Probing the $N = 32$ Shell Closure below the Magic Proton Number $Z = 20$: Mass Measurements of the Exotic Isotopes $^{52,53}\text{K}$


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The recently confirmed neutron-shell closure at $N = 32$ has been investigated for the first time below the magic proton number $Z = 20$ with mass measurements of the exotic isotopes $^{52,53}\text{K}$, the latter being the shortest-lived nuclide investigated at the online mass spectrometer ISOLTRAP. The resulting two-neutron separation energies reveal a 3 MeV shell gap at $N = 32$, slightly lower than for $^{52}\text{Ca}$, highlighting the doubly magic nature of this nuclide. Skyrme-Hartree-Fock-Bogoliubov and Gorkov-Green function calculations are challenged by the new measurements but reproduce qualitatively the observed shell effect.

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the closure of the $Z = 20$ proton core in calcium, where recently also for $^{54}$Ca a large $E(2^+)$ value has been revealed [14]. Until now, the region below $Z = 20$ was unexplored, as these extremely exotic nuclides could not be produced in sufficient quantity.

In this Letter, we present the first measurement of the masses of $^{52,53}\text{K}$ and, thus, the first investigation of the $N = 32$ shell closure towards the neutron drip line for $Z < 20$. The experimental $S_2n$ values are compared to the predictions of self-consistent \textit{ab initio} Gorkov-Green function (GGF) theory. This recent development significantly extends the reach of calculations based on realistic interactions, including three-nucleon forces, in particular to open-shell (and odd-$Z$) nuclei. In contrast, we show that the predictions of mean-field calculations with phenomenological interactions are highly dependent on the parameterization of both the particle-particle and particle-hole parts of the energy functional. To this end, we performed in addition Hartree-Fock-Bogoliubov (HFB) calculations with the Skyrme interactions SLy4 and SLy5 [15], the latter containing a phenomenological tensor term.

The exotic potassium isotopes were studied at ISOLDE/ CERN with ISOLTRAP’s multi-reflection time-of-flight mass separator (MR-TOF MS) [16,17]. Figure 1 shows the parts of the setup relevant for the present measurements: a radio-frequency quadrupole (RFQ) ion cooler and buncher [18], the MR-TOF MS, and a microchannel plate (MCP) detector.

The nuclides $^{52}\text{K}$ and $^{53}\text{K}$ were produced by impact of 1.4-GeV protons on a uranium carbide target. The atoms were surface ionized, along with stable chromium isotopes. All ions were accelerated to 30 keV, transported via the ISOLDE magnetic high-resolution mass separator to the ISOLTRAP setup [8,19], captured and cooled in the RFQ buncher, and ejected towards the MR-TOF MS. The energy of the ions was adapted by a pulsed drift tube prior to the injection into the MR-TOF MS, where in-trap-lift capturing was applied [20]. After several hundred reflections, the ions were ejected towards the MCP detector.

Figure 2 shows time-of-flight spectra of isobaric ions at mass numbers $A = 52$ and $A = 53$. The calcium peaks were identified by an element-selective laser-ionization scheme. For TOF signals at longer flight times, there are no known isobaric atomic ions except $^{52,53}\text{K}$. The expected flight times of possible molecular ions produced at ISOLDE, such as oxides, dioxides, sulfides, and hydrides, and of doubly charged ions have been compared with the observed peaks. The only species that would come close are $^{36,37}\text{Si}^{(16)O}^+$, that contain short-lived silicon isotopes. However, due to the slow chemical release from the target, the probability for short-lived silicon beam components is very low. More importantly, it can be excluded as only in the case of $A = 52$ would the flight time of $^{36}\text{Si}^{(16)O}^+$ fit the measured peak; at $A = 53$, the measured mass deviates by 1000(130) keV from the value of $^{37}\text{Si}^{(16)O}^+$ [21], which corresponds to a time-of-flight difference in the order of 40 ns.

