

Effects of pulsed potential on address electrode in a surface-discharge alternating-current plasma display panel

Jeong Chull Ahn,^{a)} Youichi Shintani, Kunihide Tachibana,^{b)} Tetsuo Sakai,^{c)}
and Naoki Kosugi^{d)}

Department of Electronic Science and Engineering, Kyoto University, Kyoto 606-8501, Japan

(Received 3 January 2003; accepted 2 April 2003)

The influence of pulsed potential application onto the address electrode of an ac-type plasma display panel was investigated from the observation of front and side views of Xe atom densities in the metastable ($1s_5$) and excited ($2p$) states in a unit discharge cell by using microscopic laser absorption spectroscopy and optical emission spectroscopy. It was seen that a predischARGE occurs between the address electrode and one of the sustain electrode worked as a preceding anode, which is a similar effect found previously by applying a higher sustain voltage. The predischARGE partially erases the surface charge accumulated in the preceding pulse, but it induces a faster main sustain discharge, bowing towards the address electrode, and enhances the production efficiency of Xe($1s_5$) atoms, if the applied potential is within an optimal range. © 2003 American Institute of Physics. [DOI: 10.1063/1.1580638]

One of the most important issues of the present alternating-current plasma display panels (ac-PDPs) is to improve luminous efficiency, which is dependent on the composition of rare gases, total pressure, cell structure, and driving wave forms. Various driving techniques using address pulses in sustain periods have been suggested to improve the luminous efficiency.¹⁻⁵ However, real phenomena occurring in the discharge by applying address pulses have not been experimentally investigated as yet. In this work, we examine the behavior of metastable Xe atoms in the $1s_5$ state, $\text{Xe}^*(1s_5)$, under this condition by using the microscopic laser absorption spectroscopy (LAS) technique, which we have developed previously.⁶⁻⁹ The near-infrared emission from excited Xe atoms in the $2p$ state, $\text{Xe}^*(2p)$, is also examined, which is composed mostly of the 823- and 828-nm lines and correlates with the vacuum ultraviolet emission at 173 and 147 nm through the cascade transitions to the metastable ($1s_5$) and resonance ($1s_4$) states.^{10,11} As the total gas pressure or the Xe concentration increases, the emission intensity at 173 nm from Xe_2^* excimers increases through the enhanced production efficiency via three-body collisions of $\text{Xe}^*(1s_5)$ atoms with Ne and Xe atoms in the ground states.^{12,13} It is, therefore, of great importance to investigate the total number of $\text{Xe}^*(1s_5)$ atoms for the estimation of the luminous efficiency in ac-PDPs.

The structure of an ac-PDP unit cell shown in Fig. 1(a) was specially designed for the three dimensional (3D) observation of spatiotemporal behaviors.^{8,9} The sustain electrodes of 200 μm width made of indium tin oxide (ITO) film were successively covered with a dielectric layer and a MgO pro-

TECTIVE layer, and set in parallel to each other on the front glass plate with a gap of 75 μm . By using transparent glass prisms of 150- μm thickness as barrier ribs, we could observe the front and side views of the discharge at the same time. On the rear glass plate, the address electrode also made of ITO was located perpendicularly to the sustain electrodes. A mixture of Ne and Xe (5%) was filled at a pressure of 500 Torr. The width, frequency, and voltage of the sustain voltage pulses (V_x and V_y) employed in this experiment were 9.2 μs , 50 kHz, and 200 V, respectively. The width of the address voltage pulses (V_a) was fixed at 800 ns. The timing of V_a at the center was adjusted to the rising edge of V_x , as shown in Fig. 1(b). Hereafter, the time origin ($t=0$) is defined at the rising edge of V_x . The values of V_a was varied from 0 to 90 V. The LAS measurement of the $\text{Xe}^*(1s_5)$ atom density was performed by tuning the laser wavelength to the 823.1 nm ($2p_6-1s_5$) transition. The IR emission from $\text{Xe}^*(2p)$ atoms

