}{m} \mathbf{1} \right] \left[ \mathbf{v} \cdot \mathbf{\sigma} \right] \) in the SL(2,\( \mathbb{C} \)) matrices. We have therefore just discussed the anomalous Zeeman effect and the spin-orbit coupling in the non-relativistic limit. Some further aspects of Eq. 55 will be discussed in the Appendix. We will give there e.g. an extremely simple derivation of the term \(-\mu \cdot \mathbf{B}\) that occurs in the literature. This derivation is exact, in contrast with wishy-washy derivations based on a treatment of a current loop.

9 Conclusion

The traditional interpretation of the anomalous \( g \)-factor in the Dirac theory might be physically attractive as it corresponds to our macroscopic intuition, but it does not agree with the true meaning of the algebra and it violates the Lorentz symmetry. We have shown this by developing an alternative approach to the physics of the anomalous \( g \)-factor by just respecting the correct geometrical interpretation of the algebra, a feat that the traditional approach is not able to accomplish as illustrated by the symmetry violation mentioned. We have proposed an interpretation that respects the Lorentz symmetry. We have thereby stuck to the working philosophy that the algebra should remain strictly the same such that only the geometrical interpretation of the algebra can be changed (in such a way that it agrees with the given geometrical meaning of the algebra) and agreement with experiment is automatically preserved.

However, this working philosophy fails when we try to apply the same methods to the algebra of the spin-orbit coupling. In searching an explanation for this failure we discover that it is not our approach but the traditional approach that contains a number of flaws. A similar analysis of Dirac’s theory of the magnetic monopole raises also some troubling issues. The origin of these problems can be traced back to a craze for abstraction whereby any feeling for the original geometrical meaning of the algebra used in the group representation theory is lost.

The whole study presented in this paper just ensues from the natural wish to make sense of Eq. 16/Eq. 21 which was obtained from a few lines of algebra based on group representation theory. This group representation theory provides also all the necessary tools to solve the many problems encountered along this search for better insight. In conclusion we think that this work shows what a powerful tool group theory can be in the quest of trying to make sense of quantum mechanics.

Acknowledgements. I wish to thank Prof. Dr. J.-E. Wegrowe for fruitful discussions.
10 Appendix. Additional Calculations

10.1 Criticism of the calculation of the energy \(-\mu \cdot B\) of a magnetic moment \(\mu\) based on a current loop

Many textbooks propose that the potential energy of a current loop within a magnetic field would be given by \(U_{\text{dipole}} = -\mu \cdot B\). This is really problematic for several reasons:

1. The force of a magnetic field \(B\) on a moving charge \(q\) with velocity \(v\) is \(F = q(v \times B)\). As \(F \cdot d\ell = F \cdot v dt = 0\), a magnetic field cannot do work on a moving charge. It is therefore puzzling how we could define a potential energy \(U_{\text{dipole}}\) with respect to a magnetic field.

2. Some textbooks argue that the magnetic field exerts a torque on the loop and on the dipole moment \(\mu\) of the loop. But the torque is calculated in a special pair of points of the loop without discussing what happens in all the other pairs of points. The special points correspond to the maximal value of the torque. By analyzing what happens in other pairs of points, one can find a pair of points where the torque is zero. This is the minimal possible value for the torque. For all other intermediate values of the torque, one can find a corresponding pair of points. That one has to select the pair of points with the maximal torque to obtain the correct value for the "potential energy" of the current loop with dipole moment \(\mu\) makes the calculation look hand-waving. Moreover, a torque is not a force such that it is not clear what its relation with a potential energy ought to be.

3. In both cases where one applies these arguments to a single electron in circular orbit around the nucleus of an atom, one has to assume that the charge \(q\) of the electron is smeared out over the whole loop. The very definition of \(\mu\) used in all the atomic calculations is built on this idea, which is certainly not correct. Such a single electron is not a current loop and not a true dipole. Moreover, only one force is acting on it, not a torque. One needs thus at least two electrons to define a dipole moment and a torque as for a macroscopic current loop. The correct definitions must thus be based on the analysis of a single electron. After that, one can put several electrons on the same orbit as one can define for a single electron to obtain the macroscopic quantities in terms of dipole moments and torques.

