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Pixel size limit of the PRIMA implants: from humans to rodents and back

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Abstract

Objective. Retinal prostheses aim at restoring sight in patients with retinal degeneration by electrically stimulating the inner retinal neurons. Clinical trials with patients blinded by atrophic age-related macular degeneration using the PRIMA subretinal implant, a 2 × 2 mm array of 100 μm-wide photovoltaic pixels, have demonstrated a prosthetic visual acuity closely matching the pixel size. Further improvement in resolution requires smaller pixels, which, with the current bipolar design, necessitates more intense stimulation.

Approach. We examine the lower limit of the pixel size for PRIMA implants by modeling the electric field, leveraging the clinical benchmarks, and using animal data to assess the stimulation strength and contrast of various patterns. Visually evoked potentials measured in Royal College of Surgeons rats with photovoltaic implants composed of 100 μm and 75 μm pixels were compared to clinical thresholds with 100 μm pixels. Electrical stimulation model calibrated by the clinical and rodent data was used to predict the performance of the implant with smaller pixels.

Main results. PRIMA implants with 75 μm bipolar pixels under the maximum safe near-infrared (880 nm) illumination of 8 mW mm$^{-2}$ with 30% duty cycle (10 ms pulses at 30 Hz) should provide a similar perceptual brightness as with 100 μm pixels under 3 mW mm$^{-2}$ irradiance, used in the current clinical trials. Contrast of the Landolt C pattern scaled down to 75 μm pixels is also similar under such illumination to that with 100 μm pixels, increasing the maximum acuity from 20/420 to 20/315.

Significance. Computational modeling defines the minimum pixel size of the PRIMA implants as 75 μm. Increasing the implant width from 2 to 3 mm and reducing the pixel size from 100 to 75 μm will nearly quadruple the number of pixels, which should be very beneficial for patients. Smaller pixels of the same bipolar flat geometry would require excessively intense illumination, and therefore a different pixel design should be considered for further improvement in resolution.

1. Introduction

Retinal degenerative diseases, such as retinitis pigmentosa (RP) and age-related macular degeneration (AMD), are the major cause of irreversible visual impairment. RP, a relatively rare class of diseases, originates from various genetic disorders, and typically affects patients in their twenties, eventually leading to profound blindness [1]. Atrophic AMD, on the other hand, causes the loss of central vision later in life due to the gradual demise of photoreceptors, a condition called geographic atrophy. This form of advanced AMD affects millions of patients: about 3% of people above the age of 75, and around 25% above 90 [2, 3]. Currently, there is no therapy for such scotomata and the loss of sight is permanent.

While photoreceptors degenerate in these diseases, the downstream neurons remain largely intact [4–7]. Electrical stimulation of these neurons allows reintroducing information into the visual pathways for restoration of sight. In RP patients, this approach has been demonstrated with an epiretinal prosthesis...
Argus I/II (Second Sight Medical Products, Inc. Sylmar, CA, USA) [8] and with a subretinal prosthesis Alpha IMS/AMS (Retina Implant AG, Reutlingen, Germany), achieving visual acuity up to 20/546 [9, 10]. In AMD patients, subretinal photovoltaic prosthesis PRIMA (Pixium Vision, Paris, France) recently demonstrated its initial safety and efficacy [11, 12]. The PRIMA implant is a photovoltaic array, where each pixel has two photodiodes connected in series between the active electrode in the center and a return electrode on the circumference (figures 1(A) and (B)). The photovoltaic pixels convert images projected from the augmented-reality glasses into patterns of electric current, stimulating the nearby bipolar cells (BCs). The first clinical trial demonstrated a letter acuity closely matching the 100 µm pixel size of the stimulation threshold (1.17 ± 0.13 pixels), corresponding to the Snellen range of 20/438–20/565 [11, 12]. Furthermore, patients simultaneously perceive prosthetic central vision and natural peripheral vision. This inspiring result indicates that smaller pixels may provide higher resolution to benefit a greater number of patients.