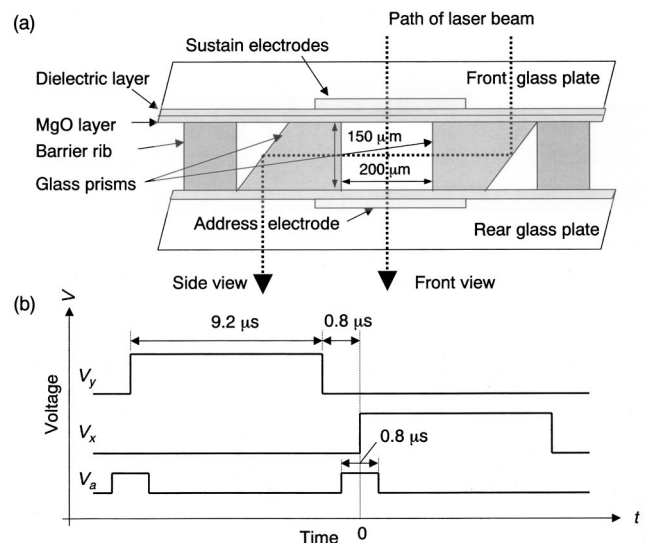


FIG. 1. (a) Structure of the 3D observation cell and (b) wave forms of sustain voltage pulses (V_x and V_y) and address voltage pulses (V_a).

^{a)}Venture Business Laboratory, Kyoto University, Yoshida-honmachi Sakyo-ku, Kyoto 606-8501, Japan.

^{b)}Author to whom correspondence should be addressed; tachibana@kuee.kyoto-u.ac.jp

^{c)}Display Research Laboratories, Co. Ltd., 2-6-12 Tamagawadai Setagaya-ku, Tokyo 158-0096, Japan.

^{d)}Plasma Display Division, Matsushita Electric Industrial Co. Ltd., 1-1 Saiwai-cho, Takatsuki, Osaka 560-1194, Japan.

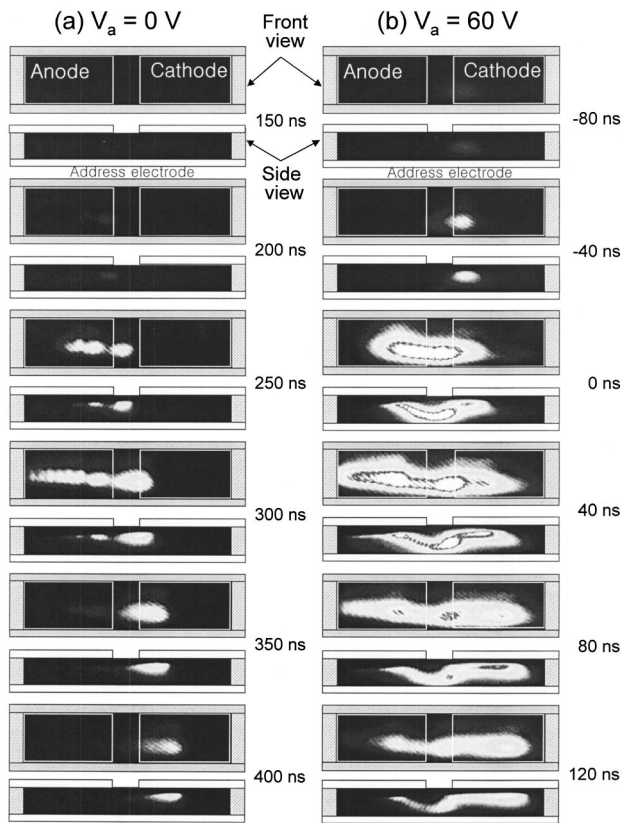


FIG. 2. Temporal sequences of IR emission images measured at address pulse voltages of (a) $V_a = 0$ V and (b) $V_a = 60$ V viewed from front (upper frame) and side (lower frame).

