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A cell-based microarray to investigate combinatorial effects of microparticle-encapsulated adjuvants on dendritic cell activation

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Experimental vaccine adjuvants are being designed to target specific toll-like receptors (TLRs) alone or in combination, expressed by antigen presenting cells, notably dendritic cells (DCs). There is a need for high-content screening (HCS) platforms to explore how DC activation is affected by adjuvant combinations. Presented is a cell-based microarray approach, "immunoarray", exposing DCs to a large number of adjuvant combinations. Microparticles encapsulating TLR ligands are printed onto arrays in a range of doses for each ligand, in all possible dose combinations. Dendritic cells are then co-localized with physisorbed microparticles on the immunoarray, adherent to isolated islands surrounded by a nonfouling background, and DC activation is quantified. Delivery of individual TLR ligands was capable of eliciting high levels of specific DC activation markers. For example, either TLR9 ligand, CpG, or TLR3 ligand, poly I:C, was capable of inducing among the highest 10% expression levels of CD86. In contrast, MHC-II expression in response to TLR4 agonist MPLA was among the highest, whereas either MPLA or poly I:C, was capable of producing among the highest levels of CCR7 expression, as well as inflammatory cytokine IL-12. However, in order to produce robust responses across all activation markers, adjuvant combinations were required, and combinations were more represented among the high responders. The immunoarray also enables investigation of interactions between adjuvants, and each TLR ligand suggested antagonism to other ligands, for various markers. Altogether, this work demonstrates feasibility of the immunoarray platform to screen microparticle-encapsulated adjuvant combinations for the development of improved and personalized vaccines.

Introduction

Modern experimental vaccines are being designed with an emphasis on specific, tailored formulations to elicit more precise and potent immunological responses¹. A critical component of a vaccine is the adjuvant, a molecule or compound that potentiates the specific type and magnitude of an immune response to co-formulated antigens²⁻⁴. Various mechanisms underlying adjuvant activity have been uncovered, providing new strategies to optimize adjuvant formulations⁵. Recognition of pathogen-associated molecular patterns (PAMPs) by the immune system is achieved via various pathogen-recognition receptors (PRRs), notably toll-like receptors (TLRs)⁶. TLRs act as sensors for different damageassociated molecular pattern (DAMP) or "danger" signals generated by TLR-agonists such as double stranded RNA, DNA, and glycolipids present on the surfaces of many pathogens⁷. Microbial ligands bind to these receptors creating diverse immune responses that are the basis for multiple adjuvants currently in development8-10.

Numerous intracellular and surface bound TLRs have been identified on dendritic cells (DCs), which are cells of the innate immune system that act as the bridge for mounting an adaptive immune response against foreign antigens^{7, 11}. Dendritic cells are the most efficient antigen presenting cells (APCs), capable of orchestrating lymphocyte function and directing the immune response toward either immunity or tolerance¹²⁻¹⁵. Exploiting this potential, DCs have been manipulated both ex vivo and in vivo through controlled release schemes¹⁶ to treat a number of diseases such as cancer¹⁷⁻¹⁹, infection^{20, 21} and autoimmunity²² such as type-1 diabetes²³⁻²⁵. A next step in DC modulation involves simultaneously providing combinations of multiple different TLR ligands, which is capable of inducing synergistic increases in antigen-specific immune responses²⁶. Targeting multiple TLRs may recapitulate in a well-controlled manner, classic adjuvants formulated from attenuated or killed viruses or bacteria, providing an opportunity to more precisely direct DC function^{27, 28}. However, while this potential has been recognized, to date there is no systematic methodology to explore dose-dependent interplay of combined TLR stimulation in DCs. Such an approach would facilitate the development of combinatorial adjuvants for tailored immune responses.

One strategy to develop immunotherapeutic treatments involves using biomaterials as modulators of DCs²⁹⁻³⁵. Polymeric,

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particle-based approaches have been developed that encapsulate combinations of antigen, adjuvant, chemokines, and other immunomodulating molecules for delivery to DCs via phagocytosis and controlled release³⁵⁻³⁷. However, when considering combinatorial adjuvant approaches, the number of possible TLR ligand combinations is large, and identifying improved formulations using current immunological methods is challenging due to limitations in cell sourcing, as well as cost, labor, and time, in particular for personalized/precision medicine applications³⁸. To address this, we developed a cellbased microarray to screen combinatorial libraries of adjuvants formulated with biodegradable microparticles (MPs). The adjuvants monophosphoryl lipid A (MPLA, TLR4 ligand)³⁹, CpG (TLR9 ligand)⁴⁰, and polyinosinic:polycytidylic acid (poly I:C, TLR3 ligand)³⁹ were encapsulated in poly(lactic-coglycolic acid) (PLGA) MPs, printed on a DC-based microarray41 in 216 unique combinations, and cell surface markers of activation and cytokine production were quantified. Improved DC responses were indicated for particular dosing combinations of TLR ligands encapsulated in PLGA MPs. This immunoarray is hereby demonstrated to provide a robust, miniaturized platform to characterize tailored adjuvant formulations.

Experimental

Materials

A 50:50 polymer composition of poly(d,l lactide-co-glycolide) (PLGA) with inherent viscosity 0.55-0.75 dL/g in hexafluoroisopropanol, HFIP (Lactel, AL, USA) was used to generate microparticles. Microparticles (MPs) were formed using a standard water-oil-water solvent evaporation technique. Briefly, the PLGA polymer was dissolved in methylene chloride at 20% concentration. To fabricate fluorescent particles, either rhodamine (RHOD) (100 µL of 5 mg/mL in PBS) (Sigma-Aldrich), fluorescein isothiocyanate (FITC) (100 μL of 5 mg/mL in PBS) (Sigma Aldrich), or 9anthracenecarboxylic acid (ACA) (100 µL of 5 mg/mL in PBS) (Sigma Aldrich) were encapsulated. For adjuvant-loaded particle fabrication, LPS (1 mg/ml Fisher Scientific), CpG ODN (0.1 mg/ml InvivoGen), polyinosinic:polycytidylic acid (poly I:C; 1 mg/ml Sigma-Aldrich), or monophosphoryl lipid A (1 mg/ml MPLA) was encapsulated in the primary water phase. The primary emulsion was generated by emulsifying 100 µL of adjuvants or dye with 1 mL of PLGA solution dissolved in methylene chloride (Fisher Scientific) using a tissue-miser homogenizer. The primary emulsion was then added to 10 mL of a 5% poly-vinyl alcohol (PVA) (MW ~ 100,000 g/mol) (Fisher Scientific) solution in PBS and homogenized to form the secondary emulsion. Afterward, the secondary emulsion was added to 100 mL of 0.5% PVA solution and the particles were agitated using a magnetic stirrer overnight to evaporate residual methylene chloride. The particles were subsequently washed, flash-frozen, and stored at -20°C until used.