Decreasing the pixel size is not trivial since the retinal stimulation threshold increases nearly quadratically with a decreasing width of bipolar pixels [13], and therefore the lower limit of the pixel size is constrained by the ocular laser safety. For an implant of 3 mm in diameter (the expected dimension in future clinical trials), the steady-state temperature increases by about 2 °C under 5 mW mm⁻² near-infrared (NIR, 880 nm) illumination [14]. To limit the temperature rise by 1 °C (half of the recommended thermal safety limit of 2 °C for active implanted medical devices, according to ISO 14708-1) with a maximum duty cycle of 30% (10 ms pulses at 30 Hz, for instance), the peak NIR irradiance should not exceed 8.25 mW mm⁻².

Clinical performance of the PRIMA implants with 100 µm pixels demonstrated comfortably bright visual perception under irradiance of 3 mW mm⁻² at the maximum pulse duration of 9.8 ms and repetition rate of 30 Hz [11, 12]. According to the square scaling of the stimulation threshold with a decreasing size of bipolar pixels in rats [13], pixels of 75 µm would require approximately doubling the light intensity, which should still fit within the ocular safety limit. However, since the human retina is thicker and, unlike the RCS rat retina, the inner nuclear layer (INL) is separated from the implant by about 35 µm [11], scaling with the pixel size might be steeper.

In this study, we evaluate the feasibility of the clinical use of 75 µm pixels by relating the implant performance in rats and humans. First, using the electric field model for PRIMA implants, we convert the measured thresholds with 100 µm and 75 µm pixels in rats from the units of irradiance to the voltage step across BCs. Next, using the clinical data with 100 µm pixels and the modeling results, we predict the corresponding thresholds and maximum stimulus strength with 75 µm pixels in human. Based on benchmarks derived from the successful 100 µm PRIMA implants and their modeling, this approach allows designing and defining the limits of the next-generation retinal prostheses.

2. Methods

All experimental procedures were conducted in accordance with the Statement for the Use of Animals in Ophthalmic and Vision research of the Association for Research in Vision and Ophthalmology (ARVO) and approved by the Stanford Administrative Panel on Laboratory Animal Care. Royal College of Surgeons (RCS) rats were used as an animal model of the inherited retinal degeneration. Animals are maintained at the Stanford Animal Facility under 12 h light/12 h dark cycles with food and water ad libitum.

2.1. Photovoltaic subretinal prosthesis design

To elicit visual perception for restoration of sight, the PRIMA subretinal photovoltaic implants electrically stimulate the second-order retinal neurons, primarily the BCs [15]. Subretinal photovoltaic pixels activated by light [14, 16] act as an optoelectronic substitute for the lost photoreceptors. To reduce the cross-talk between the neighboring pixels, each one has a local return electrode surrounding the active electrode, as illustrated in figure 1(A). Light-to-current conversion in photoreceptors includes a very strong amplification, up to six orders of magnitude: from a few photons per second to pA of current per cell. Since photodiodes do not provide any current amplification and the ambient light intensity is insufficient for electrical stimulation [16], more intense light is required. In addition, capacitive coupling between the electrodes and electrolyte necessitates pulsed charge-balanced stimulation. For these two reasons, the system includes augmented-reality (AR) glasses. Images of the visual scene captured by a video camera are processed and projected by the AR glasses onto a subretinal photodiode array using intense pulsed light. Photovoltaic pixels in the array convert this light into biphasic pulses of electric current, which stimulate the second-order neurons in the INL—primarily the BCs. To avoid perception of this light by the remaining photoreceptors, a near-infrared (NIR, 880 nm) wavelength is used.

This approach offers multiple advantages: (a) thousands of pixels in the implant can be activated simultaneously and independently; (b) since the pixels are activated by light, no wires are involved, which enables reliable encapsulation of the wireless implant and greatly simplifies the surgical procedure; (c) external camera provides autofocusing and adaptation over a wide range of ambient illumination, as well as adjustable image processing optimized for the dynamic range of the implant (the range of stimulus...
strength between the activation threshold and the maximum value supported); (d) the optical nature of the implant maintains the natural link between the eye movements and image perception; (e) network-mediated retinal stimulation retains many features of the natural signal processing, including the antagonistic center-surround [17], flicker fusion at high frequencies and nonlinear summation of the retinal ganglion cell (RGC) subunits [14], among others.

