Co-localization of cell death with antigen deposition in skin enhances vaccine immunogenicity

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Abstract

Vaccines delivered to the skin by microneedles – with and without adjuvants – have increased immunogenicity with lower doses than standard vaccine delivery techniques such as intramuscular (i.m.) or intradermal (i.d.) injection. However, the mechanisms behind this skin-mediated ‘adjuvant’ effect are not clear. Here, we show that the dynamic application of a microprojection array (the Nanopatch) to skin generates localized transient stresses invoking cell death around each projection. Nanopatch application caused significantly higher levels (~65-fold) of cell death in murine ear skin than i.d. injection using a hypodermic needle. Measured skin cell death is associated with modeled stresses ~1–10 MPa. Nanopatch-immunized groups also yielded consistently higher anti-IgG endpoint titers (up to 50-fold higher) than i.d. groups after delivery of a split virion influenza vaccine. Importantly, co-localization of cell death with nearby live skin cells and delivered antigen was necessary for immunogenicity enhancement. These results suggest a correlation between cell death caused by the Nanopatch with increased immunogenicity. We propose that the localized cell death serves as a ‘physical immune enhancer’ for the adjacent viable skin cells, which also receive antigen from the projections. This natural immune enhancer effect has the potential to mitigate or replace chemical-based adjuvants in vaccines.

Keywords

vaccine; skin; microneedles; intradermal vaccination; cell death

Conflict of interest

MAFK is a consultant, advisory board member, shareholder of Vaxxas, and inventor on patents licensed to Vaxxas, a company developing the Nanopatch for drug delivery applications. This company has not yet a microprojection-based product, and the microprojection device presented in this study is not directly related to any microprojection devices currently under development at Vaxxas. CJF is employed by Vaxxas through UQ. CJF, ACID and SCM are inventors on a patent application that is licensed to Vaxxas. The other authors state no conflict of interest and have no competing financial interests.

For full method please refer to Supplementary Information online.
Introduction

As conventional vaccine delivery techniques such as intramuscular (i.m.) and subcutaneous (s.c.) injections bypass the skin’s immune cells, the skin has increasingly become the target for vaccine delivery (Gutowska-Owsiak and Ogg, 2012; Jiang et al., 2012; Kupper, 2012). Immunizations via skin by intradermal (i.d.) injection have demonstrated similar or improved immunogenicity with commonly reported ~5–10-fold dose-sparing, i.e. a significant reduction of antigen required to elicit equivalent immunogenicity compared to other needle-based vaccinations such as i.m. injections against influenza (Auewarakul et al., 2007; Belshe et al., 2004; Hung et al., 2012a; Kenney et al., 2004; Quan et al., 2010; Van Damme et al., 2009), compared to standard i.m. routes. However, difficulties exist in effectively and consistently delivering vaccines into the skin. Therefore, new cutaneous vaccine delivery devices such as microneedles and the Nanopatch (NP) have been developed (Kim et al., 2012a; Koutsonanos et al., 2013; Chen et al., 2009), which may also reduce needle-stick injuries, disease transmission of blood-borne diseases (Ekwueme et al., 2002) and could allow for self-administration (Prausnitz et al., 2009). In our previous work, delivery of a conventional influenza vaccine by NP to skin with antigen alone has achieved ~100-fold dose-sparing compared to i.m. delivery (Fernando et al., 2010), while co-delivery of adjuvant generated synergistically-improved antibody and T-cell immune responses (Fernando et al., 2012; Ng et al., 2012).

The NP is a microprojection array with >20,000 cm⁻² of 100 μm long projections (illustrated in Figure 1a and b) that is applied onto skin at 2.3 ms⁻¹ using a spring-loaded applicator, ensuring consistent penetration across the array (Crichton et al., 2010). While i.d. injection deposits vaccine within the vicinity of dermal APCs, previous studies showed that the NP deposits antigen into both viable epidermis (VE) and dermis (Fernando et al., 2010). A contributing factor to improved immunogenicity may be in the mechanical interaction with the NP with skin. Recently, we have shown that NP projections impacting upon the skin and decelerating upon penetration generate significant stresses within the skin (up to 50 MPa) within a ~20 μm radius around each projection (Meliga et al., 2013). We postulated that such stresses invoke localized cell death and inflammation in the skin, contributing to the improved immune responses generated by NP application by activating APCs to take up antigen.