Dendritic Cell Culture

Dendritic cells were isolated from the bone marrow of 7 week old C57Bl6/j mice using a 10 day protocol²⁹. Briefly, bone marrow was isolated from the femur and tibia. Red blood cells were lysed by ACK lysing buffer (Whittaker) and the isolated precursor cells were incubated with DC-media consisting of 20 ng/mL of GM-CSF (R&D Systems), DMEM/F12 (1:1) with Lglutamine (Cellgro, Herndon, VA) and 10% fetal bovine serum (Bio-Whittaker), 1% sodium pyruvate (Lonza, Walkersville, MD) and 1% non-essential amino acid (Lonza, Walkersville, MD) for 2 days in a T-75 flask. The floating cells were collected after 48 h and re-seeded with fresh DC-media in a 6well low-attachment plate (Corning Inc., NY) for 6 days. After 6 days of culture the cells were re-suspended in fresh media and seeded onto tissue-culture treated 6-well plates (Corning, Inc., NY) for 2 days. After 10 total days of culture DCs were lifted using 10 mM solution of Na₂EDTA (Fisher Scientific) in PBS. Dendritic cells thus isolated were then tested via flow cytometry for immaturity (MHC-II+ < 6% and CD86+ < 6%), purity (CD11c+ > 90%) and viability (Trypan Blue > 99%).

Immunoarray fabrication

Microarrays were manufactured as described previously⁴¹. Briefly, glass coverslips were cleaned in an oxygen plasma etcher (Terra Universal, Fullerton, CA). Arrays of (3-Aminopropyl) trimethoxysilane (NH₂-terminated silane) (Sigma-Aldrich, St. Louis, MO) were printed on clean coverslips using a Calligrapher Miniarrayer printer (Bio-Rad, Hercules, CA) with a pin diameter of 400 µm. The silane printed coverslips were then coated with 200 Å of titanium (Ti; 99.995% pure) and 200 Å of gold (Au; 99.999% pure) (Williams Advanced Materials, Buffalo, NY). Following coating, gold-coated arrays were sonicated to remove gold from the amine spots, exposing NH2-terminated silane islands. The coverslips were incubated with 0.1 M, methyl-terminated alkanethiol (CH₃(CH₂)₁₁SH) (Sigma) for 30 min. Substrates were incubated in 10% Pluronic® F-127 (BASF Corporation, USA) for 3 h to create a non-fouling surface around the adhesive amine islands (Fig. 1b). The immunoarrays were generated with 648 cell adhesive islands to provide 216 different conditions in triplicate. Fabricated MP solutions were printed onto cell adhesive islands using standard contact pin miniarraying (BioRad) equipment Following MP printing, the microarrays were desiccated for 30 min then stored at 4°C. For adjuvant optimization experiments, particles encapsulated with CpG, poly I:C, or MPLA were used. Six dilutions of each MP formulation were generated and printed on the immunoarrays in all possible combinations, thus providing 216 different conditions. Dried MP arrays were rinsed and rehydrated for 30 minutes with PBS before seeding immature DCs. Following seeding, the arrays were rinsed with PBS to remove nonadherent cells. Fresh DC-media was added to the immunoarrays and adherent DCs were cultured for 24 h before performing immunocytochemistry.

Scanning Electron Microscope Imaging

Microparticle morphology was characterized by scanning electron microscope (FEG-SEM JEOL JSM - 6335F, Major

Analytical Instrumentation Center, University of Florida). Particles were dried for 16 h at room temperature, then coated with 5-10 nm thickness of gold and imaged at magnifications ranging from 30X to 60,000X. Dendritic cells cultured on the immunoarrays were fixed and imaged using scanning electron microscope by ICBR core electron microscopy facility at University of Florida, Gainesville.

Microparticle to Dendritic Cell Ratio Optimization

Prescribed numbers of particles can be printed by varying the concentration of MP solutions (in diH₂O) in the source plate using the following experimentally determined relationship:

Desired Number of MPs / spot = 21.3 x (Source Plate Concentration) -0.1

where: "source plate concentration" is the concentration of MP solutions in the wells of a 384-well plate from which the particles were lifted by the miniarrayer pin and printed on the immunoarray. Rhodamine encapsulated MPs (300, 600, 1500, 3000 and 9000 MPs / spot) were printed onto different islands. Dendritic cells were cultured on the vaccine array chip and cell number was quantified via nuclear staining (Hoechst 34580, Invitrogen) and fluorescent microscopy. The number of DCs was found to be 300±33 per spot, thus providing MP:DC ratios of 1, 2, 5, 10 and 30. The DCs were then cultured on the vaccine array chip for 24 h upon which the cells were lifted and transferred to 96-well plates. A standard curve for rhodamineloaded MPs was generated and the number of particles internalized by DCs was quantified for each condition. Particleprinted arrays without DCs was used as a control to analyze the lifting of particles in the absence of DCs. It was observed that Trypsin/EDTA did not dislodge the physisorbed particles.

Immunocytochemistry

Following 24 h incubation on the immunoarrays, DCs were fixed with 4% paraformaldehyde (USB Corporation). Cells were then fluorescently stained either for surface markers CCR7 (Santa Cruz Biotechnology), CD86 (BD Biosciences), MHC-II (BD Biosciences), or intracellular cytokines IL-10 (BD Biosciences), or IL-12p40 (BD Biosciences) as previously reported⁴². Intracellular staining for cytokines IL-10 and IL-12p40 was performed by first incubating seeded immunoarrays with monensin (0.7 µL/ml) for the final 8 h of culture in order to block protein secretion. The arrays were then incubated with 4% paraformaldehyde followed by 0.01% triton X-100 (Fisher Scientific). Arrays were next washed and incubated with a blocking solution consisting of 1% goat serum along with 0.01% triton X-100 in PBS for 40 min to block the background before incubation with primary antibodies. Afterward, solutions of biotinylated secondary antibodies (Invitrogen) were incubated, followed by streptavidin-cross-linked alkaline phosphatase, and lastly the precipitating fluorescent substrate, ELF97 (Invitrogen). Nuclei were stained with Hoechst 34580 dye. Immunoarrays were then mounted and imaged.

