

Revised α^4 Term of Lepton $g - 2$ from the Feynman Diagrams Containing an Internal Light-By-Light Scattering Subdiagram

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The α^4 contribution to the lepton $g - 2$ from a gauge-invariant set of 18 Feynman diagrams containing a light-by-light scattering subdiagram internally has been reevaluated by a method independent of the previous approach. Comparison of two methods revealed a program error in the first version. Correcting this error, the contributions of these 18 diagrams become $-0.990\,72(10)(\alpha/\pi)^4$ and $-4.432\,43(58)(\alpha/\pi)^4$ for the electron and muon $g - 2$, respectively. The correction is not large enough to affect the comparison between theory and experiment for the muon $g - 2$, but it does alter the inferred value for the fine structure constant α^{-1} by 6 ppb.

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Precise theoretical and experimental values of lepton anomalous magnetic moments [$a_l = (g_l - 2)/2$] provide one of the most stringent tests of QED [1]. In case of the electron (positron), the experimental value did reach the precision of 4.3 ppb [2]. Currently, the theoretical uncertainty is dominated by that of the fine structure constant α . The most precise α available now is from the atom interferometry experiment [3], which has 7.4 ppb precision. Since the muon anomalous magnetic moment a_μ is sensitive to short-distance physics, high precision measurement (1.3 ppm) of a_μ at Brookhaven National Laboratory may be able to open the first window to “new physics” [4]. Before taking the discovery of new physics in the muon $g - 2$ seriously, however, we must make sure that the old physics, namely, the standard model, is known with sufficiently high precision.

The largest source of theoretical uncertainty (0.7 ppm) for a_μ is the hadronic contribution [5,6]. Unfortunately, we are currently unable to deal with the hadronic correction from first principles because of the nonperturbative nature of QCD. On the other hand, the QED correction can be treated precisely by perturbation theory. In order to achieve the precision comparable to that of measurements, the QED calculation for lepton $g - 2$ must include terms of up to eighth order of perturbation theory. Leading contributions of tenth order are also relevant for a_μ [7,8].

The purpose of this Letter is to correct a program error in the previous calculation of the gauge-invariant set of 18 Feynman diagrams contributing to the α^4 QED term [7]. This was accomplished by constructing alternative forms of integrals for these diagrams. As a consequence, all 891 Feynman diagrams contributing to the eighth-order term of a_e , and additional diagrams contributing to $a_\mu - a_e$, have now been verified by independent calculations and/or checked by analytic comparison with

lower-order integrals. This enables us to pursue with confidence an order of magnitude improvement in numerical precision of all α^4 terms of a_μ and a_e . The results will be reported shortly elsewhere [9,10].

Let us now describe how the error was discovered and corrected. The contribution of the QED diagrams to a_μ can be written in the general form

$$a_\mu(\text{QED}) = A_1 + A_2(m_\mu/m_e) + A_2(m_\mu/m_\tau) + A_3(m_\mu/m_e, m_\mu/m_\tau), \quad (1)$$

where m_e , m_μ , and m_τ are the masses of the electron, muon, and tau, respectively. A similar equation holds for a_e . Throughout this Letter, we use the values $m_e = 0.510\,998\,902(21)$ MeV/ c^2 , $m_\mu = 105.658\,357(5)$ MeV/ c^2 , and $m_\tau = 1776.99(+29, -26)$ MeV/ c^2 [11].

The renormalizability of QED guarantees that the functions A_1 , A_2 , and A_3 can be expanded in power series in α/π with finite calculable coefficients:

$$A_i = A_i^{(2)}(\frac{\alpha}{\pi}) + A_i^{(4)}(\frac{\alpha}{\pi})^2 + A_i^{(6)}(\frac{\alpha}{\pi})^3 + \dots, \quad i = 1, 2, 3. \quad (2)$$

$A_1^{(2)}$, $A_1^{(4)}$, and $A_1^{(6)}$ are known analytically [12]. Most terms contributing to $A_1^{(8)}$ have not yet been obtained by analytic means. The current uncertainty in the value of $A_1^{(8)}$ is a consequence of the fact that, at present, it must be obtained by numerical integration. Its precision is being improved by an extensive computer calculation right now [9]. For the purpose of evaluating $a_\mu(\text{QED})$, however, it is sufficient to use $A_1^{(8)}$ derived from the measured value of the electron anomaly a_e [13] corrected for small contributions of muon, hadron, and weak interactions. The terms $A_2^{(4)}(m_\mu/m_e)$ and $A_2^{(6)}(m_\mu/m_e)$ are known exactly [14]. The situation is quite different for $A_2^{(8)}(m_\mu/m_e)$ since most terms contributing to it are known only by numerical means.

