Hyperfine Field and Hyperfine Anomalies of Copper Impurities in Iron

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A new value for the hyperfine magnetic field of copper impurities in iron is obtained by combining resonance frequencies from β-NMR/ON experiments on 59Cu, 65Cu and 71Cu with magnetic moment values from collinear laser spectroscopy measurements on these isotopes. The resulting value, i.e. $B_{hf}(\text{CuFe}) = -21.794(10)$ T, is in agreement with the value adopted until now but is an order of magnitude more precise. It is consistent with predictions from $ab$ initio calculations. Comparing the hyperfine field values obtained for the individual isotopes, the hyperfine anomalies in Fe were determined to be $59\Delta_{69} = 0.15(9)$ % and $71\Delta_{69} = 0.07(11)$ %.

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I. INTRODUCTION

Precise values of magnetic hyperfine fields [1] allow the determination of nuclear magnetic moments by experimental methods such as integral perturbed angular correlation (IPAC) [2] and time-differential perturbed angular distribution (TDPAD) [3], or low-temperature nuclear orientation (LTNO) and nuclear magnetic resonance on oriented nuclei (NMR/ON) [4]. Precise results, in addition, allow a detailed comparison with theory.

The magnetic hyperfine fields of substitutional impurities in bcc Fe are at present well understood for most of the elements in the periodic table [5–11], with sizeable differences between theory and experiment remaining mainly for the heavier 5d impurities [12, 13], the alkaline elements [14–17], and the actinides [11]. Still, for a few elements no precise experimental results are available yet. A special case is copper, for which the currently accepted value of the hyperfine field in iron is $B_{hf}(\text{CuFe}) = -21.81(1)$ T [18]. This value resulted from a spin-echo NMR (nuclear magnetic resonance) measurement with the sample containing the stable isotopes 63Cu and 65Cu cooled cooled to a temperature of 4.2 K. The error of 0.1 T was not given in the original publication but was provided later as a private communication by one of the original authors [19]. The sign was obtained from the field shift in NMR measurements [20] and was confirmed by theoretical calculations (Ref. [21] and Sec. VI).

In the past few years the magnetic hyperfine interaction frequencies $\nu_{\text{res}} \sim \mu B_{\text{tot}}$, with $\mu$ the nuclear magnetic moment and $B_{\text{tot}}$ the total magnetic field the nuclei experience, have been determined for the Cu isotopes 59Cu [22], 67Cu [19], 69Cu [23], and 71Cu [24] with the β-NMR/ON method, i.e. NMR/ON with β-particle detection. In these measurements, performed on samples that were cooled to millikelvin temperatures, the above mentioned value for the hyperfine field of Cu impurities in Fe host, viz. $B_{hf}(\text{CuFe}) = -21.81(1)$ T, was used in order to extract the nuclear magnetic moments for the isotopes studied. Recently, the magnetic moments of the Cu isotopes with $A = 61$ to 75, i.e. including the ones mentioned above, have been determined by collinear laser spectroscopy measurements at ISOLDE-CERN [25–27]. As results for the β-NMR/ON resonance frequencies and for the magnetic moments from the laser spectroscopy measurements for these isotopes all have similarly high precisions, ranging from 2 to $6 \times 10^{-4}$, combining these results allows determining a new and precise value for the hyperfine magnetic field of Cu in Fe at 0 K.

Note that the resonance frequencies and magnetic moment values that will be used here were all obtained with the same experimental methods and setups. This reduces the risk for possible systematic errors related to calibration issues.

II. NUCLEAR MAGNETIC RESONANCE ON ORIENTED NUCLEI

The NMR/ON method has been applied widely for the precise determination of the magnetic hyperfine splitting of radioactive nuclei in the ferromagnetic host lattices Fe, Co and Ni. In most cases, the primary goal was to deduce the nuclear magnetic moments of the impurity isotopes [28]. The same technique is also used to ground state spins [29], hyperfine fields [1, 30], nuclear relaxation times [31, 32], and quadrupole splittings [33, 34], as well as to obtain information on the lattice location and implantation behavior of implanted impurities [13, 35–37].
NMR/ON experiments require the nuclei to be oriented, which is done using the LTNO method [4] and requires cooling down the radioactive samples to temperatures in the millikelvin region and subjecting them to high magnetic fields, either hyperfine magnetic fields [4] or externally applied fields [38].