Following Matzinger’s ‘danger hypothesis’ (Matzinger, 1994), apoptotic and necrotic cells have been reported to act as immunostimulants by releasing damage associated molecular patterns (DAMPs), enhancing cellular and humoral immune responses to antigen (Green et al., 2009; Kinsey et al., 2004; Kono and Rock, 2008; Marichal et al., 2011; Rovere-Querini et al., 2004). Chemical adjuvants have shown to induce cell death and danger signals upon administration which can act as strong endogenous immunostimulatory agents (Marichal et al., 2011; Yang and Shen, 2007; Yang et al., 2004a, b). Disruption of the epidermis by scarification has also been demonstrated as essential for the elicitation of the vaccinia virus-mediated immunity and protection against challenge, whereas i.m. delivery was not protective (Liu et al., 2010).
Here, we hypothesized that localized cell death caused by NP penetrating epidermal and dermal tissue would initiate a cascade of events acting as a ‘natural immune enhancer’, mediated by the release of DAMPS concurrent with antigen delivery. To test this hypothesis, we measured cell death induced in situ by a conventional i.d. Mantoux injection and compared it with NP. Systemic immunogenicity was compared using delivered split virion influenza vaccine (Fluvax®). To our knowledge, this is the only quantitative and spatial study published investigating cell death induced by skin immunization on immunogenicity.

Results

Key physical differences in skin surface morphology following Nanopatch application to and intradermal needle injection into skin

We examined the skin surface pre and post NP and 31G needle (hereafter referred to as needle) application in vivo and ex vivo to gain insights into the skin perforation induced by each device. Using scanning electron microscopy (SEM), we measured the contact area between each device with skin surface (Figure 1b, 1d-f). Untreated murine ear skin revealed an intact surface (Figure 1c), while NP application produced microchannels, consistent with the spacing of projections on an NP (Figure 1d), similar to previous observations (Crichton et al., 2010; Haq et al., 2009; Milewski et al., 2010). I.d. insertion using a needle resulted in a single perforation at the needle insertion site (Figure 1e and 1f, arrows), while saline delivery resulted in the formation of a bleb (Figure 1g). The calculated contact surface area between skin and NP projections was 17.7 ± 2.3 mm$^2$ (n=18), while 1.6 ± 0.1 mm$^2$ (n=10) of the needle tip made contact with skin. Coating of Coomassie blue dye allowed visualization of microchannels inflicted by NP, and confirmed coated payload delivery discretely into each microchannel, where it had dissolved off the projection upon application due to rehydration within the skin (Figure 1h, left). Coomassie blue dye injected i.d. allowed further visualization of the bleb (Figure 1h, right). The affected skin area following NP application (Figure 1h inset and 1i) measured 1.9 ± 0.1 mm$^2$, significantly less than the delivery area of a needle: 27 ± 11 mm$^2$ ($p$<0.001; Figure 1h and 1i). Combined, these data demonstrate significant perforation of skin and discrete microchannel delivery following NP application in contrast to a single i.d. delivery to a much larger area in the form of a bleb when using a needle to inject i.d.

Nanopatch™ application to skin induces significantly higher and localized cell death, compared to standard needle-based i.d. injection

We next examined and quantified cellular damage caused within the skin by NP and i.d. needle over a 16 mm$^2$ area (an area equivalent to a single NP) by Confocal and Multiphoton Microscopy (CLSM/MPM). (Figure 1e–g). We freshly excised skin immediately after NP application or i.d. injection of PBS, and stained with an acridine orange/ ethidium bromide (AOEB) cocktail to distinguish live and dead cells, respectively (Figure 2a and 2b) (Raju et al., 2006). Untreated skin contained 3.5%±1.9% (n=18) damaged cells (Figure 2c). After NP application, 16.5%±4.1% (n=18) of epithelial cells were dead (Figure 2c), significantly more than untreated skin ($p$<0.0001). Following NP application, dead cells were restricted to a zone of up to 5 cell radii, or 26±13 μm (n=81 projections), around each microchannel perforation with interspersed viable cells. High levels of cell death were observed where the
edges of the NP and skin interacted, consistent with the theoretically-modelled stress peaks at the array boundaries (Meliga et al., 2013).

