An accurate Rb density measurement method for a plasma wakefield accelerator experiment using a novel Rb reservoir

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Abstract
A method to accurately measure the density of Rb vapor is described. We plan on using this method for the Advanced Wakefield (AWAKE) project at CERN, which will be the world’s first proton driven plasma wakefield experiment. The method is similar to the hook method and has been described in great detail in the work by W. Tendell Hill et. al. In this method a cosine fit is applied to the interferogram to obtain a relative accuracy on the order of 1% for the vapor density-length product. A single-mode, fiber-based, Mach-Zehnder interferometer will be built and used near the ends of the 10 meter-long AWAKE plasma source to be able to make accurate relative density measurement between these two locations. This can then be used to infer the vapor density gradient along the AWAKE plasma source and also change it to the value desired for the plasma wakefield experiment. Here we describe the plan in detail and show preliminary results obtained using a prototype 8 cm long novel Rb vapor cell.

Keywords: proton driven plasma wakefield, AWAKE, plasma source, accurate density measurement, rubidium vapor source, Mach-Zehnder interferometer, cosine fit

1. Introduction
The AWAKE collaboration will be conducting the world’s first proton driven plasma wakefield experiment at CERN starting in 2016. A unique plasma source is required for this experiment. It consists of a ∼ 10 m long rubidium (Rb) vapor confined in a 4 cm diameter tube. A simple schematic is shown in Fig. 1. The vapor is fully laser ionized (first Rb electron), therefore the plasma density is equal to the vapor density. The Rb reservoirs located at both ends continuously provide Rb vapor. The middle 10 m section consists of a fluid heat exchanger and keeps the temperature (T) within T ±0.5 K along the source; therefore, the vapor density profile (with a constant positive, negative or no gradient, also the plasma density profile) along the source is determined by the Rb vapor flux from the ends or by the temperature uniformity in absence of flow. By measuring the Rb density accurately at the ends one can infer the value of the density gradient.

2. The Method
Density measurements of alkali metal vapors can be made by using the anomalous dispersion around the atomic transition from their ground state and in the visible range. A Mach-Zehnder type interferometer is used (see Fig. 2), where a beam of white light (i.e. broad spectrum visible light) is split and travels through the two arms of the interferometer and is sent to an optical imaging spectrometer. Near the resonance line, the light is subject to a large phase change that is caused by the significant change of index of refraction of the alkali metal vapor. Therefore, when an alkali metal vapor column is placed in one of the arms of the interferometer the vapor density from the interferogram can be determined to a high degree of accuracy.

If the two arms of the interferometer are not parallel then the interfering beams have different phase shifts for different positions along the spectrometer slit (vertical axis of the interfero-
Here, $\lambda$ is the wavelength, $l$ is the length of the vapor column, $e$ is the optical path length difference between the two arms without the alkali metal vapor, the sign depends whether the vapor is in the long or short arm of the interferometer. The complex index of refraction of a gas at density $N_i$, assuming all the electrons are in the ground state (i.e. no excitation), in mks units is given by

$$n = \sqrt{1 + \frac{f_1 e^2 N_i / (m_i e_0)}{\omega_1^2 - \omega^2 - i\gamma_1 \omega}}$$  \hspace{1cm} (1)$$

and in cgs

$$n = \sqrt{1 + \frac{4\pi f_1 e^2 N_i / m_e}{\omega_1^2 - \omega^2 - i\gamma \omega}}$$  \hspace{1cm} (2)$$

The constants are given in table 1. The sum is over all the atomic lines; however, the main contributions come from two lines at $\lambda_1 \sim 780$ (known as the D2 line) and $\lambda_2 \sim 795$ nm (known as the D1 line). Therefore we write with $\omega_i = 2\pi c / \lambda_i$

$$n = \sqrt{1 + \frac{f_1 e^2 N_i / (m_e e_0)}{(2\pi c/\lambda_1)^2 - \omega^2 - i\gamma_1 \omega} + \frac{f_2 e^2 N_i / (m_e e_0)}{(2\pi c/\lambda_2)^2 - \omega^2 - i\gamma_2 \omega}}$$  \hspace{1cm} (3)$$

where $N$ is the electron number density of the ground state and is equal to the vapor density. The formula above ignores line broadening. For our temperature range ($< \sim 200$ °C) the Doppler broadening effect dominates over other broadening mechanisms. The index of refraction including Doppler broadening is [9]:

$$n = \left[1 + \int_{-\infty}^{+\infty} \frac{e^{-\nu t}}{2\sqrt{\pi} \Omega} \frac{d\Omega}{\omega_i + \Omega} \frac{1}{(\omega_i + \Omega)^2 - \omega^2 - i\gamma_1 \omega} + \int_{-\infty}^{+\infty} \frac{e^{-\nu t}}{2\sqrt{\pi} \Omega} \frac{d\Omega}{\omega_i + \Omega} \frac{1}{(\omega_i + \Omega)^2 - \omega^2 - i\gamma_2 \omega} \right]^{1/2}$$  \hspace{1cm} (4)$$