10.2 Magnetic moment of a current carried by a single electron

We will try to remove here these weird conceptual problems by a better description of the problem. First we address the recurring issue that the charge \(q\) of the electron is smeared out over the whole loop. In an atom, the charge density will not be uniform but singular \(\rho(\ell) = q\delta(\ell - \ell_0)\), where \(\ell_0\) is the position of the charge, and \(\ell\) denoted the curvilinear length along the orbit. This leads to a singular current loop, which we can use to describe the real situation of the moving charge. The singular loop will have a singular magnetic moment. Relativistically, we will have \(I = \gamma I_0\).

\[
\mu = \oint \gamma I_0 \frac{1}{2} r \times dr = \oint I_0 \frac{1}{2} r \times \gamma v dt = \oint \frac{1}{m_0} \frac{1}{2} r \times p dq = \frac{q}{2m_0} r_0 \times p_0 = \frac{q}{2m_0} L_0. \tag{56}
\]

We have used here \(I_0 dt = dq\). We can consider \(r_0\) as the centre of the circular orbit that defines the loop that can be associated with it. In an atom, it would be the position vector of the electron with respect to the nucleus. While it is obvious that \(I\) is singular, and that also the charge distribution \(dq = \rho(\ell) dt = q\delta(\ell - \ell_0) dt\) is singular; it is obvious that \(\oint dq = \oint q\delta(\ell - \ell_0) dt = q\). We obtain then \(\mu = -\frac{q}{2m_0} L_0\). We can replace \(L_0\) by \(L\) as \(L\) is a constant of motion. We may note that it is also possible to define \(\mu\) classically such that it does not account for \(\gamma\) in \(L\). One must then write \(\gamma\mu\) instead of \(\mu\). The quantity \(\mu\) is not a dipole moment, because there is only one moving charge. It is a monopole moment.

This calculation of \(\mu\) is exact, while from most presentations one gets the impression that it would a back-of-the-envelope calculation that is only a rough estimation. It may be noted that we do not need the loop. All we have to do is to integrate over a small segment \(dr\). In fact, in the reasoning followed above, the singular magnetic moment was only defined for a line segment \(dr\) and we could choose the loop at will. The rest of the loop that one may add can be chosen arbitrarily as it will give a zero contribution to the contour integral. And this is true at any moment. But taking the orbit for the contour integral has the advantage that we never have to change the arbitrary choice and that the definition will be valid over the whole orbit. It will then be a definition that suits the description of a stationary situation. The further manipulations introduce \(L\) which is a constant. And in the end we integrate \(dq\). In conclusion, with the necessary provisos, the current density of a moving single charged particle is singular and can always be considered as giving rise to a magnetic charge \(\mu\) which is a magnetic monopole moment (without hyphen).

As explained by Griffiths and Hnizdo [19], in Gilbert’s description a magnetic dipole moment consists just of two monopole moments. When we consider the current loop from the viewpoint of a moving Lorentz frame, the symmetry between the two monopole moments becomes broken, which leads to a “hidden momentum”. The conclusions reached in our approach are completely in line with these ideas and show that a single electron in circular motion corresponds to a magnetic monopole moment.

We still have to address the issue (1). The solution of this conundrum is that \(-\mu \cdot B\) does not represent the potential energy, but the kinetic energy as already pointed out in the main text. The following subsection will show how we
get it into the formalism by applying the correct substitution for the coupling of the charge to the free-space Dirac equation.

10.3 Further discussion of Eq. 55 and the origin of the term $-\mu \cdot B$

We consider the terms in Eq. 55 with the opposite sign, such that the signs will be those that will appear in the equation after making the substitution $E \to E' = qV, p \to p - qA$. The term $-\gamma qA \cdot \sigma$ has already been discussed in terms of a velocity field with a vorticity. As it is a vector, it can not contribute to the potential energy. However the vorticity of this term can affect the energy. The term $-\gamma q(V \cdot A)I$ can be rewritten as $-j \cdot A$, such that it can be understood as the "potential energy" of a current, while in reality it describes its kinetic energy, as verified on a simple example in the main text. By introducing $A = -\frac{1}{2}v \wedge B$, the term $-\gamma q(V \cdot A)I$ can also be written as $-L \frac{\omega v}{2m_0}$, where $L = r \wedge \gamma m_0 v$. This angular momentum $L$ is defined with respect to an arbitrary origin, as $r$ is defined with respect to an arbitrary origin. The term $-L \frac{\omega v}{2m_0}$ has the dimension of an energy. We can write it in terms of the "potential energy" term considered above:

$$-U_{dipole} = -\frac{q}{2m_0} L B = \gamma \mu \cdot B = -L \cdot \omega v.$$  \quad (57)