There are altogether 469 Feynman diagrams contributing to $A_2^{(8)}(m_\mu/m_e)$. Of these diagrams, 343 have been checked by more than one independent method, some of which even being analytic. The other 108 diagrams, all of which contain an *external* light-by-light scattering subdiagram, have been checked analytically by comparison with the exactly known sixth-order vertices [10]. Unfortunately, this was not the case for the remaining 18, all generated by inserting a light-by-light scattering subdiagram *internally* in a fourth-order vertex diagram (see Fig. 1). Here *external* and *internal* mean whether one of the attached photon lines represents an external magnetic field or not.

These 18 diagrams form a gauge-invariant set and share the same basic algebraic structure. Unfortunately, it was not possible to examine their analytic structure by comparison with lower-order diagrams since they are not reducible to such diagrams in the UV and/or IR limits. Besides having been checked by two people working independently [15], the only check made was mutual consistency among these 18 diagrams. This is not sufficient to eliminate the possibility that they share the same program error. Clearly, in order to enhance the credibility of the QED calculation of a_l , it is highly desirable to reevaluate these 18 diagrams by more than one method.

This Letter reports the consequence of our effort to reexamine the previous result by construction of new and independent integrals. Before describing the new approach, let us quickly go over the initial approach. In that approach (which will be called version A) [15], we put together vertex diagrams, for instance, $llj(1)$, $llj(2)$, $llj(3)$, all obtained from the self-energy-like diagram llj of Fig. 2 as terms linear in the magnetic field in the weak field expansion. This is to take advantage of their shared structure and the tendency of partial cancellation among them. With the help of Ward-Takahashi identity, the sum $\Lambda^\nu(p, q)$ of all vertex diagrams thus related to a self-

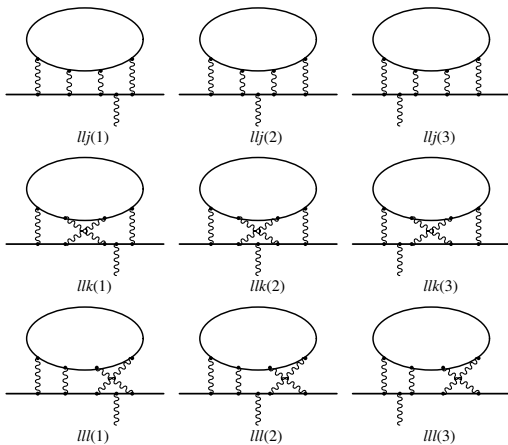


FIG. 1. Vertex diagrams containing a light-by-light scattering subdiagram internally. There are altogether 18 such diagrams.

energy diagram $\Sigma(p)$ can be expressed as

$$\Lambda^\nu(p, q) \approx -q^\mu \left[\frac{\partial \Lambda_\mu(p, q)}{\partial q_\nu} \right]_{q=0} - \frac{\partial \Sigma(p)}{\partial p_\nu}, \quad (3)$$

where $(p + q/2)^2 = (p - q/2)^2 = m_l^2$. The $g - 2$ term is projected out from the right-hand side of (3). In terms of Feynman parameters z_1, z_2, \dots, z_N , the n th order magnetic moment derived from (3) has a form

$$M^{(2n)} = \left(\frac{-1}{4} \right)^n (n-1)! \int (dz) \left[\frac{\mathbf{E} + \mathbf{C}}{n-1} \frac{1}{U^2 V^{n-1}} + (N + \mathbf{Z}) \frac{1}{U^2 V^n} \right], \quad (4)$$

where bold letters \mathbf{E} , \mathbf{C} , \mathbf{N} , and \mathbf{Z} stand for parts of the projection operator adapted to the first and second terms on the right-hand side of (3). U is the Jacobian of transformation from momentum-space variables to Feynman parameters. V^{-1} is obtained by combining all propagators into one with the help of Feynman parameters. See [16] for precise definitions of projection operators, U , V , and (dz) .