The cellular damage caused by i.d. injection of saline was restricted to the individual needle perforation site (Figure 2a and 2b), inducing 3.7% ± 1.8% (n=18) of dead cells, similar to untreated skin control. Cell death within the i.d. bleb was statistically similar in comparison to untreated control groups (2.5% ± 1.1%; p=0.16, n=5). Normalized against untreated control groups, i.d. injection generated 65-fold fewer dead cells than NP application. The area of dead cells within the bleb area skin represented <0.5% of the total bleb area (~0.07 mm²). Cross-sectional MPM images demonstrated that damaged tissue in NP-treated samples was mostly confined to the VE, while delivery by needle resulted in cell death spread throughout the VE and dermal layers (Figure 2b). The observed dead cells exhibited predominantly necrotic characteristics, which was further supported by the absence of apoptotic cell morphology (i.e. fragmented nuclei or crescent-shaped nuclei (Cobb et al., 1996; Hotchkiss et al., 2009). Further morphological, biochemical and protein-based analyses are required to further distinguish the cell death types. Collectively, these data highlight key differences between the location of live/ dead cells and a delivered payload (e.g. vaccine).

Delivery of vaccine to skin by Nanopatch induces a significantly greater antibody response than standard intradermal delivery by needle

With the local skin cell death and delivered payload profiles quantified, we next compared the resultant systemic immune responses elicited by NP and i.d. immunization routes (Figure 3) by comparing antibody responses to Fluvax® spanning 3 logs of delivered antigen doses. Endpoint titers from all doses of Fluvax® tested were higher after NP delivery than those elicited by i.d. delivery with statistical significant differences detected with delivered doses between 1–100 ng (Figure 3). In addition, all mice given a dose of 5 ng to 100 ng by NP seroconverted, and mean titers (10–100 ng) did not differ statistically from those of the i.m. control group (Supplementary Figure S1). In contrast, ≥50 ng of Fluvax® delivered i.d. was required to achieve seroconversion in all mice. (p<0.001; Figure 3 and Supplementary Figure S1). Together, these data demonstrated robust antibody titers in mice immunized with NP-delivered Fluvax®, compared to i.d. injected mice, illustrating a ≥10-fold dose-sparing effect.

Co-localized Nanopatch-mediated skin damage, adjacent live cells and vaccine enhances systemic immunogenicity

From our immunogenicity results, we hypothesized that co-localization of antigen with Nanopatch-mediated skin damage adjacent to live cells was a key contributing factor to enhanced systemic immunogenicity. To test this hypothesis, we imaged co-localization of Nanopatch-mediated skin damage in vivo, adjacent to live cells with fluorescently-labeled (Dylight® 755) Fluvax® (FV) antigen. Labeled antigen was administered by either NP or i.d. injection and animals were imaged within 5 min post immunization in vivo to limit diffusion of the antigen (rainbow-colored; Figure 4). In NP-treated samples, 100% of dead cells (orange) co-localized completely with fluorescently-labeled antigen by NP delivery, covering an area of 16 mm². Based on the cell death profiles, Coomassie Blue data and
diffusion profiles (Raphael et al., 2013), co-localization of antigen with dead cells was much less pronounced with i.d. delivered antigen (<0.5%) as indicated by the arrows (Figure 4), suggesting lower co-localization of antigen with dead cells (and therefore less DAMPs). These observations of co-localizing cell death with antigen are consistent with our previous results (Figure 1h and 2).