2.2. Surgical procedure and animal handling

The 1.5 mm diameter photovoltaic devices were implanted in the subretinal space of the Retinal College of Surgeons (RCS) rats, typically at 6 months of age, when the outer nuclear layer has degenerated completely. The total loss of the outer nuclear layer was confirmed by the optical coherence tomography (OCT; HRA2-Spectralis; Heidelberg Engineering, Heidelberg, Germany) in each animal prior to surgery. The implants were placed in the temporal-dorsal region, approximately 1 mm away from the optic nerve. The fundus image in figure 1(C) demonstrates the typical location of an implant, and the optical coherence tomography (OCT) in figure 1(D) illustrates the retinal reattachment after surgery. A total of ten animals were implanted with the PRIMA devices consisting of 100 µm (n = 5) and 75 µm (n = 5) bipolar pixels, as described earlier [18]. Animals were anesthetized with a mixture of ketamine (75 mg kg^{-1}) and xylazine (5 mg kg^{-1}) injected intraperitoneally. To visualize the retina and the implant, animals were monitored over time using OCT. For measurements of the visually evoked potentials (VEP), each animal was implanted with three transcranial screw electrodes: one electrode over each hemisphere of the visual cortex (4 mm lateral from midline, 6 mm caudal to bregma), and a reference electrode (2 mm right of midline and 2 mm anterior to bregma).

2.3. Electrophysiological measurements

Following anesthesia and pupil dilation, the cornea was covered with a viscoelastic gel and a cover slip to cancel its optical power and maintain good retinal visibility. The implant was illuminated with 880 nm NIR laser (MF-880 nm-400 µm, DILAS, Tucson, AZ) and a digital micromirror display (DMD; DLP Light Commander, LOGIC PD, Carlsbad, CA) was used for a pattern formation. The customized optical system for the pattern projection into the eye was integrated with a slit lamp (Zeiss SL-120; Carl Zeiss, Thornwood, NY) for real-time observation of the retina via a charged-coupled device (CCD) camera (acA1300-60gmNIR; Basler, Ahrensburg, Germany). Light intensity at the cornea was calibrated before and after each measurement session and scaled by the ocular magnification squared to retrieve the irradiance on the implant. Ocular magnification was defined as the ratio between the size of a projected square on the retina and in air.

VEP were recorded across the ipsilateral and contralateral visual cortices via the Espion E3 system.
Figure 2. (A) Example VEP waveforms measured in an RCS rat with a 75 µm implant. The top waveform, measured with no NIR stimulus, illustrates the noise, while the middle and bottom traces demonstrate above-threshold waveforms. The arrow indicates the amplitude of the bottom VEP waveform. (B) The peak-to-peak VEP amplitude as a function of irradiance. The arrows point to the first irradiance level yielding an amplitude above the noise band for a given pixel size.

Stimulation thresholds were measured with full-field and partial-field (0.65 × 0.65 mm) illumination (figure 3(C)) with 10 ms pulses at 2 Hz repetition rate and irradiances ranging from 0.024 to 4.7 mW mm\(^{-2}\) on the retina. The threshold was defined as the first sampling point where VEP amplitude rises above the noise floor, without any interpolation. To determine the strength-duration relationship of the stimulation threshold, we found the minimum stimulation strength necessary to elicit a significant VEP response at pulse durations ranging from 0.5 to 20 ms.
2.4. Electric field modeling