where $\Omega_i = \frac{2\pi c}{\lambda_i} \sqrt{\frac{M_i}{\hbar^2}}$. $T$ is the temperature of Rb, $M$ is the atomic mass of Rb, $M = 85.4678 / 6.02214 \times 10^{23}$ g, $k$ is the

| Table 1: Various constants |
|---------------------------|
| constant | mks | cgs |
| mass,$m_i$ | 1.000023215 × 10^{-26} kg | 1.000023215 × 10^{-28} g |
| charge,$e$ | 1.602176487 × 10^{-19} coulombs | 4.80320989 × 10^{-10} esu |
| permittivity,$\epsilon_0$ | 8.854187817 × 10^{-12} Farad m^{-1} | |
| wavelength,$\lambda_1$ | 780.24 × 10^{-7} m | 780.24 × 10^{-9} cm |
| wavelength,$\lambda_2$ | 794.98 × 10^{-7} m | 794.98 × 10^{-9} cm |
| oscillator strength,$f_1$ | 6.06777 × 10^{-36} | 6.06777 × 10^{-36} |
| oscillator strength,$f_2$ | 3.42102 × 10^{-36} | 3.42102 × 10^{-36} |
| level life time,$\gamma_1$ | 26.24 × 10^{-9} s | 26.24 × 10^{-9} s |
| level life time,$\gamma_2$ | 27.50 × 10^{-9} s | 27.50 × 10^{-9} s |
| atomic absorption coefficient,$g_1$ | 3.813107 m^{-1} | 3.813107 m^{-1} |
| atomic absorption coefficient,$g_2$ | 3.613107 m^{-1} | 3.613107 m^{-1} |
Boltzmann constant, \( k = 1.3807 \times 10^{-23} \text{ JK}^{-1} \). For \( T = 500 \text{ K} \), \( \Omega_1 = 2.30 \times 10^{10} \text{ s}^{-1} \) and \( \Omega_2 = 2.26 \times 10^{10} \text{ s}^{-1} \), \( c \) is the speed of light, \( c = 299792458 \text{ m/s} \). The doppler coefficients are determined by

\[
\int_{-\infty}^{+\infty} \frac{1}{2 \sqrt{\pi}} e^{-\frac{\omega_1^2}{\Omega_1^2}} d\Omega + \int_{-\infty}^{+\infty} \frac{1}{2 \sqrt{\pi}} e^{-\frac{\omega_2^2}{\Omega_2^2}} d\Omega = 1.
\]

There is no analytical solution to the integral therefore we numerically integrate it after separating the complex and real parts. We plot the doppler broadened index and susceptibility in the range of \( \Omega_1 \) in Fig. 5.

As can be seen from the Fig. 5 the effect of the doppler broadening on the real part of index of refraction more than 1 nm away from the absorption line on both sides is negligible. The absorption also plays no role for the purpose of this calculation so far away from the resonance line; therefore, the index of refraction in cgs is approximately given by

\[
n_n(\lambda) = 1 + \frac{e^2 N f_1 \lambda_1^3}{4 \pi n_0 (\lambda - \lambda_1)} + \frac{e^2 N f_2 \lambda_2^3}{4 \pi n_0 (\lambda - \lambda_2)}.
\]

In order to determine the density-length product one needs to record three spectra, i.e. \( I_1(\lambda), I_2(\lambda), I_T(\lambda) \) and then apply a cosine fit to \( S(\lambda) \) given below. The amplitude, \( A \), is also kept as a fitting parameter because of the imperfect of the fringes.

\[
S(\lambda) = A \frac{I - I_1 - I_2}{2 \sqrt{I_1 I_2}} \quad (5)
\]

\[
S(\lambda) = A \cos \left( \frac{2\pi}{\lambda} \left( \frac{e^2 N f_1 \lambda_1^3}{4 \pi n_0 (\lambda - \lambda_1)} + \frac{e^2 N f_2 \lambda_2^3}{4 \pi n_0 (\lambda - \lambda_2)} \pm \epsilon \right) \right) \quad (6)
\]

3. Experimental Set-up and Results

A novel oil heated reservoir was developed and used for this study (see Fig. 6). An oil bath with a circulating pump is used to heat the Rb reservoir. Being immersed in circulating oil keeps

**Figure 5:** Real and imaginary parts of the index of refraction is plotted with (dashed lines) and without (continuous lines) Doppler broadening. The y axis is cut in order to better see the Doppler broadening effect.