While we identified that more antigen overlapped (co-localized) with dead cells following NP than i.d. immunizations (Figure 4), it was not clear whether co-localization of antigen with dead cells interspersed with live cells was essential for enhanced antibody production. To explore this, we compared antibody responses (10 ng Fluvax®) between mice using NP, i.d. or with a combination, as depicted in Figure 5a). In accordance with Figure 3, Fluvax® delivered by NP elicited higher antibody responses than i.d. (Figure 5b; \( p < 0.05 \)). Application of an NP or an FP (inducing ~3.8% of cell death; Supplementary Figure S2) immediately prior to i.d. delivered Fluvax® did not enhance the resultant immunogenicity of i.d. generated beyond that obtained with i.d. delivery of Fluvax® alone. We interpret this result as: co-localization of tissue damage caused by NP with antigen is important for enhanced immunogenicity. Similar results were obtained when repeated with a 1 ng Fluvax® dose (Supplementary Figure S3). Taken together, with the results in Figure 4, these data demonstrated that co-localization of antigen with NP-induced cellular damage correlated with enhanced immunogenicity of NP-delivered Fluvax®.

Since i.d. injections resulted in a ~14-fold larger antigen delivery area than NPs (Figure 1h and 1i), we then evaluated whether altering i.d. injection volumes and hence concentrations/area while maintaining the same delivered dose (5 ng), may affect the systemic immunogenicity. Administering 5 ng Fluvax® by i.d. in different volumes (5–50 μl), with the highest concentration tested injected i.d. was similar to the concentration of a dry-coated NP (2.5 ng/μl). This difference in antigen concentration did not affect immunogenicity (data not shown). Antigen-specific endpoint titers did not significantly differ between any i.d. groups (\( p > 0.05 \); Figure 5c). These results indicated that vaccine concentration (ranging 1–0.1 ng/μl) did not affect the immunogenicity in i.d. immunizations.

To further examine the correlation between cell death and enhanced immunogenicity, we extended NP conditions to provide lower and higher levels of cell death – by changing the number of projections cm\(^{-2}\) while maintaining all other parameters constant including the delivered dose (assessed as previously described, Fernando et al., 2012). Cell death was quantified while influenza-specific IgG was measured after delivering 1 ng Fluvax® by NPs of varying projections cm\(^{-2}\) (ranging from 10,000, 21,000 to 3x 21,000) or by i.d. injection. As expected, cell death increased with increasing numbers of projections cm\(^{-2}\) almost linearly (\( R^2=0.89 \)), resulting in 2.3% ± 1.0% (untreated), 3.1% ± 1.0% (i.d.), 8.1% ± 1.1% (10,000), 16.2% ± 1.7% (21,000) or 45.4% ± 4.0% (3x 21,000 projections cm\(^{-2}\)) (Figure 5d–f left). This indicated that the localized cell death per projection was consistent. Figure 5f (right) illustrated that increased cell death correlated with increased immune responses, with one notable exception: extending to the highest level of cell death (induced by 3x 21,000 NP projections cm\(^{-2}\)) led to a significant decrease (\( p=0.0074 \)) in antibody response compared to a single NP with 21,000 projections.

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Discussion

Several different mechanical methods exist for delivering vaccines into skin such as either needle-based or needle-free approaches (Kim et al., 2012a; Kim et al., 2012b; Kis et al., 2012). The delivery of conventional protein and live viral vaccines using these devices has resulted in comparable systemic immune responses to those achieved by standard i.d. injection (Carey et al., 2011; Hung et al., 2012b; Kim et al., 2012b). Here, we compared the application two transcutaneous immunization devices into skin. We used either NP, standard needle-based i.d. injection, or combinations of both methods, and quantified the resultant cell death (including magnitude and location within skin) as well as the systemic immunogenicity generated by influenza vaccination. Overall, we found that NP immunization consistently produced higher antibody titers than standard needle-based i.d. immunizations, which correlated with higher cell death levels after NP than i.d. injections. We propose that the observed improved immunogenicity in NP groups is linked to antigen co-localizing within defined areas of live cells adjacent to dead cells within epidermal and dermal tissue (Figure 4); this was not seen with i.d. injection.