was observed by an intensified CCD camera at a gate width of 10 ns. The CCD camera was equipped with a band-pass filter centered at 820 nm with a full width at half maximum of 20 nm to observe mostly the 823 nm ($2p_6-1s_5$) and 828 nm ($2p_5-1s_4$) lines.⁹

Figures 2(a) and 2(b) show the images of IR emission from $Xe^*(2p)$ atoms observed by the CCD camera from $t = 150$ to 400 ns and $t = -80$ to 120 ns at (a) $V_a = 0$ V and (b) $V_a = 60$ V, respectively. It is noted that the cathode and the anode in these figures are right and left sides, respectively, with respect to the central gap. Front and side views of the discharge image are shown on the upper and lower parts of each frame, respectively. It is seen that the discharge starts faster as the address voltage is applied. The discharge spreads to each end of sustain electrodes but the appearance is different on each side. The emission intensity above cathode is stronger and it expands outward according to the ion mobility, while it becomes weaker and shrinks inward due to charge accumulation on the electrode surface.^{14,15} The occurrence of a predischage between the previous anode and the address electrode is clearly seen at $V_a = 60$ V. The predischage may partially erase the surface charge accumulated on the previous anode, but it induces the main discharge between the sustain electrodes as soon as the next anode pulse is applied. In this discharge mode, the main discharge tends to take a path bowed towards the address electrode. This is apparently quite similar to the case which we observed previously by applying a higher sustain voltage of 250 V, where the charge accumulated on the address electrode due to the

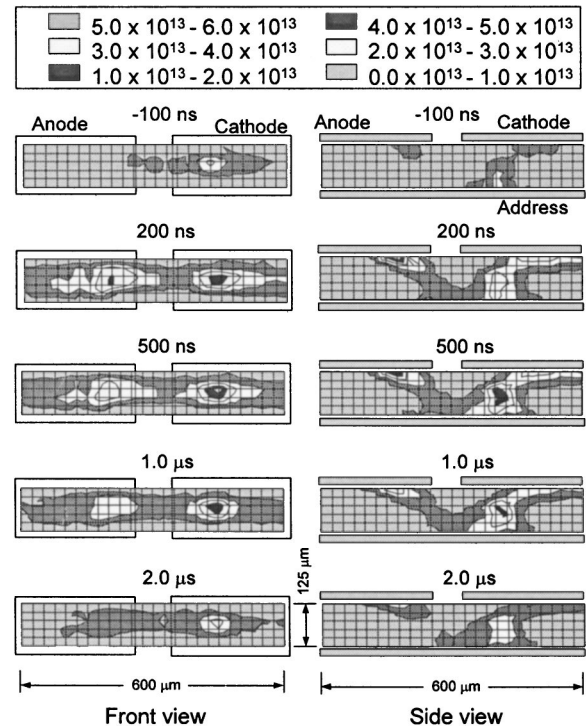


FIG. 3. Temporal sequence of contour diagram of $Xe^*(1s_5)$ atom density (in units of cm^{-3}) at an address pulse voltage of $V_a = 60$ V viewed from front (left column) and side (right column).

stronger discharge causes a similar predischage after the falling edge of the preceding sustain pulse.⁹

Figure 3 shows the measured spatiotemporal behaviors of the $Xe^*(1s_5)$ atom density at the address voltage of $V_a = 60$ V. The graphs on the left side are the distribution of $Xe^*(1s_5)$ atoms in the front view. In each graph, the left-side electrode is working as anode. In a usual operation mode without the application of V_a , $Xe^*(1s_5)$ atoms begin to be observed at the anode side of the gap and a striated pattern is formed.⁷⁻⁹ In this case, however, the excited Xe atoms appear first at the cathode side. From the side view, it is found that the predischage occurs between address electrode and the previous anode (therefore, the succeeding cathode) at $t < 0$, where the address electrode works as the anode, and stimulates the main discharge between the new anode and cathode at $t > 0$. The maximum density of $Xe^*(1s_5)$ atoms reaches $5.5 \times 10^{13} cm^{-3}$ on the cathode side.