To relate the optical stimuli to the electric potential across BCs, which is considered the figure of merit for the network-mediated retinal stimulation [20], we represented the electric field in the retina by the weighted sum of the pre-computed basis of elementary fields, each of which corresponds to one electrode injecting a unitary current [21]. Such summation is based on linearity of the Poisson’s equation describing the volume conduction of electric current in biological tissues. The weights are the current injections of the pixels, calculated from the light pattern on the implant as a function of time, with RPSim, a circuit simulator based on Xyce [22]. Besides the active electrodes, each segment of the return mesh in bipolar pixels array also corresponds to an elementary field, where the nominal ground is set at infinity. Due to conservation of charge on the implant, after superposing the elementary fields from the active and return electrodes, the net current to the infinity is always zero. More details regarding the computational framework for assessing the dynamics of the photovoltaic implants [23] are described in the companion paper [24]. The voltage differences between the somatic and the axonal ends of the BCs govern their stimulus strength [20], and we assume their average span is from the bottom of the INL to the middle of the inner plexiform layer (IPL), which was about 52 µm, as measured by OCT.

3. Results

3.1. Electrophysiological performance of PRIMA implants in rodents

To compare the implants’ performance in humans and rodents, we examined the electrophysiological response in rats elicited by 100 µm PRIMA devices. To better fit the size of the rat eye, implants were 1.5 mm in width instead of the 2 mm ones used in clinical trials [11, 12]. Similar devices with 75 µm pixels (figures 1(A) and (B)) were also implanted and evaluated in-vivo. Figure 2(A) shows some typical VEP waveforms measured in RCS rats with 75 µm pixels at various light intensities. Below the stimulation threshold, the waveforms appear indistinguishable from noise. As NIR (880 nm) light intensity increases, the signature peaks start appearing and grow in amplitude, while retaining the timing. To determine the stimulation threshold, we swept across a wide range of irradiances with 10 ms pulses at 2 Hz repetition rate. Figure 2(B) summarizes the peak-to-peak VEP amplitude as a function of irradiance on the retina. The average stimulation threshold was 0.23 ± 0.02 mW mm⁻² with 100 µm pixels, and 0.38 ± 0.06 mW mm⁻² with 75 µm pixels. These are conservative estimates, as we take the first points
where VEP amplitudes rise above the noise floor, without extrapolation. Note that the VEP amplitudes at the highest irradiance (5 mW mm\(^{-2}\)) [12] with 75 \(\mu\)m pixels are similar to those measured at half that irradiance (2.5 mW mm\(^{-2}\)) with 100 \(\mu\)m pixels. This indicates that perceptual brightness with 75 \(\mu\)m pixels under twice higher irradiance might be similar to that with 100 \(\mu\)m pixels, which in the current clinical trials produce comfortably bright percepts with irradiance of 3 mW mm\(^{-2}\) [11, 12].

The perceptual threshold in human patients was measured using a 16-pixel-wide circular spot (1.6 mm in diameter), with 9.8 ms pulses at 10 Hz repetition rate [12, 25]. In rodents, we apply similar pulse duration and repetition rate, but using full-field illumination to maximize the signal and simplify the alignment. To verify whether full-field and partial-field illumination provide similar thresholds, we measured the VEP also using a local square illumination of 0.65 \(\times\) 0.65 mm, as illustrated in figure 3(C). The illuminated local square is approximately half of the total implant area, just as the case in clinical trials. Despite the lower corneal signals with a partial-field illumination (figures 3(D) and (B)), the cortical response amplitude was found to be similar (figures 3(A) and (C)). This indicates some additional signal amplification in the brain, which is not as sensitive to the size of large stimuli on the retina. Therefore, electrophysiological stimulation thresholds derived from full-field illumination may be used for comparison with the perceptual thresholds detected with a large spot illumination.

For further comparison with the clinical results, we also characterized the strength-duration (S-D) relationship of the stimulation threshold with 75 and 100 \(\mu\)m pixels in rats. Figure 4(A) illustrates the stimulation thresholds in terms of light intensity as a function of pulse duration, fitted with the Weiss function for 100 \(\mu\)m and 75 \(\mu\)m implants. The resulting chronaxie for 100 \(\mu\)m pixels was 2.3 ms, in excellent agreement with the chronaxies measured in the PRIMA clinical trial, two examples of which are shown in figure 5(A). Rheobase in the rat measurements (0.17 mW mm\(^{-2}\)), however, was significantly lower than even in the best case of human patients (0.4 mW mm\(^{-2}\)). This is likely due to the fact that, unlike in RCS rats, INL in human patients is separated from the implant by about 35 \(\mu\)m of the debris [12].