In order to check the validity of version A based on Fig. 2, we evaluated these diagrams by a second method, called version B, which is actually a straightforward parametrization of individual vertex diagrams of Fig. 1, without relying on the Ward-Takahashi identity. In this approach, it is convenient to evaluate the sum $ijkl(n) \equiv llj(n) + llk(n) + ll(1)$, where $n = 1, 2, 3$, since partial cancellation of singular terms occurs resulting in less singular behavior.

Diagrams of Fig. 1 (or Fig. 2) form a (formal) gauge-invariant set. But individual diagrams are UV divergent and must be regularized in advance, for instance, by dimensional regularization, to enforce gauge invariance. For numerical evaluation, however, it turns out to be more convenient to combine it with a subtractive regularization. Let $F_m(d)$ be one of the integrals defined in d dimensions, where m is any one of $llj(1), \dots, ll(3)$. Let $G_m(d)$ be the subtraction term containing the light-by-light scattering tensor with zero external momenta $\Pi_{\mu\nu\sigma\rho}(0, 0, 0, 0)$ as well as terms containing a sixth-order charge renormalization diagram. Let us rewrite $F_m(d)$ symbolically as

$$[F_m(d) - G_m(d)] + G_m(d), \quad (5)$$

where “symbolically” means that subtraction is performed on the integrand before the integration is carried

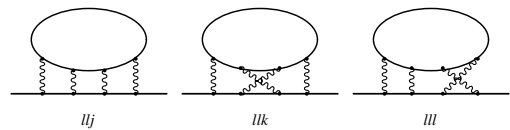


FIG. 2. Self-energy-like diagrams in which lepton lines propagate in the magnetic field. First-order terms in the weak magnetic field expansion correspond to the diagrams of Fig. 1.

out. Now we can safely take the limit $d \rightarrow 4$ for the first term since its integrand does not cause UV divergence. Of course, the second term $G_m(d)$ is singular for $d \rightarrow 4$. However, gauge invariance guarantees that the sum of $G_m(d)$ over all diagrams of Fig. 1 vanishes for any value of dimension d :

$$\sum_m G_m(d) = 0. \quad (6)$$

Thus, we have to compute only $[F_m(4) - G_m(4)]$ in the end.

The integrands of both versions *A* and *B* were generated by an algebraic program FORM [17]. Numerical integration is carried out by an adaptive-iterative Monte Carlo routine VEGAS [18].

When we compared the numerical results of versions *A* and *B*, we were surprised to find that their values were significantly different. After extensive detective work, we located a programming error in version *A*, which resulted from an incomplete implementation of the *E* operation of (4) in the algebraic manipulation program: It left out some terms referring to the light-by-light loop subdiagram. Such a program error can be readily detected if the integral exhibits UV or IR divergence after renormalization is carried out. Unfortunately, the particular error in the 18 diagrams caused no divergence and escaped scrutiny of two people. Once this error was corrected, both approaches gave identical numerical results. Thus, we now have two sets of independent codes for the 18 diagrams that have been fully verified.

Numerical evaluation of these diagrams requires an enormous amount of computational effort. A systematic algorithm of computation to minimize human error is indispensable. Such a scheme was developed originally for the calculation of the sixth-order lepton $g - 2$ [16,19] and was thoroughly tested over the years [20]. It was later extended to the eighth order [21]. Early results for the eighth-order term remained rather crude for many years. This is mainly due to the enormous size of the integrands which could not be handled adequately by the computers then available. More precise values have become available only in this decade thanks to the development of massively parallel computer, which enabled us to vastly increase the sampling statistics of VEGAS.

Enlarging sampling statistics, however, amplified the difficulty caused by a previously poorly understood problem in estimating errors in a computer calculation. This arises from the fact that computer calculation always deals with a finite number of digits. This means that the error estimate based on the assumption of normal distribution of errors must be modified to take the effect of rounding-off of digits into account. In our formulation in which subtractive renormalization of UV divergence as well as separation of IR divergences are carried out on the computer, this *digit deficiency problem* can seriously distort error estimates, or even prevent further iteration.

TABLE I. Muon $g - 2$ contributions from the diagrams of Fig. 1. In version *A* the Ward-Takahashi-summed $llj \equiv llj(1) + llj(2) + llj(3)$, $llk \equiv llk(1) + llk(2) + llk(3)$, and $lll \equiv lll(1) + lll(2) + lll(3)$ are calculated, while in version *B* $jkl(1,3) \equiv llj(1) + llj(3) + llk(1) + llk(3) + lll(1) + lll(3)$ and $jkl(2) \equiv llj(2) + llk(2) + lll(2)$ are calculated.