In NMR/ON, the resonant depolarization of the radioactive probe nuclei is detected as a function of the frequency of the applied radio-frequency field via the resulting destruction of anisotropy in the anisotropic emission of the decay radiation. The resonance frequency is related to the hyperfine field, \( B_{hf} \), through the relation

\[
\nu_{\text{res}}[\text{MHz}] = \frac{7.6226 \cdot \mu[\mu_N] B_{\text{tot}}[\text{Tesla}]}{\hbar[\text{MHz}]} \quad (1)
\]

with \( I \) the nuclear spin of the isotope studied and

\[
B_{\text{tot}} = B_{hf} + B_{\text{app}}(1 + K) - B_{\text{dem}}. \quad (2)
\]

Here the hyperfine field \( B_{hf} \) includes the Lorentz field of 0.742 T for bcc iron at 0 K, \( B_{\text{app}} \) is the externally applied magnetic field, \( B_{\text{dem}} \) is the demagnetization field and, \( K \) is the Knight shift. The constant factor is the ratio of the fundamental constants \( \mu_N/\hbar \) and does not contribute to the error budgets here.

### III. COLLINEAR LASER SPECTROSCOPY

Collinear laser spectroscopy experiments determine the hyperfine structure of atomic ground- and excited states, yielding precise values for the hyperfine parameters \( A \) and \( B \), which in turn provide accurate values for magnetic and quadrupole moments of the isotopes studied [41].

Recently, collinear laser spectroscopy experiments were performed on a series of Cu isotopes at ISOLDE with the COLLAPS setup, including the four isotopes for which resonance frequencies are available from recent \( \beta \)-NMR/ON measurements. For several of these isotopes laser spectroscopy could only be performed after the installation of the ISCOOL cooler and buncher radio-frequency quadrupole Paul trap [42, 43]. This allowed the collinear laser setup to be operated in bunched mode, reducing the background photon counts from scattered laser light by more than three orders of magnitude, and permitting to perform measurements on isotopes that were previously not accessible due to their low yields. The \( \text{Cu}^+ \) bunches from ISCOOL were sent through a sodium vapor cell which neutralized the ions through charge-exchange collisions. A voltage was applied to the vapor cell for tuning the velocity of the ions and bringing them onto resonance with the laser beam that was overlapped with the Cu beam in the co-propagating direction. Resonances were located by measuring the photon yield as a function of the voltage with two photomultiplier tubes, the voltage of which was gated so that photons were only recorded when an atom bunch was within the light collection region [25]. An example of a collinear resonance fluorescence spectrum obtained in these measurements is shown in Fig. 1 of Ref. [25].

### IV. HYPERFINE MAGNETIC FIELD FOR COPPER IN IRON

In the following the magnetic moments from the collinear laser spectroscopy measurements on the isotopes \( ^{59}\text{Cu} \) [27], \( ^{67}\text{Cu} \) [26], \( ^{69}\text{Cu} \) [26] and \( ^{71}\text{Cu} \) [25] are combined with the \( \beta \)-NMR/ON measurements that were performed on these isotopes [19, 22–24]. The experimental magnetic moment values and \( \beta \)-NMR/ON resonance frequencies for \( ^{59}\text{Cu} \), \( ^{69}\text{Cu} \), and \( ^{71}\text{Cu} \) are listed in Table I and will be discussed in Sec. IV B. The data for \( ^{67}\text{Cu} \) are discussed separately in Sec. IV C.

As the extraction of the hyperfine magnetic field using Eqs. 1 and 2 requires that a possible Knight shift and the demagnetization field are taken into account as well, the \( \beta \)-NMR/ON measurements and these two factors will be addressed in some detail first.