Inter-Spot Cross-Talk

In order to investigate cross-talk, or paracrine signaling between spots, center to center spot distances of 1,000 or 1,200

µm were investigated. Poly I:C loaded MPs were printed in dilutions to provide MP:DC ratio of 10, 5, 2.5, 1 and 0 (no MPs). Randomized array configurations generated by MATLAB were used. Dendritic cells were cultured for 24 h and stained for intracellular IL-12p40. Production levels of IL-12p40 were

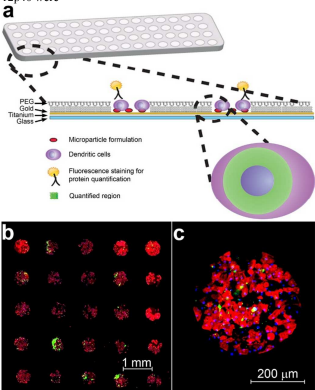


Fig 1. Schematic of immunoarray design and representative images of a previously described immunoarray of dendritic cells (DCs) cultured with co-localized microparticles⁴¹. **(a)** PLGA microparticles encapsulated with adjuvants are printed on isolated microarray islands. Dendritic cells are co-cultured with particles on the immunoarrays and characterized via immunocytochemistry. Arrays are imaged and quantified for expression of proteins that define DC activation and function. **(b)** Fluorescence microscopy mosaic image of a 5x5 section of a microarray seeded with murine bone-marrow derived dendritic cells illustrating fidelity of cell adhesion to isolated islands of adjuvant-encapsulated microparticles. Actin filaments stained using rhodamine phalloidin (red), FITC encapsulated particles demonstrating colocalization of microparticles (green) with DCs, and nuclear staining with Hoechs 34580 (blue). **(c)** Shown is a detail of a single island with DCs co-localized with microparticles.

quantified using Axiovision as described below.

Data Acquisition - Optical Microscopy and Image Processing

Following immunocytochemistry, mounted immunoarrays were imaged using an Axiovert 200M Carl Zeiss inverted fluorescence microscope. The images were collected at 12-bit resolution (4096 different colors / pixel) with a filter sensitive for DAPI and a long pass filter (EX 365/12 EM 397) appropriate for the ELF 97 fluorescent precipitate. Image analysis was performed using

AxioVision software. Briefly, nuclei were selected using DAPI staining, and a 4 µm rings distance of concentric circles were drawn around the nucleus (**Figure 1a**). Next the mean relative

fluorescence intensity of the area between the concentric circles were determined using the AxioVision software for each cell and utilized to determine the expression of different proteins. Cell nuclei were identified by DAPI staining and DC numbers per island were quantified by Carl Zeiss Axiovision automatic count object routine.

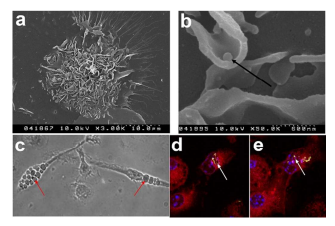


Fig 2. Characterization of microparticle phagocytosis by dendritic cells (DCs) on immunoarrays. (a) Scanning electron microscope micrograph of a DC on an island surrounded by particles (3,000x). (b) Scanning electron microscope micrograph indicating DCs can lift physisorbed microparticles (arrow) from the array surface (50,000x). (c) Phase contrast image of DCs with phagocytosed particles localized in the cytosol. Particle internalization is highlighted (arrows). (d, e) Fluorescence confocal microscopy images of DCs phagocytosing microparticles. Confocal microscopy showing the co-localization of dye-loaded particles (arrows) with the cytosol (red). Multiple z-stacked images were acquired to confirm the internalization of the microparticles (data not shown). Actin filaments stained using rhodamine-conjugated phalloidin (red), FITC encapsulated particles demonstrating phagocytosis (yellow-green), and nuclear staining with Hoechst 34580 (blue).

Statistical Analyses

Overall statistical significance, least mean squares values and standard error were obtained using two-way ANOVA via Systat (Version 12, Systat Software, Inc., San Jose, CA), with independent variables being the individual experimental run identifier and the particle condition identifier. Pair-wise comparisons were made by Tukey's post-hoc analysis. A p-value of p < 0.05 was considered significant. Data was plotted using Sigmaplot (Version 10, Systat Software, Inc., San Jose, CA).

Results and discussion

Dendritic cell characterization on the immunoarray

The immunoarray developed here represents a high-content, cell-based array approach to screen combinatorial adjuvant microparticle formulations. PLGA MPs encapsulating three TLR ligands were printed onto solid substrates with discrete islands in hundreds of dosing combinations. A non-fouling background surrounding these cell-adhesive islands enabled dendritic cells (DCs) to be co-localized as isolated samples with unique particle/adjuvant formulations⁴¹. Microparticles were printed on immunoarrays in a reproducible and quantitative manner, with minimal cross-contamination⁴¹. Using these

printing methods we could obtain homogenous deposition with as few as 16±2 particles per spot. This miniaturized platform can decrease the use of reagents, time, and cost while providing the capability of screening greater than 30-fold more immunomodulatory molecules, based on the number of cells required, compared to traditional methods. Further, since the only complimentary equipment required is a fluorescent microscope, the immunoarray platform can easily be adopted into many biomedical laboratories.

Expression of cell surface markers and cytokine production upon interaction with particle formulations is quantified using standard immunocytochemistry and fluorescent microscopy (Fig 1a). Cell attachment is tightly confined to the particle-coated silane islands as confirmed via actin (shown in red) and nuclei (shown in blue) staining of adherent DCs (Fig 1b, 1c). Dendritic cell viability was assessed using 7-AAD on the immunoarrays and found to be 99% viable. Scanning electron microscope (SEM) micrographs revealed the interaction between DCs, microparticles on immunoarrays (Fig 2).

Cell attachment was seen to occur with the formation of lamellipodia like structures, numerous dendritic processes and membrane ruffling (Fig 2a). The ability to lift physisorbed MPs from the array surface was evident (Fig 2b), with apparent internalization in the cytosol (Fig 2c). Fluorescence confocal microscopy shows microparticles were localized internally, adjacent to the nucleus (Fig 2d).

Particle uptake

In order to further characterize DC-MP interactions on the array, particle uptake was quantified (**Fig 3a**). Immunoarrays were generated, varying the number of fluorescent dye-loaded particles per spot. DCs were then incubated on the immunoarrays for 24 h followed by nuclear staining. The time point of 24 h was chosen both for convenience as an incubation time for subsequent experiments, and because others have suggested that DC phagocytosis can saturate at this time 43 . The percentage of particles phagocytosed plotted against the particle to DC ratio was found to exhibit a linear relationship. Approximately 90% of the physisorbed MPs on the array were taken up by DCs after 24 h incubation at a MP:DC ratio of 10 ± 2 . In subsequent studies, this ratio was used to provide a high MP uptake level while limiting the number of non-phagocytosed particles.