The term $\gamma$ associated with $\mu \cdot B$ is motivated by the fact that $\mu$ has been defined with $L = m_0 r \wedge v$. The minimal substitution will lead to $E \to E' = U_{dipole} = E - \gamma qB$. This quantity $U_{dipole}$ is determined up to an arbitrary constant just like a Coulomb potential is determined up to an arbitrary constant. The arbitrary constant is related to the fact that we can define $r$ in $A = -\frac{1}{2}v \wedge B$ with respect to an arbitrarily chosen origin. The constant can be fixed by choosing an origin. When we choose an origin to calculate the Coulomb potential, this will simultaneously determine an origin for the potential $-\mu \cdot B$. If there is no Coulomb potential, then we must find a different criterion to define the origin. The term $\frac{q}{2m_0}L$ can be rewritten as $\frac{1}{2m_0}r \wedge \omega v$. Here $\frac{1}{2m_0}r$ contains $r$ as a position vector with respect to the arbitrary origin, such that $\mu$ is the monopole moment of the magnetic charge with respect to this origin. Magnetic moments can be defined with respect to arbitrary points, just like angular momentum is in principle defined with respect to an arbitrary origin. This magnetic moment is just the product of the magnetic charge $qV$ with a position vector $r$. Just as only differences of potential energy make physical sense, only differences of magnetic moments make physical sense. These differences will occur from the moment on we have two magnetic charges. It is thus when we have two moving charges in points $r_1$ and $r_2$ that the arbitrariness disappears because $r_2 - r_1$ does then no longer depend on the choice of the origin, such that we then can really define a magnetic dipole moment. It is because it consists of many magnetic moments that a macroscopic current loop can be identified with a magnetic dipole moment. But the singular magnetic moment of a single moving charge is not a magnetic dipole moment. One could call it an arbitrary magnetic monopole moment (which is not the same as a magnetic-monopole moment). One can split the magnetic dipole moment into two such magnetic moments, just as we can split a dipole into two monopoles.

Finally, $-\gamma q(V \cdot A) \cdot \sigma = \frac{1}{2}[(v \cdot A) B - (B \cdot A)v] \cdot \sigma$. This term is also determined up to an arbitrary constant. While it has here the dimension of an energy, it’s pseudo-vector character precludes using it as a potential energy. For a circular motion of the charge due to the Lorentz force exerted by the magnetic field, this term reduces zero if we choose the origin at the centre of the circle. In fact, $r, v, B$ are then all mutually orthogonal. Once again, the case $v \parallel A$ can only have meaning in a context of forced motion. In the non-relativistic limit, where we can neglect $\gamma$, the force term $[\nabla \sigma] [\gamma q(V \cdot A) \cdot \sigma]$ will lead to four terms: two pseudo-scalar terms and two vector terms. One of the vector terms will be the Lorentz force $q(V \cdot (\nabla \wedge A)) \cdot \sigma$. This will lead to the torque on a non-aligned current loop considered by many authors, from the moment on we consider two or more charges rather than one in the current loop. The other vector term will be $q((\nabla \wedge v) \wedge A) \cdot \sigma$. The two pseudo-scalar terms can be written together as $-iq \nabla \cdot (V \wedge A) I = iq (v \cdot B) I - iq A \cdot (\nabla \wedge v) I$

We can analyze this non-aligned situation again in terms of two mutually orthogonal fields $B_1$ and $B_2$. The motion of a single charged particle in the field $B$ will then just be a circular motion in the field $B = B_1 + B_2$. It will take place in the plane orthogonal to $B$ with velocity $v$. We can decompose $A = A_1 + A_2$. The global term $-\gamma q(V \cdot A) \cdot \sigma$ will be zero, as $v \parallel A$. However, the two terms $-\gamma q\gamma q(v \wedge A) \cdot \sigma$ will both be different from zero, with their sum adding up to zero. The terms $\gamma q(V \wedge A_1)$ and $\gamma v \cdot A_1$ have norms $\gamma v A_1 \sin \chi$ and $\gamma v A_1 \cos \chi$, where $\chi$ is the angle between $v$ and $A_1$. The term $\gamma q(V \wedge A_1)$ accounts for that part of the energy that can be associated with $v$, other than the kinetic energy $\gamma v \cdot A_1$ which has to be attributed to the motion in the field $B_1$.

To conclude, we can state that in the absence of an electric field we obtain in the non-relativistic limit for the substitution required:

$$E \to E - \mu \cdot B, \quad cp \to cp - cqA.$$  \quad (58)

This calculation is exact and the picture is clear. We do not understand why the electron spin must be aligned (such that the energies are quantized and the imaginary terms become zero), because we do not understand the electron
spin. The real vector term $cqA$ in $cp \rightarrow cp - cqA$ corresponds in principle to the minimal substitution for a spin-less particle but its vorticity yields the frequency of a spinning motion that after multiplication by $\hbar/2$ can represent energy pumped into the spin. Finally, we can see from all this that in the non-relativistic limit the Zeeman effect is given by:

$$(L + 1 + 2S)qB/2m_0$$

where $L \geq 0$.

References

8. In the experiments of Couder et al. (Y. Couder, S. Protière, A. Fort, and A. Boudaoud, Nature 437(7056), 208-208 (2005)), one observes an association of a particle with a wave, which gives a more intuitive reading for the possibility of having particles and waves at the same time. This is reminiscent of de Broglie’s theory of the pilot wave, but it is still different. Following these ideas, an electron is not both a particle and a wave. The physical situation is just that of a particle associated with a wave. In [2] we describe a natural association of this type without having to postulate that the wave would guide the particle.