#### A. \( \beta \)-NMR/ON measurements

All four \( \text{Cu} \) isotopes on which \( \beta \)-NMR/ON measurements have been performed were produced at the ISOLDE isotope separator facility. The NMR/ON measurements were performed either on-line using the NICOLE LTNO setup [44] at ISOLDE (for \( ^{59}\text{Cu} \) [22], \( ^{69}\text{Cu} \) [23] and \( ^{71}\text{Cu} \) [24], or off-line with the nuclear-orientation facility at Oxford University (for \( ^{67}\text{Cu} \) [19]). As an example, the NMR/ON result for \( ^{59}\text{CuFe} \) is shown in Fig. 1.

##### 1. Demagnetization field

The calculation of the demagnetization field is not straightforward, but for the simple shapes and thin foils (magnetized in the plane of the foil) that were used in the \( \beta \)-NMR/ON measurements discussed here, analytical expressions can be obtained [40]. At the current level of precision, the small demagnetization fields in the foils used, which are of the order of 0.01 T, cannot be neglected.

The foil used by the Leuven group for the measurement with \( ^{59}\text{Cu} \) had a size of 9 mm by 14 mm and an initial thickness of 125 \( \mu \text{m} \). It was polished with diamond-base paste with grain sizes of 3 \( \mu \text{m} \) and 1 \( \mu \text{m} \). It is estimated that this reduced the thickness of this foil to (95 \( \pm \) 20) \( \mu \text{m} \). For this a demagnetization field \( B_{\text{dem}} = 0.020(4) \text{T} \) is calculated. The foils used by the Oxford group for the measurements with \( ^{67}\text{Cu} \), \( ^{69}\text{Cu} \) and \( ^{71}\text{Cu} \) had an initial thickness of 250 \( \mu \text{m} \) [45]. They were first polished with sand paper with CAMI (Coated Abrasive Manufacturers Institute) grit designation 600 (i.e.
165 is the ratio of the count rates for two detectors at 15° and 165° with respect to the quantization axis as a function of frequency. At the bottom, the count rate at 0° is shown for the 136 keV frequency. (arrows indicate the vertical scale that applies; from ref. [22])

FIG. 1. On-line NMR/ON curve for 69Cu isotope. Plotted is the ratio of the count rates for two detectors at 15° and 165° with respect to the quantization axis as a function of frequency. At the bottom, the count rate at 0° is shown for the 136 keV γ ray of the 57CoFe thermometer (corresponding to a sample temperature of about 10 mK) for the same frequency region, showing no effect at the position of the 59Cu resonance. The slope in the count rates is caused by a small increase of the sample temperature with the resonance frequency. (arrows indicate the vertical scale that applies; from ref. [22])

with an average particle diameter of 16 μm) and thereafter also with diamond-base paste. Due to the larger particle diameter used in the first step, this procedure removed more material from the foils leading to an estimated thickness of (190 ± 40) μm. For typical dimensions of 10 mm by 15 mm for such foils, a demagnetization field $B_{dem} = 0.036(8)$ T is then calculated for the measurements with 67Cu, 69Cu and 71Cu.

2. Knight shift

The Knight shift for copper in iron has never been determined at low temperatures. For other elements in ferromagnetic host materials Knight shift values ranging from zero to about 5% have been reported [39]. Thus, for the low external fields, $B_{app}$, of 0.1 T and 0.2 T that were used in the $\beta$-NMR/ON measurements discussed here, the Knight shift corrections could amount up to about 0.005 T and 0.010 T, respectively. These values were then used as the 1σ error bars on the values for the externally applied magnetic fields. Note that the precision to which these external fields could be set is an order of magnitude better and can therefore be neglected at the current level of precision.