	Version A		Version B
llj	6.389 802(460)	$jkl(1,3)$	-3.509 978(802)
llk	-7.763 474(537)	$jkl(2)$	-0.921 589(89)
lll	-3.059 704(452)		
Sum	-4.433 376(840)	Sum	-4.431 567(806)

Besides increasing the number of effective digits from real*8 to real*16 arithmetic, which was the most obvious and effective cure, various methods had to be devised to deal with the *digit deficiency error* [22].

New results of numerical integration of a_μ by versions *A* and *B* are listed in Table I. The values of llj , llk , and lll listed were obtained using 5×10^9 sampling points per iteration and iterated 110, 219, and 220 times, respectively. They were evaluated on *v1* cluster at Cornell Theory Center. The calculation of version *B* was carried out on Fujitsu VPP700E at the Computer and Information Division of RIKEN. For $jkl(1,3)$, 4.6×10^9 sampling points per iteration were used for 131 iterations. The program $jkl(2)$ shows less singular behavior. It was evaluated using 4.6×10^9 sampling points per iteration and iterated 60 times.

Results for the electron are listed in Table II. The values of llj , llk , and lll listed were obtained using 2×10^9 sampling points per iteration and iterated 160, 220, and 180 times, respectively, on *v1* cluster at Cornell Theory Center. For $jkl(1,3)$, 4.6×10^9 sampling points per iteration were used for 60 iterations on VPP700E. For $jkl(2)$, 4.6×10^9 sampling points per iteration were used and iterated 60 times.

Combining the results of version *A* and version *B* from Table I, and similarly for Table II, which we treat as statistically independent, we obtain the best estimate of the contribution to a_μ from the 18 Feynman diagrams of Fig. 1:

TABLE II. Electron $g - 2$ contributions from the diagrams of Fig. 1. In version *A* the Ward-Takahashi-summed $llj \equiv llj(1) + llj(2) + llj(3)$, $llk \equiv llk(1) + llk(2) + llk(3)$, and $lll \equiv lll(1) + lll(2) + lll(3)$ are calculated, while in version *B* $jkl(1,3) \equiv llj(1) + llj(3) + llk(1) + llk(3) + lll(1) + lll(3)$ and $jkl(2) \equiv llj(2) + llk(2) + lll(2)$ are calculated.

	Version A		Version B
llj	2.551 223(78)	$jkl(1,3)$	-0.872 717(138)
llk	-1.873 801(72)	$jkl(2)$	-0.117 959(28)
lll	-1.668 182(80)		
Sum	-0.990 760(133)	Sum	-0.990 675(141)

$$a_{IV(d)}^{(8)\mu} = -4.432\,43(58)(\frac{\alpha}{\pi})^4, \quad (7)$$

and a corresponding result for a_e ,

$$a_{IV(d)}^{(8)e} = -0.990\,72(10)(\frac{\alpha}{\pi})^4. \quad (8)$$

Here, superscripts $(8)\mu$ and $(8)e$ refer to the eight-order muon and electron $g - 2$, the subscript IV to the group of all diagrams containing light-by-light scattering subdiagrams, and (d) to its subgroup shown in Fig. 1, which consists of all diagrams containing an internal light-by-light scattering diagram.

The new results (7) and (8) supersede the earlier values $-3.4387(533)(\alpha/\pi)^4$ [7] and $-0.7503(60)(\alpha/\pi)^4$ [23], respectively. The effect of this modification on $a_{\mu}(\text{QED})$ is less than 1% of the overall eighth-order term [which is of the order of $130(\alpha/\pi)^4$], and thus does not affect comparison of experiment and theory significantly. On the other hand, the effect on a_e is $\sim -7.0 \times 10^{-12}$ which is about 16% of the entire eighth-order term and is larger than the measurement uncertainty 4.3×10^{-12} . As a consequence, it reduces the inverse fine structure constant α^{-1} obtained from theory and measurement of a_e by $\sim 0.82 \times 10^{-6}$ or ~ 6 ppb. Currently, all α^4 terms are being upgraded by an extensive numerical integration. Precise values of $a_{\mu}(\text{QED})$ and $a_e(\text{QED})$ including these terms will be reported in [9,10].

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