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HIGH RESOLUTION LASER SPECTROS-COPY OF THE D LINES OF ON-LINE PRODUCED 21,22,24,25 Na USING A NEW HIGH SENSITIVITY METHOD OF DETECTION OF OPTICAL RESONANCES

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OPTICAL RESUMANCES

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ABSTRACT

A polyisotopic sodium beam of 21-25 Na, produced by spallation of Al, was illuminated with a tunable sw dye laser. The atomic beam, analyzed by a sizpole magnet is then ionized and detected after a mass spectrometer.

The results are : isotopic shifts relative to 23 Na : 21 Na = -53.5 mK, 22 Ma = -25.3 mK, 24 Na = +23.1 mK, 25 Na = +44.6 mK ; y_{I} (25 Na) = +3.685 (22/n.m.; j b-factor in the $P_{3/2}$ state of 25 Na = -16 (8) MHz.

ANALYTIC SUBJECT INDEX : 11.4 ; 54.1

Half-lives ¹, misses ², and other properties ³, of short-lived sodium isotopes have recently been determined using a mass spectrometer on-line with an accelerator. An an extension of this work, we wished to determine other static properties of these nuclei. We have thus attempted to measure the hyperfine structure (hfs) and isotope chift (IS) of the atomic C lines of sodium isotopes. The purpose of this paper is to present the first results that have been obtained using a newly developed method that could eventually extend cur knowledge on the shape and magnetic properties of nuclei far from stability.

The hfs of the ground state of 2^{9-24} has has already beer measured by different methods ⁴; recently a determination of the absolute value of the magnetic moment of 2^{5} ha has been published ⁵. However the hfs of the excited 3P states are unknown for the unstable isotopes and there are no data available for isotope shifts.

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This type of measurement on a series of isotopes can give interesting indications on nuclear shapes. The observation of irregularities of the hfs and IS of short lived Hg isotopes ⁶ has been especially fruitful in that context. In the case of sodium, however, the same method as in ref. 6 would fail because of the Doppler broadening, and it is necessary to use one of the new methods of the Doppler free high resolution spectroscopy ⁷⁻⁹.

The special requirements of our on-line experiment have in fact led to the development of a variant of one 7 of these methods of detection of optical transitions in an atomic beam 10 . In essence this method rests upon the magnetic detection of the optical pumping which occurs when a laser beam is tuned to the frequency of one of the hyperfine components of the D lines.

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This optical pumping changes the magnetic state $(n_{ij}=\pm 1/2)$ of the atoms of an atomic beam and this change is detected by beans of a magnetic filter, a six pole magnet which focuses the atom, with $n_j = \pm 1/2$ and defocuses the atoms with $m_j = \pm 1/2$.

In the actual experiment, the isotopes of sodium are produced in the spallation of aluminum by 150 MeV protons from the Orsay synchrocycletron. Since this nuclear reaction results in a mixture of the different isotopes and their optical resonances might overlap, it is necessary to separate them on-line by a mass spectrometer.

A highly efficient target has been developed in order to have the best sodium isotopes production. We have used a molten metal target in a graphite container. It was loaded with 10 g of high purity aluminum and heated by alternating current up to 900° C. The yield of the 27 Al (p, 3pxn) Ka spallation reaction with 150 MeV protons was studied by directly connecting the target to the thermoionic source of the mass spectrometer. The counting rate after mass separation is shown in Fig. 1. The diffusion delay time in the molten metal is strongly temperature dependent and decreases from 1 mn to 10 s between 850° C and 900° C.

This target is used in the present experiment as the oven of the atomic beam. A schematic view is given in Fig. 2. The collimation of the beam is defined by a nitrogen-cooled diaphragm to 30/1. The sodium atoms effusing from the oven interact with the laser beam before passing the sixpole magnet. The outgoing atoms implies on a hot rhenium surface acting as the ion source of a small magnetic mass spectrometer. Mass separated ions are counted after the exit slit with an electron multiplier. This device detects equally well atoms of stable or short lived isotopes.

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The laser is a single mode tunable by laser of high stability which has been used in other experiments ¹¹. A new frequency control and scanning system can provide a frequency of angle by increments of 7.5 MHz (0.25 mK) initiated by pulses from , pilot clock : this system, called " sigma-meter " will be described clocwhere ¹². A small part of the laser light is used to observe the fluorescence is signal from an auxiliary ²³Hz atomic beam which serves as a reference.

For each isotope, the hfs of its D_1 line is recorded in a single scan, together with the corresponding line of 23 Na. The scanning time is about 10 mm. The signals are directly monitored on a 2-channel chart recorder and fed simultaneously, by a specially built routing device, into the different subgroups of a multichannel analyzer.

The total beam leaving the oven contains up to 10^8 atoms/s of 25 Na. Because of the different losses, the corresponding counting rate at the detector is then 3000 per second. The background of the electron multiplier due to nuclear reaction is only 30-50 counts/s. It should be noted however that tailing of 23 Na ions at the mass 22 results in up to 1000 counts/s. This could certainly be improved in the future by adding an electrostatic deviation after the magnetic mass spectrometer.

Fig. 3 shows the spectra obtained. Our results on the hfs of the ${}^{2}S_{1/2}$ ground state are in agreement with the known values obtained by radiofrequency techniques for 21 , 22 , 24 Na within the limits of the statistical uncertainties of the signal and accidental irregularities of the scanning by the signa-meter. Our value for the magnetic moment of 25 Na is $\mu_{1} = +3.635$ (22)m.m.. The value $|\nu_{1}| = 3.710$ (70) given by M. Deimling et al. 5 is in agreement with our number. The measured values of the isotope shifts are given in table 1.

It appears that the measured values follow the law of the normal plus specific mass effect within the experimental errors. This type of experiment can also lead to a determination of the nuclear electric quadrupple moment if the hyperfine structure of the ${}^{2}P_{3/2}$ state can be measured with a good precision. For that purpose, we have also observed the four (2,3 - 2,3) transitions in the D_2 line of 25 Ka, which were only partially resolved with the present collimation of the atomic beam. From this measurement we have estimated the b factor for the ${}^{2}P_{3/2}$ state to be b = -15 (8) Min. An experiment with a better collimation for the measurement of b is in progress.

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FOOTNOTES AND REFERENCES

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Isotopes	Isotopic shift (mK)
²¹ Na - ²³ Na	- 53,5 (2)
²² Na - ²³ Na	- 25,3 (2)
²⁴ Na - ²³ Na	+ 23,1 (4)
25 _{Na -} 23 _{Na}	+ 44,6 (2)

TABLE 1

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Relative isotopic shifts of 21-25

FIGURE CAPTIONS

Figure 1 - Isotopic yield of the ²²Al (p. 3pxr)^{25-x} We spalletion reaction with 150 MeV protono.

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Figure 2 - Schematic view of the experiment.

Figure 3 - hfs and IS of the D₁ lines of $^{21-25}$ Na isotopes. These lines, measured individually relative to 23 Na are here shown together or a common wavenumber scale.

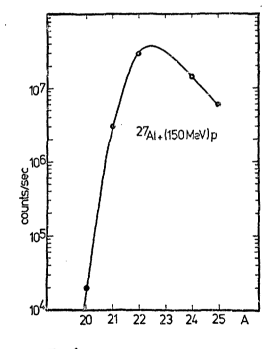
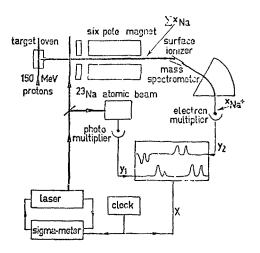


Fig 1

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Fig. 2





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