A novel fast timing micropattern gaseous detector: FTM

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Abstract

In recent years important progress in micropattern gaseous detectors has been achieved in the use of resistive material to build compact spark-protected devices. The novel idea presented here consists of the polarisation of WELL structures using only resistive electrodes. This allows a new device to be built with an architecture based on a stack of several coupled layers where drift and WELL multiplication stages alternate in the structure. The signals from each multiplication stage can be read out from any external readout boards through the capacitive couplings. Each layer provides a signal with a gain of $10^4 - 10^5$. The main advantage of this new device is the dramatic improvement of the timing provided by the competition of the ionisation processes in the different drift regions, which can be exploited for fast timing at the high luminosity accelerators (e.g. HL-LHC upgrade) as well as applications outside particle physics.
1 Introduction

Recent years have seen much progress on micropattern gaseous detectors (MPGD) [1, 2], mostly under the RD51 umbrella [3]. These detectors have been introduced using well established photolithographic technology on PCB supports. However, they might suffer discharges due to inherent design feature: the very small distance between anode and cathode electrodes limiting the gain of a single structure to $\sim 10^3$ [4]. In Micromegas [5] the problem of the spark occurrence between the metallic mesh and the readout PCB has been solved with the introduction of a resistive layer deposition on top of the readout [6]. Recently, resistive layers have been adopted in WELLs [7]. Instead in GEMs [8] the adopted solution is to share the gain among different multiplication stages [9, 10] without the need of a resistive layer.

This class of detectors is now being exploited in many applications since: they exhibit good spatial and time resolution, high rate capability [11]; they are cost-effective and can be used for large sensitive area [12]; they are flexible and have been used for different geometries [13]. Though time resolution of a few nanoseconds is perfectly adequate in several applications, it would represent a limiting factor in others, where more precise timing is required, for example triggering at high luminosity and medical imaging. To improve the time resolution, the novel detector, described in this paper and named Fast Timing Micropattern (FTM) detector, is based on a series of fully resistive WELL. The use of only resistive layers to polarise the drift and the multiplication volumes allows the construction of consecutive drift-multiplication volumes. In fact, the extraction of signals in any multiplication stage is possible since the overall structure is transparent to the signal, which can be picked up by external electrodes located on top and bottom of the new device. The timing of the ionisation processes in the respective drift volumes will then compete leading to a decrease of the arrival time of nearest ionisation to any multiplication volume and consequently, a decrease of the fluctuations and consequently an improvement of the time resolution.

In this paper we describe the working principle of this new detector, the FTM, of which a first prototype has been realised in the TE-MPE-EM Workshop at CERN and operated successfully for the first time in December 2014. This is one of the implementations described in the European Patent Application number 14200153.6.

2 The working principle

Micropattern detectors are characterised by a clear division of the various phases of the detection process, in particular the primary ionisation creation and drift is separated by the multiplication phase and so the gain of the signal. Thus, the time resolution is dominated by fluctuations of the nearest distance of the primary ionisation processes to the region where the gain is acquired, $d_{\text{near}}$. Defining $\lambda$ as the average number of primary clusters generated by an ionising particle inside the gas, this distance follows a classical exponential distribution $d_{\text{near}} = \exp(-\lambda x)/\lambda$. The drift velocity of the gas $v_d$ determines the arrival time, the exponential behaviour is shown in Fig. 1 and contribution of $v_d$ to the time resolution is

$$\sigma_t = (\lambda v_d)^{-1}.$$  (1)

Both gas parameters depend mainly on the gas mixture used in the device and, in addition, $v_d$ is also a function of the electric field. Typical values for gases employed in MPGDs are $\lambda = 3 \text{ mm}^{-1}$ and $v_d$ up to 0.1 mm/ns leading to few ns time resolution with the best choice of gas mixtures and operating voltages. By comparison, the contribution of the gain fluctuation is governed by the gas time constant $(\eta_{\text{eff}} v_d)^{-1}$, where $\eta_{\text{eff}}$ is the effective Townsend coefficient. In the multiplication volume the field is very high ($\sim 50 - 100 \text{ kV/cm}$) and consequently $\eta_{\text{eff}} > 100 \text{ mm}^{-1}$ and $v_d > 0.25 \text{ mm/ns}$ with a resulting contribution to the time resolution much below 100 ps.

In order to improve the time resolution a new configuration is proposed and is shown in Fig. 2. The improvement is obtained by using several drift regions each one coupled to its multiplication stage,
which is realised with a fully resistive WELL structure. In Fig. 1 results of a simulation are shown for configurations using up to four drift regions, which already demonstrate the benefit. The reduction of the time resolution, in fact, is proportional to the number of the layers \( N_D \) employed, leading Eq. 1 to be modified as follows:

\[
\sigma_t = (\lambda v_d N_D)^{-1};
\]

The behaviour is also evident in Fig. 3, where the results of a simulation of timing response, taking into account also the micro behaviour of the drift and avalanche formation with GARFIELD [14] is demonstrated.