**Figure 6:** (a) Drawing of the oil heated Rb reservoir, oil bath with heater not shown. (b) Photo of the Rb reservoir.

**Figure 7:** Rubidium vapor density (blue line) and vapor pressure (green dashed line) as a function of temperature. Region between 1 cm and 2 mm, and the corresponding temperature shows the parameter range of interest for the PDPWFA.
The temperature of the entire system uniform and one can determine the temperature of the Rb liquid, which determines the vapor density, using the probe inserted in the temperature pocket (a thin stainless steel tube) as shown in the schematic (Fig. 6 (a)). The reservoir contains an in-house modified all metal angle valve. The modification allows additional oil sealing and the elongated arm reduces heat losses. The reservoir is sealed using the lid, filled with silicon oil and then insulated against heat losses using insulation wool on the outside. When heated the Rb vapor fills the region of the DN 16 cross which has sapphire diagnostic viewports.

For the measurements, the reservoir is placed in one of the arms of the single mode fiber-based, Mach-Zehnder set-up as shown in Fig. 8. A white light laser is used as the light source. The high intensity laser is chosen since for the AWAKE experiment light is transported away from the radiation area (see Fig. 9). One can also extend the fit to include the central region by using a convolved fit function. The density corresponding to the vapor pressure curve for this temperature is $5.9 \times 10^{14}$ cm$^{-3}$ and the fit gives $6.0 \times 10^{14}$ cm$^{-3}$, which is within expected absolute accuracy (5%) of vapor curve expression [10]. For the AWAKE experiment, absolute accuracy is not so important since the required quantity for determining the density gradient is a relative measurement between the two locations.

We assess the accuracy of the density measurements in two ways. One is from a single spectra including the experimental errors in y axis as follows. We generate multiple random data sets (each corresponds to a virtual experiment) according to the experimental errors. Uncertainty in the y axis is the poisson noise which is the square root of the number of photoelectrons (note that number of CCD counts must be converted to photoelectron count, the specified number by the manufacturer is ~ 1 counts/photoelectron). Then non-linear fitting is applied to each random data set, the standard deviation ($\sigma$ ~0.2%) that results from these data sets gives the uncertainty in the density measurement. This statistical analysis of calculating uncertainty using experimental errors from a single spectra is ideally equivalent to taking multiple shots and taking the standard deviation of the data set. Since the multiple shots will be randomly distributed according to the experimental uncertainties. Doing so for multiple spectra taken consecutively we find that the density fluctuation remains small (namely $\sigma$ ~0.5%). Vibrations contribute random phase shifts, but this is handled by the fitting algorithm.

4. Implementation to the AWAKE Proton Driven Plasma Wakefield Accelerator Experiment

For the AWAKE experiment two ports at the ends of the plasma source are planned to be used. Each Rb vapor source has an independent temperature controller. The relative accuracy of the temperature measurement between the two reservoirs is planned to be around ±0.05 K (using platinum resistor temperature sensors, pt-111, or silicon diode sensors, DT670). Although the temperature can be adjusted to such high accuracy, the temperature of the coldest point of liquid Rb may not correspond to the set temperature. Since the coldest point determines the vapor density this introduces a challenge. This can be handled by combining temperature measurement with the high accuracy interferometric density measurement. To see this we calculate the fractional change in the density ($\frac{d\rho}{\rho}$) corresponding to the fractional temperature adjustment $\frac{dT}{T}$ using the Clausius-Clapeyron relation. The vapor pressure is given by

$$P = P_i e^{-\frac{dV}{k_B T_i}}$$

and

$$\frac{dP}{P} = \frac{h}{k_B T}$$

(7)
where \( h = 1.1 \times 10^{-19} \) joules is the heat of vaporization per Rb atom and \( T \) is the temperature of the liquid. According to the ideal gas equation:

\[
p = nkT
\]

where \( p \) is pressure and \( n \) is density. Therefore at constant temperature

\[
\frac{dp}{p} = \frac{dn}{n}
\]

and using eq. 7 one can calculate the effect of change of liquid Rb temperature on the density. For example for the optimum AWAKE plasma density \( n = 7 \times 10^{14} \text{ cm}^{-3} \) which corresponds to a temperature \( T \sim 466 \text{ K} \) (given by the vapor pressure curve of Rb [10]) (see Fig. 7), we get:

\[
\frac{dn}{n} = 17.09 \frac{dT}{T}.
\]

Therefore, a temperature change of 0.1 K, changes the Rb vapor density by 0.36%. Hence, using this temperature control knob which is capable of such fine adjustments with the interferometric measurement described here one can adjust the two Rb sources relative two each other and independently verify that the relative density measurement is consistent.

Note that in this method one measures the density length product. Therefore the length of the vapor column of the two different reservoirs needs to be known to a high accuracy. This can be done by using commercial mechanical micrometers with 1 micrometer accuracy. Since the vapor column is planned to be a few centimeters long, the uncertainty in length measurement is negligible.

5. Conclusion

In conclusion, an interferometric method to measure Rb vapor density length product to high relative accuracy \( (n \pm 0.5\%) \) is presented. Initial measurements performed using a novel Rb reservoir confirms the feasibility of the method. This diagnostic is planned to be used for the world’s first proton driven plasma wakefield experiment, AWAKE. The unique 10 m long AWAKE plasma source density profile can be determined using this method.

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