The concept that cell death or co-delivering dead cells with antigens enhances immunogenicity is well known (Green et al., 2009; Kinsey et al., 2004; Kojima et al., 2007; Kono and Rock, 2008; Rovere-Querini et al., 2004). Similarly, others found that DNA released by dead cells mediates adjuvanticity of aluminium salts (Marichal et al., 2011). Increasing the number of NP projections penetrating the skin and thus increasing localized cell death led to improved immunogenicity, until a plateau was reached, after which the immunogenicity decreased (Figure 5d–f). This type of curve is homologous to dose ranging vs. immunogenicity that shows similarity with titration curves of chemical adjuvant doses (unpublished observations and Ng et al., 2012). A key difference however is that in our study, we propose the immune enhancing effect is generated by the mechanical stimulus of the NP being applied to the skin, generating localized cell death.

Recently, laser-based adjuvantation gained interest as potential immune enhancer (Chen et al., 2010) that worked synergistically with other chemical adjuvants (Chen et al., 2012). A ‘physical adjuvantation effect’ of the laser was stipulated (Chen et al., 2013). In agreement with our study, colocalization of the laser beam and immunization site was essential for enhanced immunogenicity. We speculate that laser-induced cell damage/ death (and associated DAMP release) colocalized with antigen and thus enhanced immunogenicity similar to the synergistic effect with chemical adjuvants. A key difference in our study in comparison to others was however that the immune enhancer effect was generated by the NP application itself, generating localized cell death. I.d. injection generated significantly lower cell death, therefore DAMPS may have co-localized less with antigen and live cells; this is summarized schematically in Figure 6a. With live cells immediately adjacent to dead cells sensing danger signals and chemokine/ cytokine gradients required for their activation, and with interspersed vaccine, these live cells may be considered as the target cells to which antigen should be delivered to for enhanced immune responses. Therefore, we believe that a certain threshold of cell death is beneficial to augment immunogenicity to an antigen, beyond which cell death becomes detrimental. This would further support the hypothesis of
the NP-induced cell death acting as ‘natural immune enhancer’ contributing to enhanced immunogenicity, compared to standard i.d. injection.

The markedly lower cell death after i.d. injection was localized to a single needle insertion site. Previous investigations of skin cell death induced by mechanical vaccine delivery devices such as tattooing devices were reported to induce necrosis and inflammatory cell infiltration (Gopee et al., 2005) with increased cellular and humoral immune responses (Pokorná et al., 2008). Further, ballistic delivery of micro-particles found >90% dead cells in the center of the target site with only the skin target periphery containing both live/ dead cells (Raju et al., 2006). Our experimental findings contrast Raju et al.’s findings most likely due to the inherently different device and application velocities (2.3 ms\(^{-1}\) vs. >100 ms\(^{-1}\)). However, our experimental findings are consistent with our modeled stress-contours induced by NP application to skin (Figure 6b), where cell death and highest simulated stress levels (red) overlap at the site of projections penetrating the skin (applying previously published mechanical models; Meliga et al., 2013). Qualitative comparison of the spatial distributions of theoretical stress with measured cell death suggests that cells die in regions where Von-Mises stresses of at least 1–10 MPa are generated, subject to modelling assumptions (Meliga et al., 2013). This is the only time that skin cell death has been linked to stress distribution and magnitude to our knowledge.

The type of induced cell death is likely to be important as necrotic cells are generally more immunogenic than apoptotic cells (Rock and Kono, 2008). We speculate that the visualized dead cells are predominantly of necrotic origin, which is in agreement with Gopee et al., who observed necrotic cell death following tattooing in skin (Gopee et al., 2005). Necrotic cell death is further supported by the dynamic application of the NP generating instantaneous stresses and stress-induced trauma in Figure 6b (detailed analysis on the stresses in Meliga et al., 2013) and the absence of apoptotic cell characteristics (Cobb et al., 1996; Hotchkiss et al., 2009) post NP/ i.d. application. Further studies are required to distinguish the type of cell death.