Island spacing optimization

A potential design constraint to miniaturized biological array platforms is the possibility of interactions between neighboring samples. Dendritic cells in particular have been shown to induce functional modulations in neighboring cells via paracrine signaling^{30, 44, 45}. While this phenomenon is generally accepted, the spatio-temporal modulations are not well characterized in culture. We explored the functional range of paracrine signaling between DCs by comparing distances in which a dose-dependent response was achievable (**Fig 3b**). Doses were obtained by printing different numbers of poly I:C

MPs to achieve a range of MP:DC ratios. Poly I:C is a potent inducer of IL-12 production in DCs⁴⁶. We therefore quantified IL-12p40, an active subunit of IL-12, for platform optimization studies. The 1,000 μm spacing (**Fig 3b, left panel**) does not provide the high fidelity dose response seen with the 1,200 mm spacing (i.e., all doses are significant from each other), which suggests negligible cross-talk between neighboring islands at the larger inter-island distance (**Fig 3b, right panel**). In subsequent studies, 1,200 μm spacing

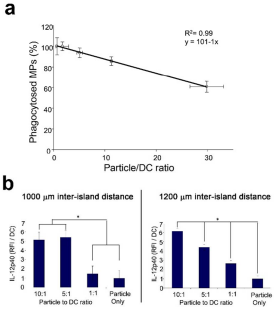


Fig 3. Quantification and optimization of microparticle (MP) to dendritic cell (DC) ratio and immunoarray island spacing. (a) Dendritic cells were cultured with rhodamine-encapsulated microparticles at varying ratios of MPs:DCs and the percentage of MPs phagocytosed was quantified. Dendritic cells were shown to phagocytose ~90% of the printed particles within the first 24 h when the particle to DC ratio was maintained at 10:1. A linear relationship was observed between the percent of particles phagocytosed and the MP:DC ratio. (b) Inter-island spacing was investigated to limit paracrine signaling. Poly I:C encapsulated MPs were printed with a range of MP:DC ratios. IL-12p40 production was quantified between immunoarrays printed in randomized configurations at either 1,000 μm or 1,200 μm center-to-center inter-island distances. Production of IL-12p40 at the 1,000 μm spacing (b, left panel) does not provide the dose response fidelity seen with the 1,200 μm spacing (b, right panel) suggesting negligible cross-talk between neighboring islands at the larger inter-island distance. (*:p<0.05)

was incorporated in the immunoarrays limit paracrine signaling between neighboring groups. The ability to elicit dose dependent cellular responses on the immunoarray is central to the platform validation. Having established island spacing, arrays were next fabricated with MPs loaded with a range of loading doses of LPS, an activating signal. Equal amounts of MPs were then printed on each island and DCs were cultured on the immunoarrays for 16 h before quantifying activation. To determine DC activation, several markers were explored including CD86 (co-stimulatory signal providing protein), MHC-II (antigen expressing molecule), IL-12p40 (proinflammatory cytokine), and IL-10 (anti-inflammatory molecule). Utilizing this approach confirmed that an increase in the MP loading dose of LPS increased production of all four

markers investigated (**Fig 4a-d**). Further, no significant increase in activation was observed from DCs co-localized with unloaded blank particles compared to the no particle control, confirming that these PLGA MPs are not stimulatory on their own.

Combination adjuvant effects on dendritic cells

Increasing adjuvant quantity has been shown to increase DC activation^{45, 47} until saturated levels are reached²⁶. The effect of

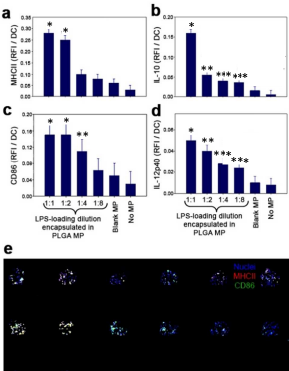


Fig 4. Dose-dependent dendritic cell activation on immunoarrays. Microparticles were fabricated with a range of LPS loading doses and printed in equal numbers onto the immunoarrays. DCs were then cultured on the platform for 16 h before fixation and immunocytochemistry was performed. (a) MHC-II expression was upregulated 6-fold at the highest dose of LPS compared to the unloaded MP control. * significant from 1:4, 1:8, Blank MP and No MP. (b) IL-10 production was shown to increase 10-fold for the highest LPS dose compared to unloaded MPs. * significant from all other groups; ** significant from 1:1, 1:8, Blank MP and No MP; *** significant from 1:1, Blank MP and No MP (c) Expression of the co-stimulatory signal CD86 revealed a dose-dependent response to LPS that reached a maximum at the second highest dose, a 3-fold increase compared to the unloaded control. * significant from 1:4, 1:8, Blank MP and No MP; ** significant from Blank MP and No MP. (d) Pro-inflammatory cytokine IL-12 showed increased production for each dose of LPS. * significant from 1:4, 1:8, Blank MP and No MP; ** significant from 1:8, Blank MP and No MP; *** significant from 1:1, Blank MP and No MP. No differences were found between the unloaded blank MPs and the no MP control for all markers.

varying combined adjuvant formulations on DC activation states is less explored but more intriguing, particularly in the pursuit of tunable immunotherapeutics. It is known that combining adjuvants may lead to differential activation of DCs, a requisite for successful induction of cellular and humoral responses^{26, 48, 49}. For instance, Th1 immune responses are

500 μm

stimulated through TLR3, TLR 4, TLR 7, TLR 8, and TLR 9 whereas TLR5 and $\,$

TLR2 together with TLR1 or TLR6 supports a Th2 immune response ⁵⁰⁻⁵². Utilizing the immunoarray platform facilitates investigation of the interplay between TLR signaling on DCs, a critical component to controlling the type and magnitude of antigen-specific immune responses. Microparticles encapsulated with MPLA, CpG, and poly I:C were printed on arrays consisting of 648 cell adhesive islands, testing six dilutions of each adjuvant

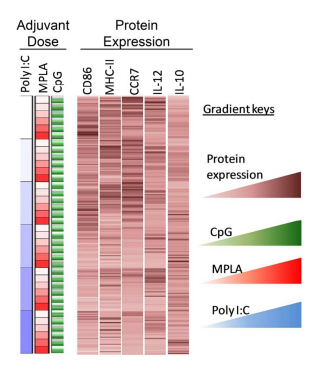


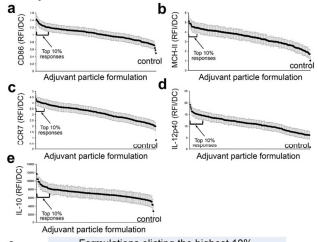
Fig 5. Protein expression and cytokine production from dendritic cells (DCs) upon exposure to different combinations of microparticle-encapsulated adjuvants. A color-coded map connecting external stimulation (in rows) to protein expression from DCs (columns). Columns display relative fluorescent intensity of each marker measured. Each row corresponds to a particular formulation of adjuvant combination. Average RFI per cell were measured across three array experiments per marker, averaged across experiments, and normalized across separate microarrays to each array's no particle control. Gradients are represented by lighter to darker color for CpG (green), MPLA (red), poly I:C (blue) and protein expression (brown).