B. Results for 59Cu, 69Cu and 71Cu

The magnetic moment values and the resonance frequencies for the isotopes 59Cu, 69Cu and 71Cu are listed in Table I (the case of 67Cu will be discussed separately in the next section). Also listed there are the values for the total magnetic field, $B_{tot}$, obtained for each isotope using Eq. 1, as well as the hyperfine field, $B_{hf}$, that is then obtained from Eq. 2 using the values for the externally applied field, $B_{app}$, and the demagnetization field, $B_{dem}$.

As can be seen, the hyperfine field values obtained for these three isotopes are in very good agreement with each other (see also Fig. 2). Combining all three results yields a weighted average value of

$$B_{hf}(CuFe) = -21.794(10) \text{ T}.$$ (3)

The differences between the hyperfine field values obtained for the three isotopes listed in Table I could be due to small differences in the distribution of the nuclear magnetization over the nuclear volume for the three isotopes, i.e. hyperfine anomalies [46]. The usual definition for the hyperfine anomaly, $\epsilon_i$, for a single nuclear state $i$ is

$$B_{eff} = B_0 (1 + \epsilon_i).$$ (4)

where $B_{eff}$ is the hyperfine field averaged over the distribution of nuclear magnetization of the state $i$ and $B_0$ is the hyperfine field at $r = 0$. The difference of the hyperfine anomalies of two nuclear states in the same host metal is then given by

$$\Delta^2 = \epsilon_1 - \epsilon_2 = \frac{B_i}{B_{eff}} - 1.$$ (5)

When considering the possible presence of hyperfine anomalies the values for $B_{hf}$ listed in Table I have to be interpreted as values for $B_{app}$. It then turns out that the differences of the hyperfine anomalies of these three isotopes are less than $3 \times 10^{-3}$ (see Table I; 90% C.L.). Note that previously the hyperfine anomaly between 64Cu and 66Cu was estimated to be less than $5 \times 10^{-5}$ [47], while in the recent collinear laser spectroscopy measurements on Cu isotopes [26], no indication for hyperfine anomalies was observed either for the series of isotopes ranging from $A = 61$ to 75.

For 59Cu, two other values for the magnetic moment were recently quoted as well, i.e. $\mu(59Cu) = 1.83(4) \mu_N$, from in-source laser spectroscopy [48], and $\mu(59Cu) = 1.910(4) \mu_N$, from an in-gas-cell laser spectroscopy experiment [49, 50]. As the first value is much less precise than the one obtained from collinear laser spectroscopy it is not further used here. The second value differs well outside error bars from the collinear laser spectroscopy result (see Table I). Combining it with the $\beta$-NMR/ON resonance frequency of 208.79(4) MHz for 59Cu reported in Ref. [22], a hyperfine field value
of \(B_{hf}(\text{CuFe}) = -21.59(5) \text{T}\) is obtained. This differs by about 4 standard deviations from the values listed in Table I that resulted from combining resonance frequencies and magnetic moment values that were each obtained with the same experimental setups. This difference might be the due to unforeseen systematic effects in this measurement [49] which was performed with a different experimental setup, as is currently being investigated [51].

C. The case of \(^{67}\text{Cu}\)

The situation for \(^{67}\text{Cu}\) turns out to be more complicated. In Ref. [23], the magnetic moment of \(^{67}\text{Cu}\) obtained with the Oxford LTNO setup is given as \(\mu(\text{Cu}^{67}) = +2.542(2) \mu_N\); in Ref. [52] this result is quoted as \(\mu = +2.536(3) \mu_N\) (this smaller error bar most probably only reflects the statistical precision). The authors do not quote a resonance frequency value, but analysis of the resonance curve shown in Ref. [23] yields a central value of \(\nu_{res} = 278.38(6) \text{MHz}\). In combination with the magnetic moment value of \(\mu = +2.536 \mu_N\), this yields a total magnetic field of \(-21.60 \text{T}\). Comparing this with the hyperfine field value of \(-21.8(1) \text{T}\) for copper in iron that was used in Ref. [23] and neglecting the demagnetization field (as the authors of Ref. [23] also did), yields a value of \(+0.20 \text{T}\) for the externally applied magnetic field instead of the value of \(+0.10 \text{T}\) mentioned in Refs. [23, 52]. The latter value is therefore most probably a typographical error.