In conclusion, our work herein proposes that physically induced cell death by an immunization device co-delivering antigen into the skin can significantly improve the immunogenicity of vaccines by physical immune enhancement. This is a significant departure from the current adjuvant paradigm, generated by chemical or biological reagents added to antigens to enhance immune responses that are delivered into the body.

**Materials and Methods**

**Materials**

Seasonal human trivalent influenza vaccine (Fluvax® 2010) was manufactured by CSL Limited (Parkville, VIC, Australia), containing 15 μg haemagglutinin per strain per dose of purified, inactivated, detergent-disrupted split virion influenza virus with antigen of the following strains: A/California/7/2009, A/Wisconsin/15/2009 and B/Brisbane/60/2008.
Mice

Female 6–8 week old BALB/c or C57BL/6J mice were obtained from the Australian Research Council (Perth, WA). Anesthesia was performed by intraperitoneal (i.p.) injection of ketamine and xylazil anesthetic (50 mg/ kg and 10 mg/ kg; Troy laboratories Pty., Ltd., Smithfield, Australia) prior to immunizations. Mice were euthanized by cervical dislocation either immediately or at indicated time points after treatment. All animal experiments were conducted according to the University of Queensland Anatomical Biosciences Ethics Committee regulations.

Immunizations

The NP was fabricated from silicon using a deep reactive ion etching process at the University of Queensland (Jenkins et al., 2012). Coating solutions for NP were prepared as previously described (Chen et al., 2011; Chen et al., 2009), with coated projections examined by SEM and delivery efficiency verified (Fernando et al., 2012). The NP was applied using a spring-loaded applicator (Crichton et al., 2010) at ~2.3 ± 0.2 ms⁻¹ and applied for 2 min. NP and i.d. injections (31G, 20 μl) were administered into the ventral site of murine ears while i.m. injections were given into both hind legs into the caudal thigh muscle (29G; 33 μl).

Viability staining of tissue

To discriminate between live and dead cells, a mixture of metachromatic acridine orange (AO) and ethidium bromide (EB; both Fluka, Steinheim, Germany) in PBS was used (Franklin and Locker, 1981; Raju et al., 2006). AO and EB are routinely used to discriminate between live and dead cells in cell suspensions and tissue by intercalating with DNA and RNA (Baskic et al., 2006; Darzynkiewicz et al., 1992). Tissues were split at the dermis-cartilage-dermis junction with cartilage carefully removed prior to staining in AO (0.03 μg/ ml) and EB (0.1 μg/ ml). Positive controls for cell death were pre-treated with ice-cold methanol prior to staining (Skala et al., 2005). Confocal and Multiphoton Microscopy was used for visualization purposes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations used

| Abbreviation | Description     |
|--------------|-----------------|
| AO           | acridine orange |
| EB           | ethidium bromide|
AOEB  acridine orange and ethidium bromide
SC  stratum corneum
VE  viable epidermis
i.d  intradermal
i.m  intramuscular
APCs  antigen presenting cells
NP  Nanopatch
FP  flat patch
DAMPs  damage associated molecular patterns
CLSM  confocal laser scanning microscope
MPM  multiphoton microscopy
MIP  maximum intensity projection