in all possible combinations resulting in 216 unique combinations (i.e., 6³=216, and a replicate number of n=3 on each array). DC activation was quantified via the markers CD86, MHC-II, IL-12p40, IL-10, and also CCR7 (chemokine receptor for lymphatic homing) (Fig 5). A pro-inflammatory response is characterized by high expression of CD86, MHC-II, IL-12p40, CCR7 and low expression of IL-10. In all adjuvant combinations, it was observed that expression of MHC-II, CD86, CCR7, IL-12p40, and IL-10 were significantly different from the negative control condition of no-particles (Fig 6 a-e). This confirms that adjuvant loaded particles have a measureable effect on DC activation using this approach. Furthermore, it was observed that the condition consisting of CpG:MPLA:Poly I:C at a ratio of 1c:2m:2p (c = CpG, m = MPLA, p = Poly I:C) generated the highest pro-inflammatory state of DCs on the immunoarray, as indicated by high expression of MHC-II, CD86, IL-12, and CCR7 and low expression of IL-10 (Fig 6 f).

Adjuvant interactions

To gain insight into the interactions between the three MP encapsulated adjuvants, analysis of the combinatorial effects was performed. The results are presented by arranging data in blocks of dosing categories along the x-axis (Fig 7a – darker shades represent larger doses). Thus, this data is arranged in

large (blue) blocks in order of increasing poly I:C dose; within this superset



Formulations elicting the highest 10% of responses for each marker				
CD86	MHC-II	IL-12p40	CCR7	IL-10
5c5m0p	2c3m5p	3c5m4p	4c5m1p	5c5m5p
5c5m2p	2c3m1p	3c4m3p	4c0m2p	5c5m4p
5c3m0p	2c2m4p	3c2m0p	3c2m2p	5c5m3p
5c2m0p	2c0m2p	3c1m0p	2c2m0p	5c3m5p
5c2m2p	1c5m2p	1c5m4p	2c0m2p	5c3m3p
5c1m0p	1c5m1p	1c5m3p	1c5m1p	5c1m2p
5c0m2p	1c3m5p	1c5m1p	1c4m0p	5c0m4p
5c0m0p	1c2m2p	1c5m0p	1c2m2p	5c0m3p
4c5m0p	1c2m1p	1c4m4p	1c2m0p	4c4m5p
4c3m2p	1c2m0p	1c2m4p	1c0m2p	4c3m3p
4c2m0p	1c1m1p	1c2m2p	0c5m1p	4c3m2p
4c1m0p	1c0m5p	1c2m0p	0c4m2p	4c2m5p
4c0m0p	1c0m4p	1c1m5p	0c3m0p	4c1m5p
3c5m2p	1c0m2p	1c1m4p	0c2m0p	4c1m3p
3c4m2p	1c0m1p	1c1m2p	0c1m2p	4c0m5p
3c2m0p	0c4m2p	1c1m1p	0c1m1p	4c0m4p
3c0m2p	0c4m0p	1c0m1p	0c1m0p	4c0m2p
3c0m0p	0c1m1p	0c5m0p	0c0m2p	3c3m4p
2c4m2p		0c2m4p		2c0m3p
1c2m2p		0c0m4p		1c1m3p
0c0m5p				1c0m4p

Fig 6. Combinations of microparticle-loaded adjuvants up-regulate dendritic cell activation markers to varying degrees. Microparticle formulations were organized left to right by the level of expression/production of markers from highest to lowest (mean +/- standard error) for a) CD86, b) MHC-II c) CCR7 d) IL-12 e) IL-10, where the x-axis represents each of the 216 unique formulations. Brackets identify formulations eliciting the highest 10% responses, and are listed in **(f)**. Highlighted are adjuvant combinations with particularly high expression of more than one marker, with the condition consisting of CpG:MPLA:Poly I:C at a ratio of 1c:2m:2p generating the highest pro-inflammatory state of DCs indicated by MHC-II^{HI}CD86^{HI}IL-12^{HI}CCR7^{HI}IL-10^{LI}. Key: c – CpG; m – MPLA; p – poly I:C; 0 = No particle; 1 = 62 particles; 2 – 125 particles; 3 – 250 particles; 5 – 1,000 particles.

are smaller (red) blocks in order of increasing MPLA dose; and within this dataset is individual CpG doses (green) in increasing order. This hierarchical data organization was rearranged in all possible configurations to identify potential trends (data not shown).

Linear regressions were generated for every adjuvant dosing range in the presence of a fixed amount of the other two adjuvants for all possible configurations in order to elucidate

the effect of each adjuvant on surface expression of DC activation marker, costimulatory molecule CD86 (**Fig 7b-g**).

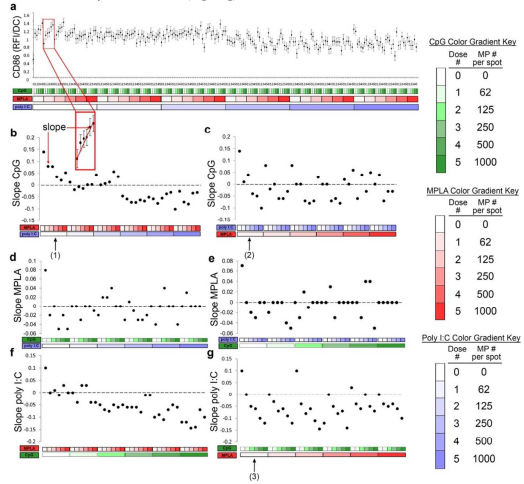


Fig 7. Analysis of dendritic cell (DC) CD86 expression (RFI) in response to combinatorial adjuvant-loaded microparticles (MPs). (a) CD86 expression (RFI) per DC is plotted against varying amounts of three MP-encapsulated adjuvants (CpG, MPLA and poly I:C; n = 9, with 3 replicates per array on 3 separate arrays). Data is arranged by grouping increasing particle numbers of CpG (green), MPLA (red), and lastly poly I:C (blue) with doses of 0, 62, 125, 250, 500 and 1,000 particles per spot, represented by increased shading from lighter to darker colors. The x-axis represents different adjuvant concentrations, and the y-axis represents the expression of CD86 per DC, expressed in RFI. (b, c) Linear fits were calculated for each set of CpG dosing range (lowest to highest number of particles) for every combination of MPLA and poly I:C doses. An example showing the mapping of a single slope is illustrated in the inset. The slope of each linear fit was then calculated and plotted as a function of MPLA and poly I:C doses. Dotted lines represent a slope of zero. (b, arrow # 1) In the absence of poly I:C, the slope of CD86 expression demonstrates a positive response to CpG MPs, and introducing MPLA (red) dampens this effect as seen by the decreasing slope values plotted on the y-axis, with increasing MPLA dose. (b) Introducing poly I:C blocks the dampening effect MPLA has on CpG at lower doses and reverses the positive CD86 response to CpG, as evidenced by the negative slope values, at high poly I:C doses. (c, arrow # 2) In the absence of MPLA, poly I:C decreases CD86 responsiveness to CpG. (c) As MPLA is introduced, it mitigates some of the dampening effect of poly I:C on CpG dependent CD86 expression. (d, e) The adjuvant formulations were then reorganized for every combination dose of CpG and poly I:C. Linear fits were created for the slope of the MPLA response associated with the specific dose of CpG and poly I:C. appears to mitigate this response as most combination doses are near or equal to zero with no apparent trends