In 4 h, 12 000 counts were collected for $^{52}\text{K}$, and in 12 h, 2300 counts for $^{53}\text{K}$. Note that with a half-life of just 30 ms, $^{52}\text{K}$ is the shortest-lived nuclide ever investigated with ISOLTRAP. For mass calibration, stable $^{39}\text{K}$ from the ISOLTRAP reference ion source as well as $^{52,53}\text{Cr}$ ions from ISOLDE were used. With the reference-ion masses $m_{1,2}$ and their corresponding flight times $t_{1,2}$, the ion mass of interest $m$ is determined from its time of flight $t$ by [11]

$$m^{1/2} = C_{\text{TOF}}(m_{1}^{1/2} - m_{2}^{1/2}) + (m_{1}^{1/2} + m_{2}^{1/2})/2,$$

where the experimental values are described by the single variable $C_{\text{TOF}} = (2t - t_{1} - t_{2})/[2(t_{1} - t_{2})]$. Because of the

![FIG. 1. Sketch of the part of the ISOLTRAP setup used for the measurements reported here. For details, see the text.](image)

![FIG. 2 (color online). Time-of-flight spectra for nuclides at mass numbers $A = 52$ (top) and $A = 53$ (bottom). Vertical lines indicate the Cr$^+$ fit ranges.](image)
deviation from Gaussian shape of the intense chromium ion peaks as shown in Fig. 2, the mean flight times have been determined from a limited region around the peak maximum (as indicated by vertical lines) that cover about 95% of the measured events. Uncertainties that arise from different choices of this region have been investigated and taken into account.

The resulting mass excesses $ME = M - Au$ are listed in Table I, where $M$ is the atomic mass, $A$ is the mass number, and $u$ is the atomic mass unit. We note that, in principle, $^{52}$Ca, with its mass well known from recent Penning-trap measurements [11,22], would be the second mass reference of choice (in addition to $^{53}$Cr) for measuring $^{52}$K rather than $^{39}$K from the reference ion source, as $^{52}$Ca appears in the same spectrum with $^{52}$K and $^{52}$Cr. In general, such a procedure (used here for the first time) is preferred, because the MR-TOF data of all three isobaric ions are taken simultaneously, excluding drifts between the measurements. The mass values from both calibrations are in agreement. However, the isobaric calibration has a larger uncertainty due to significantly lower statistics of the $^{52}$Ca as compared to the $^{39}$K signal from the reference ion source.

In Fig. 3, the two-neutron separation energies $S_{2n} = B(Z, N) - B(Z, N - 2)$, where $B(N, Z)$ is the binding energy of a nuclide with $Z$ protons and $N$ neutrons, of the isotopic chains from argon to scandium are presented, including the present data. The new $S_{2n}$ values show a significant drop from $^{51}$K [22] to $^{53}$K: about 3 MeV, compared to only about 1 MeV from $^{49}$K to $^{51}$K, clearly illustrating the $N = 32$ shell effect for potassium. The large uncertainty of the $S_{2n}$ value of $^{53}$Sc (271 keV) [21,23] at $N = 32$ leaves room for some ambiguity as to whether the $S_{2n}$ values of calcium and scandium cross. Thus, new precision-mass measurements of the corresponding Sc isotopes would be desirable.

In Fig. 4, the empirical two-neutron shell gap $S_{2n}(N, Z) - S_{2n}(N + 2, Z)$ is plotted as a function of the proton number for $N = 28$ and $N = 32$, including the new point below the magic proton number $Z = 20$. A local maximum of the $N = 32$ shell gap has thus been revealed for the proton-magic $Z = 20$ core. In addition, the plot shows a similar behavior for both neutron numbers going towards the drip line with lower $Z$, where the shell gap is rising up to a maximum at $Z = 20$ and falling by about 1 MeV at $Z = 19$.

As shown already in Ref. [11] for the calcium isotopes, density-functional-theory calculations reproduce the average $S_{2n}$ trend across $N = 28$ and $N = 32$ but do not predict the sharp drops associated with the two shell closures. We extend this study to $Z < 20$ and perform Skyrme-Hartree-Fock-Bogoliubov (Skyrme-HFB) [25] calculations using the HFODD code [26] and interactions of the Skyrme-Lyon family [15], in the spherical approximation.

The calculations use the SLy4 or SLy5 interactions for the particle-hole part and a volume delta-pairing interaction for the particle-particle part of the energy functional, the latter being defined exclusively by a strength parameter $V_0$. With respect to the SLy4 interaction, SLy5 contains in addition a tensor term.
The top panel in Fig. 5 shows the experimental and computed HFB $S_{2n}$ values for the potassium and calcium isotopic chains. The theoretical $S_{2n}$ are computed for nuclei of even neutron number. Self-consistent quasiparticle blocking of the odd protons is performed for the potassium isotopes, by using the procedure described in Ref. [27].