3.2. Predictions of clinical performance with 75 \(\mu\)m PRIMA implants

BCs are the target neurons in our subretinal approach, and we have shown previously that depolarization of their axonal terminals can be estimated from the voltage step between their top and bottom boundaries [20]. To assess this voltage step, we converted the stimulation thresholds from the irradiance to voltage using the model of electric field generated by photovoltaic arrays in electrolyte [24]. Figure 4(B) shows the S-D curves for 100 \(\mu\)m and 75 \(\mu\)m pixels measured in RCS rats, converted into the voltage across BCs, estimated to extend from the bottom of INL (starting about 5 \(\mu\)m above the implant) to the middle of IPL, at 57 \(\mu\)m height. Despite the very different levels of light intensity shown in figure 4(A), the voltage S-D curves are nearly identical, confirming that the same voltage step is required for BC stimulation, regardless of the implant configuration. The rheobases for these two fits are in the range of 8–9 mV, and with 10 ms pulses, the threshold voltage is about 11 mV.

To predict the clinical performance of PRIMA implants with 75 \(\mu\)m pixels, we first convert the human stimulation thresholds with 100 \(\mu\)m pixels from light intensity into the voltage step across BCs, and then calculate the light intensity required to generate the same voltage with 75 \(\mu\)m pixels in human retina. Figure 5(A) shows the lowest and the highest stimulation thresholds in AMD patients reported to date with the 100 \(\mu\)m PRIMA implants [12], which we use as the ‘best’ and ‘worst’ cases to estimate the range of responses with future implants. The chronaxies of the S-D relationships are very similar between these two boundary cases (2.2 and 2.3 ms) while the chronaxie of the S-D relationships are very similar between these two boundary cases (2.2 and 2.3 ms) while the rheobase varies substantially: by a factor of 3.5. Conversion to the corresponding voltage thresholds in human retina, where we assume the INL to begin 35 \(\mu\)m above the implant and the middle of IPL at 87 \(\mu\)m, based on clinical OCT images [12], is shown in figure 5(B). For the worst case, a rheobase is about 16.4 mV, roughly twice the rheobase measured in rodents, which may be due to the reduced neural excitability in the badly degenerated retina. In the best case, the stimulation thresholds are more than three times lower, with a corresponding rheobase of 4.7 mV. This value is below the approximate turn-on voltage of the Ca ion channels (>8 mV) [26, 27]. This discrepancy could be reconciled if we assume that dendrites of the BCs in the best-case patient extend below the cell soma, thereby reducing the distance from the implant. If we assume that the BC stimulation threshold in human retina in the best-case scenario is similar to that in rats, the distance between the dendrites and the implant surface should be about 20 \(\mu\)m. This S-D curve, also shown in figure 5(B), lies in between the best and the worst cases, with a rheobase of 9.7 mV.

Given the voltage step across BCs, we can now calculate the light intensity required for generating such a voltage drop with 75 \(\mu\)m pixel arrays in human retina. Figure 5(C) shows the predicted stimulation thresholds in units of irradiance as a function of pulse duration for the two boundary cases. For the worst-case scenario, the rheobase is 3.5 mW mm\(^{-2}\)–2.5 times higher than in the worst case with 100 \(\mu\)m pixels. This increase closely matches the 2.59 ratio of the photosensitive area in 100 and 75 \(\mu\)m PRIMA pixels: 4075 \(\mu\)m\(^2\) vs. 1576 \(\mu\)m\(^2\), respectively. This
similarity indicates that the same current per pixel is required to produce the same voltage drop across the BCs with these two arrays. This, in turn, indicates that higher current density per unit area of the implant with smaller pixels is compensated by stronger field confinement with denser return electrodes grid. With the irradiance of 8 mW mm$^{-2}$, stimulation should exceed the worst-case threshold with pulses longer than 2.6 ms, which is similar to the minimum pulse duration with 100 µm pixels under 3 mW mm$^{-2}$ used in the clinical trial [12].