The slope of each linear fit was calculated, where a positive slope indicates increasing surface expression and a negative slope indicates decreasing expression when exposed to the adjuvant named on the y-axis, and when in combination with fixed amount of the other adjuvants. Using this method, it is thus possible to interpret interaction effects between two or three adjuvants used in combination by comparing how the

dose responses are altered upon introduction of additional adjuvants.

Several dominant interactions emerged from the linear regression analysis of the CD86 data. First, exposure to CpG MPs alone displayed the greatest capacity to upregulate expression of CD86 in DCs (**Fig 7b,d,f** leftmost data points). Both the addition of MPLA or poly I:C dampens responsiveness to CpG, but while higher dose MPLA can block

the response to CpG, higher dose poly I:C can actually reverse it. Additionally, while MPLA had a modest effect on CD86 expression by itself, in combination with CpG or poly I:C this effect was mitigated (**Fig 7d,e**). Lastly, the addition of MPLA or CpG dampens responsiveness to poly I:C, and higher CpG doses can reverse the poly I:C response, as previously reported⁵³.

1a

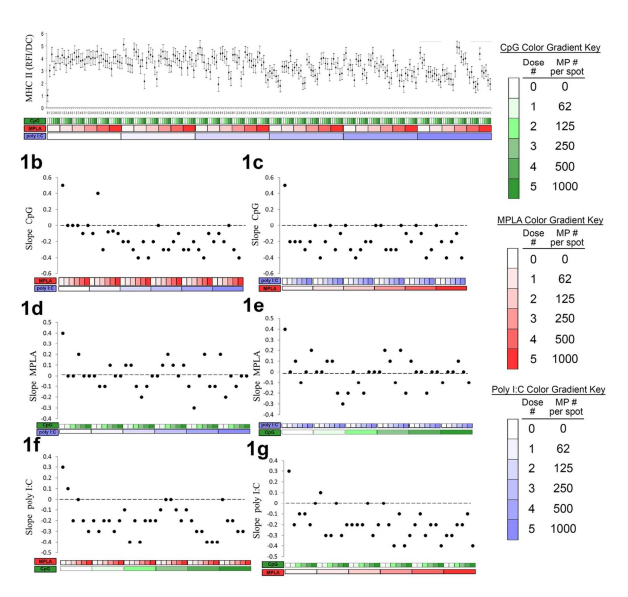


Fig 8: Analysis of dendritic cell (DC) MHC-II expression (RFI) in response to combinatorial adjuvant-loaded microparticles (MPs). (a) MHC-II expression (RFI) per cell is plotted against varying amounts of three MP-encapsulated adjuvants (CpG, MPLA and poly I:C; n = 9, with 3 replicates per array on 3 separate arrays). Data is arranged by grouping increasing particle numbers of CpG (green), MPLA (red), and lastly poly I:C (blue) with doses of 0, 62, 125, 250, 500 and 1,000 particles per spot, represented by increased shading from lighter to darker colors. (b, c) Linear fits were calculated for each set of CpG dosing range (lowest to highest number of particles) for every combination of MPLA (red) and poly I:C (blue) doses. The slope of each linear fit was then calculated and plotted as a function of MPLA and poly I:C doses. Dotted lines represent a slope of zero. (d, e) The adjuvant formulations were then reorganized for every combination dose of CpG and poly I:C. Linear fits were created for the slope of the MPLA response associated with the specific dose of CpG and poly I:C. The slope of each linear fit was then calculated and plotted as a function of CpG (green) and poly I:C (blue) doses. Dotted lines represent a slope of zero. (f, g) The adjuvant formulations were then reorganized for every combination dose of CpG and MPLA. Linear fits were created for the slope of the poly I:C response associated with the specific dose of CpG and MPLA. Linear fits were calculated for each set of poly I:C dosing range The slope of each linear fit was then calculated and plotted as a function of CpG (green) and MPLA (red) doses. Dotted lines represent a slope of zero.

Expression of MHC class II molecules in response to MP-encapsulated adjuvants was quantified (**Fig 8**). MHC-II is a critical component of DC activation, allowing presentation of foreign peptides to antigen-specific T-cells⁵⁴. Each MP encapsulated adjuvant individually increased MHC-II expression, with similar effects (**Fig 8b,d,f** leftmost points). The dose dependent response to CpG with the addition of either

MPLA or poly I:C decreased MHC-II surface expression (**Fig 8b,c**). On the other hand, dose dependent responses to MPLA were non-linearly modified by the addition of CpG or poly I:C (**Fig 8d,e**). Lastly, dose dependent responses to poly I:C with the addition of either MPLA or CpG decreased MHC-II expression (**Fig 8f,g**).