A strength of the pairing interaction of $-200$ MeV fm$^3$ reproduces the very smooth $S_{2n}$ trend observed in Ref. [11]. It describes correctly the experimental values on average but underestimates the drop at the crossing of the magic neutron numbers. A reduction of the strength of the pairing interaction (solid lines) leads to a significant improvement of the description of the experimental $S_{2n}$ trend. The addition of the tensor term with the SLy5 interaction leads to a change in the wrong direction. However, a recent work [28] has shown that the effect of the tensor term in mean-field calculations strongly depends on the way it is constrained to experimental data.

In addition to the empirical HFB approach, it is now possible to perform calculations up to the medium mass region using ab initio methods (see, e.g., Refs. [29–36]).

Thus, new mass calculations have been performed in the ab initio GGF framework [31,37,38] that allow for the study of open-shell nuclei. This method is particularly suited for the present purpose due to the ease of calculating odd-even systems, which also makes it a unique tool to investigate neighboring isotopic chains.

In our calculations, the only input are two- and three-nucleon forces derived within chiral effective field theory at next-to-next-to- and next-to-next-to-next-to-leading order (N$^2$LO and N$^3$LO), respectively [39,40], extended to the low-momentum scale $\lambda = 2.0$ fm$^{-1}$ by means of free-space similarity renormalization-group techniques. The many-body treatment is set by a second-order truncation in the GGF self-energy expansion [37]. Model spaces up to 14 harmonic oscillator shells were employed, and three-body interactions were restricted to basis states with $E_3 \leq 16$. Infrared extrapolations of the calculated ground state energies were subsequently performed following Ref. [41]. We note that, in the present case, this procedure is formally defective due to the different truncations of one- and three-body model spaces. Nevertheless, we find that the trend expected from Ref. [41] is qualitatively reproduced, although with larger extrapolation uncertainties. This is in agreement with other calculations [35]. As an example, we obtain binding energies of 439.52(0.71) MeV for $^{51}$K and 443.31(0.85) MeV for $^{53}$K. This overbinding of about 0.7 MeV/A is a general feature of currently available chiral interactions, and it is a constant effect throughout the whole isotopic chain that cancels in separation energies [31,35,36].

GGF results for $S_{2n}$ of $^{47,49,51,53}$K and $^{48,50,52,54}$Ca are shown in the bottom panel in Fig. 5 and are all resulting from the infrared extrapolation. Different sources of uncertainty affect the present theoretical results (see Refs. [31,38] for a detailed discussion). In particular, this method breaks particle-number symmetry (like HFB theory) and generates the correct expectation values for the proton and neutron numbers only on average, with a finite variance. However, the associated errors are expected to cancel with good accuracy for energy differences (such as $S_{2n}$). The uncertainties indicated in Fig. 5 are uniquely those originating from the extrapolation fit and range between 0.4 and 1.5 MeV with increasing mass number. In general, GGF calculations are in fair agreement with measured $S_{2n}$, with the mismatch at $^{53}$K being on the order of the truncation error. The significant drop from $^{51}$K to $^{53}$K is qualitatively reproduced but overestimated by theory, which also leads to an overestimation of the empirical shell gap for potassium. In contrast to the $N = 28$ gap, which is
quantitatively better reproduced, the overestimation of the $N = 32$ gap emerges as a common feature of \textit{ab initio} calculations in this mass region (see also \cite{35}). The present measurements therefore constitute an important test for the ongoing developments of chiral interactions and \textit{ab initio} methods.

In summary, direct mass measurements performed with ISOLTRAP’s multi-reflection time-of-flight mass separator show that the neutron-shell closure at $N = 32$ extends below the closed proton shell $Z = 20$. The empirical gap is about 1 MeV weaker than for calcium, indicating the stabilizing effect of the $Z = 20$ proton core and highlighting the doubly magic nature of $^{52}$Ca. Mean-field calculations for K and Ca show that interactions with effective parameters, although leading to more easily accessible calculations, depend on the way the parameters are fitted and are thus not always precisely enough constrained to yield robust predictions of shell effects. In contrast, Gorkov-Green function calculations, extended to $N > 32$ along the potassium chain for the first time, offer predictions that are essentially parameter-free and without any fitting to this mass region. By providing novel details on shell evolution, the measurements represent an important test and are instrumental for validating \textit{ab initio} theories and extending them to heavier and more exotic nuclei.

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