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Figure 1. Comparison of the area of affect following Nanopatch application to skin and i.d. injection by hypodermic needle
(a) Size comparison of a NP next to a 31G needle. (b) SEM image of 31G needle tip over NP projections (inset). Cryo-SEM images of murine skin of (c) untreated ear, (d) following NP application, (e) 31G needle i.d. in situ, (f) skin after 31G needle removal following delivery of saline, (g) bleb formation following i.d. injection (20 μl saline). Arrows depict the needle insertion site. (h) Coomassie blue administered into mouse ears by either NP (left) or i.d. injection (right). (i) Affected skin surfaces by Coomassie Blue measured in Image J (n=18 applications each). Bar (a, g) = 1 mm, (b, e, f) = 200 μm, (c, d) 50 μm. i.d., intradermal; NP, Nanopatch. Bars represent means ± SD (*** p<0.001).
Figure 2. Cell viability in untreated, NP-treated and i.d. saline-injected murine skin
Acridine orange and ethidium bromide were used to differentiate between live (green) and
dead (magenta) cells. (a) Multiphoton microscopy images of stained murine; untreated, NP-
treated and i.d. injected with high magnification insets. (b) Representative side views of a)
with collagen (blue, second harmonic generation). (c) Quantification of cell viability per 16
mm²; skin incubated in methanol prior to staining as positive cell death control (Skala et al.,
2005). Original magnification 10x and 40x. White arrow depicts i.d. injection site. Results
are representative of three independent experiments with n=5–8 replicates and up to 3
measurements per replicate. Bar (a, upper panels) = 1 mm, (a, lower panels and b) = 200 μm.
MIP, Maximum Intensity Projection; SC, Stratum corneum; VE, viable epidermis; D,
dermis. Error bars represent means ± SD (***  p<0.0001).
Figure 3. Increased dose-sensitivity to antigen delivered by Nanopatch compared to i.d. injection
Endpoint titers of Fluvax® administered by NP or i.d. injections at various doses (0.1, 1, 5, 10, 20, 50 and 100 ng; i.m. 6000 ng) and analyzed by total IgG ELISA 21 days post immunization. Dose response curve on log scale depicting the dose differences between i.d., NP and i.m. NP, Nanopatch (▲), i.d., intradermal (●) and i.m., intramuscular (◆). Symbols are means of n=5 and error bars are mean + SD.
Figure 4. Co-localization of antigen with live and dead cells using Dylight® 755-labelled Fluvax® and viability stain

*In vivo* imaging of Fluorescent Dylight® 755-labelled Fluvax® (Rainbow colored) administered by NP or i.d. and dead cells (orange) 5 min post administration, detected using a MS FX Pro system overlaid with X-rays (gray scale). Images are representative of n=3 replicates/group. White arrows depict i.d. injection sites. NP, Nanopatch; i.d., intradermal; EB, ethidium bromide; FV, Fluvax®. Color-coded Dylight® 755 fluorescence represents intensity ranging from high (white-red) to low (blue-black).
Figure 5. Immune responses to Fluvax® following different levels of induced cell death
(a) Schematic of applied immunizations (b) Fluvax® (10 ng) was delivered by coated NPs, i.d. or a combination thereof (i.d. vaccine delivery followed by NP or FP) into murine ears (control group i.m. 6000 ng). (c) Various Fluvax® concentrations administered i.d. did not affect immunogenicity following i.d. immunization delivering a total of 5 ng Fluvax®. (d) CLSM and (e) MPM imaging of cell death post NP application using various densities (10,000; 21,000; 3x 21,000 projections cm$^{-2}$). (f) Quantification of cell death linked to Fluvax® immunogenicity (1 ng). All samples were analyzed as previously described. Bar
(d) = 1 mm, (e) = 50 μm. NP, Nanopatch; FP, flat patch; i.d., intradermal; i.m., intramuscular. Depicted are means with n=5 (ns: not significant $p>0.05$, * $p<0.05$, ** $p<0.01$, *** $p<0.001$, **** $p<0.0001$).
Figure 6. Schematic of i.d. injection and NP delivery alongside live-dead cells, vaccine and proposed DAMPs release and comparison of induced cell death with theoretical stresses in skin
(a) Vaccine (blue) administered either by i.d. injection or dissolving off coated NP projections with cell death (magenta). Vaccine was found predominantly around the i.d. insertion site within the VE and dermis, causing a bleb formation with minimal co-localization of DAMPs (magenta stars) with vaccine and viable cells (green). NP induces highly localized cell death in the VE and minimally in the dermis, with co-localization of DAMPs with antigen and live cells. (b) Comparison of measured cell death with theoretical stresses induced in skin by NP application. Bar (a, upper panel) = 200 μm, (a, lower panel) = 30 μm, (b) = 50 μm. i.d., intradermal; NP, Nanopatch; DAMPs, damage associated molecular patterns; SC, stratum corneum; VE, Viable Epidermis.