When combining the above resonance frequency of \(278.38(6) \text{MHz}\) with the magnetic moment value \(\mu(\text{Cu}^{67}) = +2.5142(6) \mu_N\) from the laser spectroscopy experiments [26], a hyperfine field value of \(B_{hf}(\text{CuFe}) = -21.952(15) \text{T}\) is obtained for \(B_{ext} = 0.20(1) \text{T}\) and with \(B_{dem} = 0.036(8) \text{T}\) for the foils of the Oxford team (see Sec. IV A 1). If the demagnetization field is neglected \(B_{hf}(\text{CuFe}) = -21.988(15) \text{T}\) is obtained. Both values are at variance with the ones obtained for the other Cu isotopes (see Table I and Fig. 2).

Concluding, there seems to be a problem with the \(\beta\)-NMR/ON result for \(^{67}\text{Cu}\) that was obtained using another experimental setup than the one used for the isotopes \(^{59}\text{Cu}, ^{69}\text{Cu}\) and \(^{71}\text{Cu}\). The origin of this may e.g. be an undetected error in the frequency calibration. We therefore did not include the \(\beta\)-NMR/ON result for \(^{67}\text{Cu}\) in the hyperfine field analysis presented here.

V. PREVIOUS RESULTS

Table II lists other values for the hyperfine field of \(\text{Cu}\) in \(\text{Fe}\) that are available in the literature, most of which were obtained in classical NMR experiments at room temperature and with - unfortunately - no error being quoted. As can be seen, most values are in reasonable agreement with the new value of \(-21.794(10) \text{T}\) presented here. The value from the \(\gamma-\gamma\) perturbed angular correlation measurement, which is significantly deviating from all other results, is either wrong or might be related to a different lattice site for \(\text{Cu}\) impurities in \(\text{Fe}\).

VI. COMPARISON WITH THEORETICAL VALUES

Hyperfine fields in solids can be calculated from first principles. This allows understanding trends in those hyperfine fields as a function of the impurity element in e.g. an iron matrix [5–11], and it allows disentangling the hyperfine field into contributions with a different physical origin [10, 11, 61, 62]. We have calculated the hyperfine field of \(\text{Cu}\) in \(\text{Fe}\) within the framework of Density Functional Theory [63–65], using the Perdew-Burke-Ernzerhof (PBE) exchange-correlation functional [66]. For solving the scalar-relativistic Kohn-Sham equations we employed the Augmented Plane Waves + local orbitals (APW+lo) method [65, 67, 68] as implemented in the wien2k package [69] for periodic solids. In this method the wave functions are expanded into spherical harmonics inside nonoverlapping atomic spheres of radius \(R_{MT}\) and in plane waves in the remaining space of the unit cell, i.e. the interstitial region. We took \(R_{Fe}^{\text{MT}} = R_{Cu}^{\text{MT}} = 2.30 \text{a.u.}\) . The plane wave expansion of the wave function in the interstitial region was truncated at a large value of \(K_{max} = 8.0 / R_{MT}^{\text{min}} = 3.48 \text{a.u.}^{-1}\), which leads to very accurate values for the calculated hyperfine fields. A dense mesh of \(\text{k}\)-points, corresponding to a \(20 \times 20 \times 20\) mesh for a conventional bcc unit cell for \(\text{Fe}\), was taken. Spin-orbit coupling was taken into account by a second variational step scheme [70] using a cutoff energy \(E_{SO} = 5.0 \text{Ry}\). The

![Figure 2](image-url)
substitutional Cu impurity was modeled by a 128-atom supercell, and all atoms in the supercell were allowed to adjust their positions due to the presence of the impurity. The lattice constant for the Fe-matrix was taken to be the equilibrium lattice constant for the PBE functional (2.8404 Å). These settings allow for an excellent numerical convergence of the hyperfine field.