The discharge current at the sustain electrode I_x and address electrode I_a were measured by a differential probe through a small resistor of 680 Ω . Figures 4(a) and 4(b) show the current wave forms of I_x and I_a , respectively, which were measured at $V_a = 0, 30, 60,$ and 90 V. Here, we define the response (or delay) time τ as the time difference between the rising edge of the sustain pulse ($t = 0$) and the peak of the sustain current. At a given sustain voltage of 200 V, τ was measured to be 360, 135, 100, and 95 ns at $V_a = 0, 30, 60,$ and 90 V, respectively, showing a tendency of shortening with the increase of V_a . The occurrence of predischage is also seen in the wave forms of I_a at $V_a = 60$ and 90 V, where the address electrode is working as the anode. However, it is noticed that in each wave form of I_a , a smaller negative peak follows the positive peak just after the start of the main sustain current. It suggests that the address

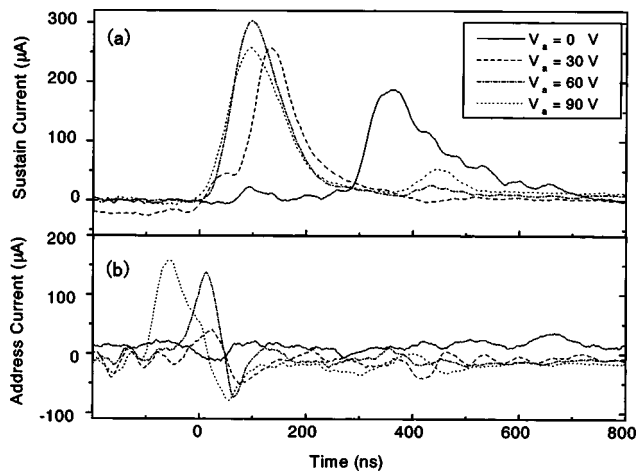


FIG. 4. Wave forms of (a) sustain electrode current I_x and (b) address electrode current I_a measured at address voltages of $V_a=0, 30, 60,$ and 90 V.

electrode turns the role from anode to cathode in the period of sustaining discharge, since the surface of the address electrode has been charged negatively within the predischage period. Therefore, the polarity of the address electrode charge changes to positive after the sustain period, which is favorable for the next predischage phase.

Figure 5 shows the total number of $\text{Xe}^*(1s_5)$ atoms, which is spatially integrated in the whole cell volume, as a function of time. The peak number of $\text{Xe}^*(1s_5)$ atoms increases from 1.3×10^8 to 2.1×10^8 as V_a increases from 0 to

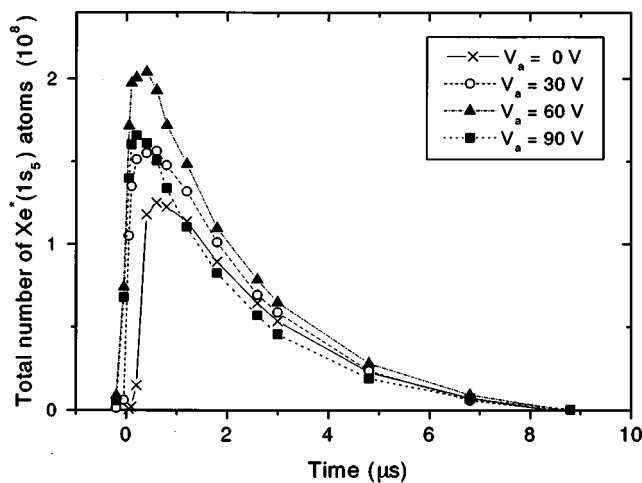


FIG. 5. Temporal behavior of total number of $\text{Xe}^*(1s_5)$ atoms in the whole unit cell volume measured at address voltages of $V_a=0, 30, 60,$ and 90 V.