For the best-case scenario, required irradiance can be predicted two ways. First, using a ‘naive’ voltage threshold, which assumes the BCs residing 35 µm above the implant and having the rheobase of 4.7 mV. The more physiologically realistic voltage thresholds, as described above, correspond to BCs extending their dendrites down to 20 µm above the implant and having the rheobase of 9.7 mV. Interestingly, both models yield very similar S-D curves in units of light intensity, as shown in figure 5(C). Here again, the ratio of the average rheobase with 75 µm pixels (1.05 mW mm$^{-2}$) to that with 100 µm pixels shown in figure 5(A) (0.4 mW mm$^{-2}$) is about 2.6, like in the worst-case scenario. Therefore, clinical performance of the 75 µm implants in the best and the worst cases under 8 mW mm$^{-2}$ irradiance should be similar to that in the clinical trial with 100 µm pixels, where irradiance was about 2.6 times lower—3 mW mm$^{-2}$ [11, 12].

Prosthetic visual acuity is fundamentally limited by the pixel pitch. We have demonstrated that
prosthetic visual acuity matches the pixel pitch of our subretinal implants with pixel size of 75 [14], 55 [28], and 40 [29] µm in rodents, and of 100 µm in patients [11]. In clinical trials, acuity was assessed using the Landolt C optotype, projected with 9.8 ms pulses at 30 Hz. Theoretically, the minimum resolvable font size corresponds to the gap of 1 pixel in the letter C. Patients resolved the fonts in the range of 1.04–1.3 pixels per gap, with the average of 1.17 pixels [12]. To check the expected resolution with 75 µm pixels, we calculated the electric potential across BCs when a Landolt C font with a gap of 1.2 pixel (90 µm) was projected onto the photovoltaic array with the irradiance of 8 mW mm$^{-2}$ and 9.8 ms pulses at 30 Hz (figure 6). As with the stimulation thresholds, we calculated the field maps for two scenarios: if the BCs are separated from the implant by 35 µm and by 20 µm. With the 35 µm separation, we outlined two thresholds for 10 ms pulses: for the best (5.6 mV) and the worst (18.9 mV) cases. For the 20 µm separation, assumed only for the best case, the threshold is 11.8 mV. As shown in figures 6(B) and (C), the predicted contours of visibility for the best case in two scenarios are very similar. The worst case, as shown in figure 6(B), represents a much narrower letter with a wavy outline. However, with the eye movements, this wavy structure may average out and hence not be noticeable by the patients. Amplitude of the electric potential in these maps (57 mV for 20 µm and 28 mV for 35 µm height) is very close to that calculated for the 100 µm pixels under 3 mW mm$^{-2}$ irradiance: 62 mV and 31 mV, respectively [24].

Under 3 mW mm$^{-2}$ illumination of the 75 µm pixels, the voltage steps across BCs are expectedly lower: 23 mV for 20 µm and 11 mV for 35 µm separation between the implant and BCs, respectively. For the worst case, such voltage drop is even below the stimulation threshold, so, clearly, much higher irradiance is required with 75 µm pixels.

Under 8 mW mm$^{-2}$ irradiance with 75 µm pixels, the 1.2 pixel gap in Landolt C is expected to have a similar perceptual brightness and contrast of the pattern as in the current clinical trials, and hence should be resolvable. If resolution of prosthetic vision will not be limited by other factors in patients, this would improve the average visual acuity from 20/500 to 20/380.