The results of the calculations are summarized in Table III. As can be seen, the distance between the Cu

![](http://example.com/image.png)

TABLE III. Calculated values for the Fermi contact field, orbital hyperfine field, dipolar hyperfine field and atomic dipolar hyperfine field for a Cu impurity in Fe. Spin and orbital contributions to the atomic magnetic moment for Cu (μ_B) and distance between Cu and its first Fe-neighbor (Å).

| B(T) | Ref. | method |
|------|------|--------|
| B_F  |      |        |
| B_{orb} |      |        |
| B_{dip} |      |        |
| B_{app}^a |      |        |
| B_{app}^b |      |        |
| B_{tot} |      |        |

impurity and its first eight Fe neighbours is expanded only slightly (0.75 Å) compared to the Fe-Fe distance of 2.460 Å in pure Fe. The dominant contribution to the hyperfine field is the Fermi contact term, caused by s-electron spin polarization, due to the small atomic (d-electron) magnetic moment at the Cu atom. This value can be further split into a core contribution due to 1s and 2s electrons (~8.54 T), a semi-core contribution by the 3s-electrons (+5.80 T) and a valence contribution by 4s electrons (~22.59 T). The result for the Fermi contact hyperfine field can be compared with the ~18.2 T that was obtained 25 years ago by the Korringa-Kohn-Rostoker Green’s function method [6]. The orbital hyperfine field of Cu (0.60 T) is an order of magnitude smaller than the corresponding quantity in pure Fe, consistent with the very small atomic orbital magnetic moment of Cu. All contributions to the hyperfine field, together with the Lorenz field (0.74 T), sum to a total hyperfine field of ~23.99 T. Although this is about 2 T larger than the experimental value determined in this work, this deviation is state-of-the-art, and is due to inherent limitations of the chosen exchange-correlation functional.

VII. CONCLUSION

Combining resonance frequencies for the isotopes ^{59}Cu, ^{69}Cu and ^{71}Cu obtained from β-NMR/ON measurements with the NICOLE LTNO setup at ISOLDE, with magnetic moment values obtained for these isotopes in collinear laser spectroscopy measurements at ISOLDE, the hyperfine field of Cu impurities in iron is found to be $B_{hf}(CuFe) = -21.794(10)$ T. This value is in agreement with but almost an order of magnitude more precise than the previously adopted value of ~21.8(1) T and in good agreement with predictions from ab initio calculations. Interpreting the differences between the hyperfine field values obtained for the individual isotopes to be due to hyperfine anomalies, the hyperfine anomalies in Fe for the isotopes considered here were found to be smaller than $3 \times 10^{-3}$ (90% C.L.).

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TABLE II. Overview of hyperfine magnetic fields for Cu in Fe host reported in the literature. Note that in most cases no error bar was quoted.

| Isotope | μ (μ_B) | Ref. | $\nu_{res}$ (MHz) | Ref. | $B_{tot}$^a (T) | $B_{app}$^b (T) | $B_{atom}$ (T) | $B_{tot}$^a (T) | $\chi^{69}$ (%) |
|---------|---------|------|------------------|------|----------------|----------------|----------------|----------------|----------------|
| ^{59}Cu | 1.89(9) | [27] | 208.79(4) | [22] | -21.72(11) | 0.10(5) | 0.02(4)^c | -21.807(13) | 0.15(9) |
| ^{69}Cu | 2.838(10) | [26] | 311.7(1) | [23] | -21.611(10) | 0.20(1) | 0.036(8)^c | -21.775(16) | - |
| ^{71}Cu | 2.27(8) | [25] | 250.00(14) | [24] | -21.627(14) | 0.20(1) | 0.036(8)^c | -21.791(19) | 0.07(11) |

Weighted average $-21.794(10)$

^aSign from [57]. ^bFrom the Refs. listed in column 3. ^cCalculated with the formulas derived in Ref. [40] (see also Sect. IV A 1).
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