60 V, but it decreases as V_a increases further to 90 V. At $V_a=90$ V, it is suggested that too much surface charge is lost from the sustain electrode in the predischage, and as the result the main discharge is weakened. At the optimal condition with $V_a=60$ V, the total input power estimated from the current wave form and the applied voltage increases by about 30% as compared to the case without V_a . However, the density of $\text{Xe}^*(1s_5)$ atoms increases by about 70% and, then, the production efficiency increases by about 25% with the power expense at the address electrode of only 6% of the sustain discharge power consumption.

In conclusion, by the application of address pulse potential with a moderate amplitude, say 60 V in our case, the spatiotemporal distribution of the electric field caused by both the surface charges and the external applied voltages becomes optimal across the whole electrode area, leading to an expanded discharge scheme with higher luminance and luminous efficiency. At higher values of V_a , however, the predischage becomes too strong to disturb the optimum balance of the electric field.

This work has been partially supported by Venture Business Laboratory of Kyoto University.

- ¹S.-T. Lo and C.-L. Chen, IEEE Trans. Electron Devices **49**, 762 (2002).
- ²B. G. Cho, H. S. Tae, S. I. Chien, S. H. Lee, and J. Y. Kim, *SID '02 Digest* (Soc. Information Display, Playa del Rey, 2002), p. 440.
- ³J. D. Yi, J. K. Han, K. H. Bu, S. J. Moon, M. S. Kim, W. J. Kim, J. T. Park, H. S. Jang, S. K. Lee, Y. H. Kwon, and J. D. Kim, *SID '02 Digest* (Soc. Information Display, Playa del Rey, 2002), p. 428.
- ⁴H. J. Seo, H. Kim, D. C. Jeong, S. H. Jang, H. S. Tae, S. I. Chien, and K. W. Whang, *Proceedings of Ninth International Display Workshop (IDW'02)*, Hiroshima (Soc. Information Display, Tokyo, 2002), p. 813.
- ⁵K. Yamamoto, H. Kajiyama, K. Suzuki, S. Ho, and Y. Kawanami, *SID '02 Digest* (Soc. Information Display, Playa del Rey, 2002), p. 856.
- ⁶K. Tachibana, N. Kosugi, and T. Sakai, Appl. Phys. Lett. **65**, 935 (1994).
- ⁷K. Tachibana, S. Feng, and T. Sakai, J. Appl. Phys. **88**, 4967 (2000).
- ⁸K. Mizokami, K. Tachibana, N. Kosugi, T. Sakai and Y. Wakabayashi, *Proceedings of Eighth International Display Workshop (IDW'01)*, Nagoya (Soc. Information Display, Tokyo, 2001), p. 1755.
- ⁹K. Tachibana, K. Mizokami, N. Kosugi, and T. Sakai, IEEE Trans. Plasma Sci. **31**, 68 (2003).
- ¹⁰R. Wendt and H. Lange, J. Phys. D **31**, 3368 (1998).
- ¹¹E. H. Choi, J. C. Ahn, M. W. Moon, Y. Jung, M. C. Choi, Y. Seo, G. S. Cho, H. S. Uhm, K. Tachibana, K. W. Whang, and M. Kristiansen, Appl. Phys. Lett. **81**, 3341 (2002).
- ¹²W. G. Lee, M. Shao, J. R. Gottschalk, M. Brown, and A. D. Compagn, J. Appl. Phys. **92**, 682 (2002).
- ¹³J. C. Ahn, T. Y. Kim, J. J. Ko, Y. Seo, G. S. Cho, and E. H. Choi, J. Appl. Phys. **87**, 8045 (2000).
- ¹⁴J. C. Ahn, S. B. Kim, T. S. Cho, M. C. Choi, D. G. Joh, M. W. Moon, Y. H. Seo, S. O. Kang, G. S. Cho, E. H. Cho, and H. S. Uhm, Jpn. J. Appl. Phys. **41**, 860 (2002).
- ¹⁵G. S. Cho, E. H. Choi, J. G. Kim, Y. G. Kim, J. J. Ko, D. I. Kim, C. W. Lee, Y. H. Seo, and H. S. Uhm, Jpn. J. Appl. Phys. **38**, L830 (1999).