4. Discussion

The predicted stimulation thresholds with 75 µm bipolar pixels are higher than those measured with 100 µm pixels by a factor of about 2.5, approximately matching the ratio of the photosensitive areas in these pixels. This means that the same current per pixel is required for stimulation, as would be with the
electric field of a point source. Such a simple approximation is expected with stimulating electrodes smaller than the distance to the target cells and with a return electrode located much further away than the target cells. However, with the BCs extending over more than 50 \( \mu \text{m} \), this is not the case even with RCS rat retina, where INL begins within a few micrometers from the implant. Even less so in AMD patients, where INL is separated from the implant by about 35 \( \mu \text{m} \) [12]. In our case, neurons are separated from the active electrodes and the mesh of circumferential return electrodes in each pixel by distances comparable to the pixel size. In this case, decreasing pixel size in a pattern activation has two counteracting effects: (a) higher current density per unit area generated with smaller pixels injecting the same current per electrode should increase the electric potential, but at the same time, (b) the electric field is diminished by the denser mesh of the return electrodes in the array. These two effects counteract each other, resulting in the threshold scaling according to the constant electric current per pixel in our geometry.

Perceptual brightness and contrast of the patterns with 75 \( \mu \text{m} \) pixels at 8 mW mm\(^{-2}\) is expected to be similar to that with 100 \( \mu \text{m} \) pixels at 3 mW mm\(^{-2}\) irradiance in humans. Since 8 mW mm\(^{-2}\) with a 30% duty cycle is close to the NIR retinal safety limit, further reduction in pixel size of the same geometry is unlikely. In rodents, where INL is closer to the implant, we demonstrated a prosthetic visual acuity matching the pixel pitch even with 55 \( \mu \text{m} \) bipolar pixels [28], but for human retina, such pixel size would require light intensity exceeding the safety limits.

Bipolar pixels effectively prevent the cross-talk between neighboring pixels, but they do it by overly constraining the electric field. Placing a local return electrode inside each pixel creates a dark ring between the adjacent bright spots in the pattern. Ideally, the electric field should be confined such that the bright

**Figure 6.** Landolt C pattern stimulation with 75 \( \mu \text{m} \) pixels. (A) A Landolt C pattern projected on top of a 75 \( \mu \text{m} \) pixel array. The width of the gap in C is 90 \( \mu \text{m} \)—1.2 times the pixel width. (B) The simulated field map for BCs separated by 35 \( \mu \text{m} \) from the implant, whose terminals end in the middle of the inner plexiform layer (87 \( \mu \text{m} \) above the implant surface). The solid line marks the base-case threshold (5.6 mV), while the dotted line, the worst-case threshold (18.9 mV). The label 28 mV indicates the highest voltage step in this map. (C) The field map for BCs separated from the implant by 20 \( \mu \text{m} \), while the location of axonal terminals remain the same as in B. The solid line marks the threshold (11.8 mV) for the best-case at this height. The label 57 mV indicates the highest voltage step here.
pixel creates a stimulation spot, while the dark pixel does not, without a dark boundary around the active pixels. This can be achieved using current steering, if the active electrodes on monopolar pixels could be used for both, injecting and collecting the current, i.e. as the active and return electrodes [18, 21]. Additionally, 3-dimensional electrodes can improve the electric field penetration into the tissue, thereby maintaining a low stimulation threshold and good contrast, such as the honeycomb and pillar layouts [13, 30].

5. Conclusions

Under the maximum safe NIR irradiance (8 mW mm⁻² with 30% duty cycle), the 75 µm PRIMA implants should provide a similar perceptual brightness to that with the 100 µm pixels under 3 mW mm⁻², which is used in the first clinical trials. It should also maintain sufficient contrast for resolving a Landolt C with a gap width of 1.2 pixel, corresponding to the visual acuity of 20/380. A decrease in the pixel size from 100 to 75 µm and an increase in the implant width from 2 to 3 mm will nearly quadruple the number of pixels in an implant, which would be beneficial for AMD patients. Further decrease of the pixel size in flat bipolar geometry would require exceedingly bright illumination, and therefore a different electrode geometry should be considered for further improvement in resolution.

Data availability statement

The data that support the findings of this study are available upon reasonable request from the authors.

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