3 The detector layout and test set-up

The detector layout of the prototype is based on the concept of the micro-Resistive WELL detector [7]: two layers are used for this first implementation, which is sketched in Fig. 4. The multiplication volume is based on a pair of polyimide foils stacked due to the electrostatic force induced by the polarisation of the foils. The perforated foils are a 50 \( \mu m \) thick Apical KANECA, which is initially coated with diamond-like carbon (DLC) techniques. This coating technique is used to provide, to the perforated polyimide foil, a potential gradient with resistive layers of the order of 800 M\( \Omega \)/\( \square \), replacing the copper coating. The holes are inverted truncated cones with the two bases of 100 \( \mu m \) and 70 \( \mu m \) diameter. The pitch between holes is of 140 \( \mu m \). The second foils are the 25 \( \mu m \) thick XC Dupont Kapton, which have a resistivity of the order of 2 M\( \Omega \)/\( \square \). The drift volumes are 250 \( \mu m \) thick. The planarity is ensured by a set of pillars obtained by PCB technique with photoimageable coverlay. The pillars’ diameter is 400 \( \mu m \) with a pitch of \( \sim 3.3 \) mm. The spacing around the active region is ensured by the use of two 125 \( \mu m \)-thick polyimide foils.

The employment of only resistive layers in the architecture allows signals coming from the layers to be externally extracted thanks to the resulting transparency of polarising electrodes. The prototype has been operated with a Ar/CO\(_2\)/CF\(_4\) = 45/15/40 gas mixture. The different regions have been polarised with the CAEN N1470 power supply in order to give an electric field of 2 kV/cm for the drift regions and 100 kV/cm for the amplification regions for a gain between \( 10^4 \) and \( 10^5 \). The signals from the readout electrode have been sent to the low-noise charge-sensitive ORTEC PC142 preamplifier. The resulting inverted signal outputs have been amplified by the ORTEC 474 NIM module and acquired with a Tektronix TDS 2024C oscilloscope. An X-ray source has been used in order to demonstrated the capability of signal extraction, and the results are shown in Fig. 5 where an average signal is shown in presence and absence of the source. This is the first signal coming from a FTM device allowing the implementation of the principles described in this paper. The detailed studies of the performance for this class of devices has started and will be the subject of future papers.

4 Conclusions and outlook

In this paper we have described a novel class of micropattern gaseous detectors for fast timing applications called the FTM. The construction feasibility has been demonstrated by building a first working prototype. We expect that this technique can be exploited for applications in high energy physics experiments, particularly for upgrades at LHC where sub nanosecond time resolutions are critical for particle identification and vertex separation. Other applications include X-ray diffraction studies and fast time-resolved measurements offer excellent medical imaging opportunities. In combination with an X-ray convertor and FTM and a visible photocathode shows great promise for use in digital mammography. Other applications include X-ray astronomy by exploiting time resolution of the FTM and selective sensitivity to soft X-rays. For further studies, the subject of future publications, we are building prototypes with several layers, and in parallel investigating larger size cost effective production techniques.
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References

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**Fig. 1:** Timing distribution of the fastest ionisation process. The dotted line histogram represents the distribution for a single drift volume. The dashed line histogram is the result obtained in a double layer configuration. Finally the full histogram is the the distribution obtained in a configuration with four layers.
Fig. 2: Schematic of the working principle of the FTM. Drift and gain processes alternate in the overall configuration that is a stack of several detection layers. Electrons from the primary ionization clouds drift towards the multiplication volumes and timing is determined by the cloud nearest to the respective gain region, which is represented by the minimum of the layer’s $d_{near}$. 
Fig. 3: Simulated time resolution of the FTM device as a function of the number of WELL layers. The simulation has been performed with two standard gas mixtures: the full squares represents the time resolutions with a mixture of $\text{Ar}/\text{CO}_2 = 70/30$; the full circles those obtained with $\text{Ar}/\text{CO}_2/\text{CF}_4 = 45/15/40$. 
Fig. 4: Drawing of the first implementation of the FTM. The basic structure used to build the prototype consists of two layers of full resistive WELL with DLC coating on the top (dark blue surface) on the perforated foils (yellow volumes) and amplification volumes are closed by the antistatic polyimide foils (brown volume). The red cylinders are the pillars. In light blue the pick-up electrode is represented. In the zoomed area is visible the detail in 2D of the fully resistive WELL.
Fig. 5: Average FTM signal with X-ray source on (full line) and off (dotted line).