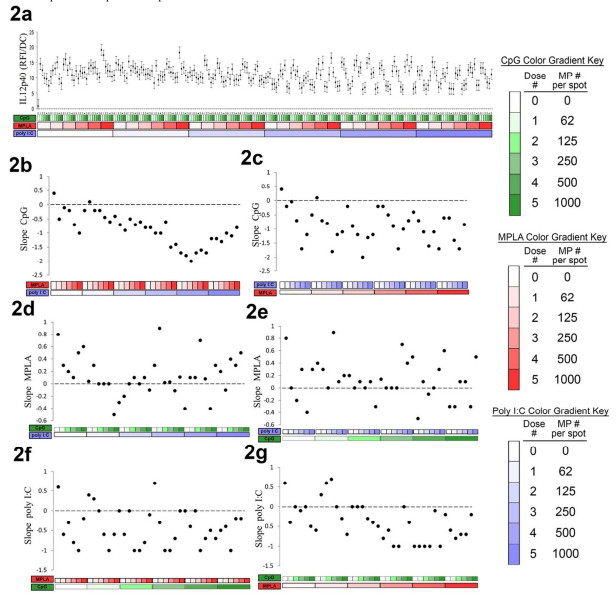


Fig 9: Analysis of dendritic cell (DC) IL-12 production in response to combinatorial adjuvant-loaded microparticles (MPs). (a) IL-12 production per cell is plotted against varying amounts of three MP-encapsulated adjuvants (CpG, MPLA and poly I:C; n = 9, with 3 replicates per array on 3 separate arrays). Data is arranged by grouping increasing particle numbers of CpG (green), MPLA (red), and lastly poly I:C (blue) with doses of 0, 62, 125, 250, 500 and 1,000 particles per spot, represented by increased shading from lighter to darker colors. (b, c) Linear fits were calculated for each set of CpG dosing range (lowest to highest number of particles) for every combination of MPLA (red) and poly I:C (blue) doses. The slope of each linear fit was then calculated and plotted as a function of MPLA and poly I:C doses. Dotted lines represent a slope of zero. (d, e) The adjuvant formulations were then reorganized for every combination dose of CpG and poly I:C. Linear fits were created for the slope of the MPLA response associated with the specific dose of CpG and MPLA. Linear fits were then reorganized for every combination dose of CpG and MPLA. Linear fits were created for the slope of the poly I:C response associated with the specific dose of CpG and MPLA. Linear fits were calculated for each set of poly I:C dosing range The slope of each linear fit was then calculated and plotted as a function of CpG (green) and MPLA (red) doses. Dotted lines represent a slope of zero.

Secretion of the IL-12, a dominant Th1-promoting cytokine⁵⁵, was also investigated (**Fig 9**). Each adjuvant individually upregulated IL-12 production, with MPLA providing a somewhat larger effect (**Fig 9b,d,f** leftmost points). Introduction of either MPLA or poly I:C was capable of reversing the dose dependent response to CpG (**Fig 9b,c**). In contrast, the addition of either CpG or poly I:C had a tendency

to maintain positive dose dependent responses to MPLA (**Fig 9d,e**). Lastly, introduction of either MPLA or CpG was capable of reversing the dose dependent responses to poly I:C (**Fig 9f, g**). Overall, these observations suggest significant interactions between MPLA (TLR4) and either poly I:C (TLR3) or CpG (TLR9) on DC production of IL-12p40, which corroborates previous

findings^{27, 39, 56}

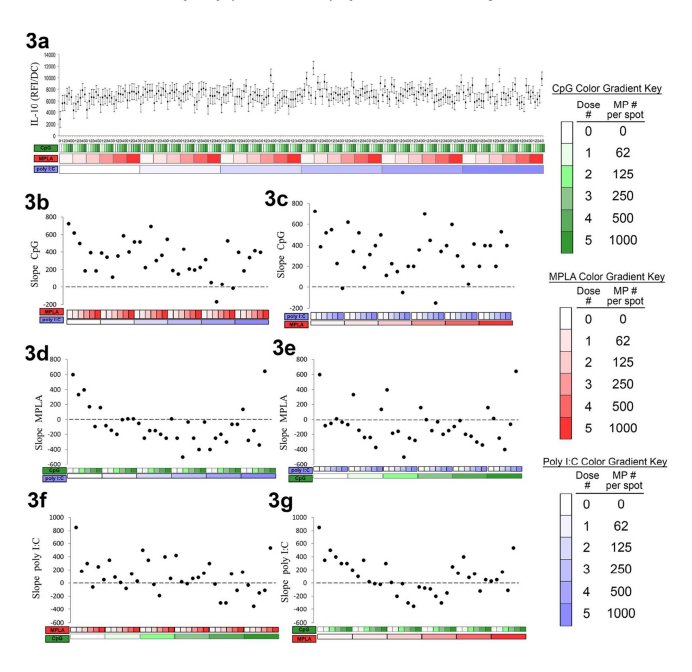


Fig 10: Analysis of dendritic cell (DC) IL-10 production in response to combinatorial adjuvant-loaded microparticles (MPs). (a) IL-10 production per cell is plotted against varying amounts of three MP-encapsulated adjuvants (CpG, MPLA and poly I:C; n = 9, with 3 replicates per array on 3 separate arrays). Data is arranged by grouping increasing particle numbers of CpG (green), MPLA (red), and lastly poly I:C (blue) with doses of 0, 62, 125, 250, 500 and 1,000 particles per spot, represented by increased shading from lighter to darker colors. (b, c) Linear fits were calculated for each set of CpG dosing range (lowest to highest number of particles) for every combination of MPLA (red) and poly I:C (blue) doses. The slope of each linear fit was then calculated and plotted as a function of MPLA and poly I:C doses. Dotted lines represent a slope of zero. (d, e) The adjuvant formulations

were then reorganized for every combination dose of CpG and poly I:C. Linear fits were created for the slope of the MPLA response associated with the specific dose of CpG and poly I:C. The slope of each linear fit was then calculated and plotted as a function of CpG (green) and poly I:C (blue) doses. Dotted lines represent a slope of zero. (f, g) The adjuvant formulations were then reorganized for every combination dose of CpG and MPLA. Linear fits were created for the slope of the poly I:C response associated with the specific dose of CpG and MPLA. Linear fits were calculated for each set of poly I:C dosing range The slope of each linear fit was then calculated and plotted as a function of CpG (green) and MPLA (red) doses. Dotted lines represent a slope of zero.

Production of the anti-inflammatory cytokine IL-10 was quantified (**Fig 10**). IL-10 was also upregulated by each adjuvant individually, as expected^{57, 58}, with similar responses across adjuvants (**Fig 10b,d,f** leftmost points). Positive dose dependent responses to CpG were maintained for nearly all combinations including MPLA and/or poly I:C (**Fig 10b, c**). In

contrast, the addition of either CpG or poly I:C dampened and reversed the dose dependent response to MPLA (**Fig 10d,e**). Positive dose dependent responses to poly I:C were also maintained for nearly all combinations including MPLA and/or CpG (**Fig 10b, c**).

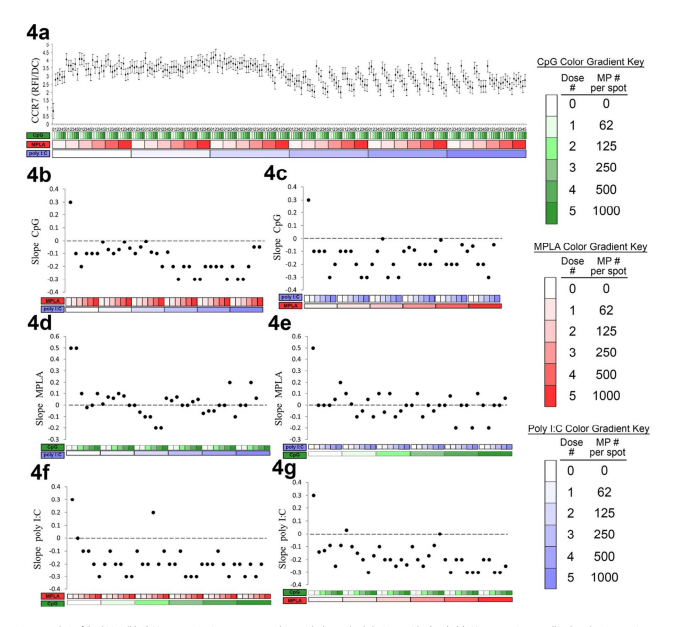


Fig 11: Analysis of dendritic cell (DC) CCR7 expression in response to combinatorial adjuvant-loaded microparticles (MPs). (a) CCR7 expression per cell is plotted against varying amounts of three MP-encapsulated adjuvants (CpG, MPLA and poly I:C; n = 9, with 3 replicates per array on 3 separate arrays). Data is arranged by grouping increasing particle numbers of CpG (green), MPLA (red), and lastly poly I:C (blue) with doses of 0, 62, 125, 250, 500 and 1,000 particles per spot, represented by increased shading from lighter to darker colors. (b, c) Linear fits were calculated for each set of CpG dosing range (lowest to highest number of particles) for every combination of MPLA (red) and poly I:C (blue) doses. The slope of each linear fit was then calculated and plotted as a function of MPLA and poly I:C doses. Dotted lines represent a slope of zero. (d, e) The adjuvant formulations

were then reorganized for every combination dose of CpG and poly I:C. Linear fits were created for the slope of the MPLA response associated with the specific dose of CpG and poly I:C. The slope of each linear fit was then calculated and plotted as a function of CpG (green) and poly I:C (blue) doses. Dotted lines represent a slope of zero. (f, g) The adjuvant formulations were then reorganized for every combination dose of CpG and MPLA. Linear fits were created for the slope of the poly I:C response associated with the specific dose of CpG and MPLA. Linear fits were calculated for each set of poly I:C dosing range The slope of each linear fit was then calculated and plotted as a function of CpG (green) and MPLA (red) doses. Dotted lines represent a slope of zero.

In addition to cytokine signaling, the chemokine receptor CCR7, important for DC lymph node homing^{40, 59}, was quantified (**Fig 11**). Expression of CCR7 was upregulated by each adjuvant alone (**Fig 11b,d,f** leftmost points). However, the addition of either MPLA and/or poly I:C reversed the dose dependent response to CpG (**Fig 11b,c**). In contrast, the dose dependent responses to MPLA were largely muted by the addition of either CpG or poly I:C (**Fig 11d, e**). Lastly, the addition of either MPLA and/or CpG reversed the dose dependent response to poly I:C (**Fig 11f, g**).

DC activation via TLR activation can occur via MyD88 dependent or MyD88 independent pathways, with simultaneous activation potentially eliciting synergistic effects on DC activation^{56, 60}. For instance, activation of TLR2/6, TLR9 (MyD88 dependent) along with TLR3 (MyD88 independent) in DCs has generated strong adaptive responses against HIV infection in mice⁶¹, highlighting the potential of signaling through non-overlapping pathways. In addition to TLRs, nucleotide-binding domain (NOD), leucine rich repeat containing family [Nod-like receptors (NLRs)], and the retinoic-acid inducible protein 1 [RIG-I-like receptors (RLRs)] can interact synergistically⁶² or antagonistically⁶³. Ultimately, the role of simultaneous activation of variable signaling pathways has been recognized, and vaccines are being developed that utilize multiple ligands to induce robust immune responses⁶².

Identifying adjuvant combinations that synergize to promote enhanced immune responses is central to the goal of developing more potent and precise vaccines. To that end, we sought to explore the interactions of TLR agonists CpG, MPLA, and poly I:C in their ability to either induce synergistic or antagonistic DC activation states. When examining surface expression of activation markers (MHC-II, CD86, CCR7) separately, there was a tendency for inhibitory interactions between adjuvants. Similar antagonistic interactions have been shown due to simultaneous activation of TLRs on DCs. For example, inhibitory interactions between TLR4 and TLR2, TLR4 and Dectin-1, TLR2 and TLR9, and TLR3 and TLR2 have been observed⁶⁴. On the other hand, cytokine production (IL-12, IL-10) maintained more positive dose dependent responses overall. Previous studies show augmentation of IL-12 production when MyD88 dependent (CpG) and MyD88 independent (poly I:C) pathways were activated simultaneously⁵⁶ which supports our findings. However this effect was not observed for surface expression of CD86, corroborating the results found here. Notably, in order to produce robust responses across all activation markers, adjuvant combinations were required, and combinations were more represented among the high responders. (Fig 6f). While mechanisms governing interaction effects between adjuvants were not investigated, considerations

may include aspects such as targeting the ligands to subcellular locations of each TLR, the competition of phagocytosis of differently formulated microparticles, the interplay between downstream signaling pathways^{56, 65}, and the dynamics of adjuvant-induced cytokine secretion^{66, 67}.

The task of resolving the interplay of TLR networks is arduous, necessitating the development of new tools, such as the immunoarrays here, which facilitate such discovery. A previous approach employed microfluidics to investigate combinatorial adjuvant signaling on DC cytokine production⁶⁸. They found MPLA to synergize with multiple TLR agonists in the production of IL-6, IL-12, and TNF-a with the strongest DC activation achieved by MPLA together with GardiquimodTM or CpG-B. While encouraging, this approach is limited to screening soluble adjuvants and quantifying cytokine secretion. In contrast, the immunoarray developed here is capable of evaluating cell surface activation markers in addition to cytokine secretion, thereby providing a more comprehensive assessment of DC activation. Further, our approach utilizes microparticle encapsulated adjuvants which have the advantage of providing vaccine stability, controlled release, and preferential targeting to DCs. While different, both approaches underscore the utility of miniaturized platforms to aid in vaccine development through major cost reductions and parallel analyses.

Conclusions

A significant barrier to high-content investigation of new drug candidates is the limited availability of compounds and relevant cells⁶⁹. Thus, technologies that aim to reduce compounds and cells required for screening greatly expedite identification of leads. Polymeric, particle-based approaches encapsulating adjuvants can provide delivery to DCs via phagocytosis and controlled release³⁵⁻³⁷. A recently described parallel particle production method describes rapid fabrication of hundreds of unique MP formulations⁷⁰. Integrating this method with the immunoarray platform may present a robust strategy to generate and quantify large libraries of factor-loaded MPs requiring limited cells and reagents. With thirteen TLR ligands identified to date, the task of resolving the combinatorial effects is challenging using standard techniques. Thus, a high-content approach mapping these interactions is enabling. This work demonstrates feasibility of a MP-based DC immunoarray platform to screen adjuvant combinations for the development of improved and personalized vaccines.

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