Immunostimulatory nanomedicines synergize with checkpoint blockade immunotherapy to eradicate colorectal tumors

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Nanoparticles can potentially stimulate tumour microenvironments to elicit antitumour immunity. Herein, we demonstrate effective immunotherapy of colorectal cancer via systemic delivery of an immunostimulatory chemotherapeutic combination in nanoscale coordination polymer (NCP) core-shell particles. Oxaliplatin and dihydroartemisinin have contrasting physicochemical properties but strong synergy in reactive oxygen species (ROS) generation and anticancer activity. The combined ROS generation is harnessed for immune activation to synergize with an anti-PD-L1 antibody for the treatment of murine colorectal cancer tumours. The favourable biodistribution and tumour uptake of NCPs and the absence of peripheral neuropathy allow for repeated dosing to afford 100% tumour eradication. The involvement of innate and adaptive immune systems elicit strong and long lasting antitumour immunity which prevents tumour formation when cured mice are challenged with cancer cells. The intrinsically biodegradable, well tolerated, and systemically available immunostimulatory NCP promises to enter clinical testing as an immunotherapy against colorectal cancer.
OxPt/DHA particles have desired surface properties to minimize uptake by the mononuclear phagocyte system (MPS), allowing for their selective accumulation in tumours after systemic injection. Comprised of two ICD-inducing therapeutics, OxPt/DHA elicits strong antitumour immunity in addition to anticancer efficacy, evidenced by early cell-surface exposure of calreticulin (CRT) and high mobility group box 1 (HMGB-1) protein release. Fragments of dead cancer cells were uptaken by phagocytes, leading to T-cell priming and antitumour vaccination. When supplemented with α-PD-L1, OxPt/DHA treatment completely eradicates CRC tumours, in addition to generating long-term tumour-specific immune memory responses to prevent the formation and growth of new CRC tumours in mouse models.

**Results**

**NCPs codeliver and stabilize synergetic OxPt and DHA.** We probed the synergy between the chemotherapeutic combination of OxPt and DHA at varying ratios on two murine CRC cell lines, CT26 and MC38. As shown in Supplementary Table 1, combining OxPt with DHA led to significant reduction of the OxPt IC50 values (from 9.1 ± 0.7 to 1.1–3.2 μM and 10.1 ± 1.1 to 1.1–3.6 μM on CT26 and MC38 cells, respectively).

To fix the molar ratios and prevent premature decomposition, we synthesized and encapsulated OxPt and DHA prodrugs, Pt(dach) (oxalate)(bisphosphoramic acid) (OxPt-bp) and cholest-5-en-3-ol (3b)-2-((2-((hydroartemesinincarbonyl)oxy)ethyl)disulfanyl)ethyl carbonate (choI-DHA) in NCPs (Supplementary Methods 1, 2 and Supplementary Figure 1, 2). OxPt/DHA particles were prepared in two steps (Fig. 1a). The NCP core of OxPt (OxPt-bare) was first synthesized by polymerization between Zn2+ ions and the phosphate groups of OxPt-bp in the presence of 1,2-dioleoyl-sn-glycerol-3-phospho-1-diesteryl-sn-glycerol-3-phospho-1-diesteryl-sn-glycerol-3-phosphate [DSPE-PEG2k] (molar ratio 2:1:1), and different amounts of chol-DHA to afford the core-shell nanoparticles OxPt/DHA with three different OxPt/DHA ratios (1:0.5, 1:1, and 1:2), with Z-average diameters of 73.8±103.4 nm, PDI of 0.12–0.17, and slightly negative surface charges of −20.8 to −13.0 mV in water (Fig. 1b, c and Supplementary Table 3).

DHA was conjugated to cholesterol via a disulphide bond to facilitate incorporation into the shell of the nanoparticles. The lipid bilayer protects chol-DHA from exposure to water and reductants, limiting decomposition and systemic release by preventing hydrolysis and reduction. The nanoparticles were stable with no changes in size or PDI at 4 °C for 1 year or 37 °C for 24 h in the presence of bovine serum albumin (BSA, Fig. 1d and Supplementary Figure 4). The monotherapy nanoparticle controls OxPt NCP and Zn/DHA prepared in the absence of chol-DHA or with pyrophosphate replacing OxPt prodrug, respectively, showed similar sizes and morphology (Fig. 1a, Supplementary Figure 5, 6 and Supplementary Table 2, 3).

Encapsulation into nanoparticles led to slightly higher IC50s than free drugs, but showed similar trends of OxPt IC50 reduction when adding chol-DHA (Supplementary Table 1), suggesting that the drugs are readily released from NCPs. The synergy between the two drugs in OxPt/DHA was most readily observed with a OxPt/DHA molar ratio of 1:0.5, with a combination index of <1 at nearly all effect levels (Supplementary Figure 7). This formulation was used for all further investigations. Using OxPt/DHA particles

**C**olorectal cancer (CRC) is the second leading cause of cancer-related deaths in the US, with an approximate lifetime risk of 1 in 20 people1. The standard therapy of surgery plus adjuvant chemotherapies is often limited by the side effects of and resistance to chemotherapy2-3. Great emphasis has thus been placed on developing immunotherapies for CRC treatment4, particularly after the Food and Drug Administration’s approval of the cytotoxic T-lymphocyte-associated protein 4 (CTLA-4) antibody ipilimumab in 2011, the programmed cell death protein 1 (PD-1) antibodies pembrolizumab and nivolumab in 2014, and the PD-1 ligand (PD-L1) antibody atezolizumab in 2015. Clinical trials of immune checkpoint inhibitors (α-CTLA-4, α-PD-1, α-PD-L1) have shown efficacy against many cancers, but limited effect in CRCs. A small subset of CRC patients having tumours with inherently high CD8+ T-cell infiltration and regulatory immune checkpoint overexpression have benefitted from α-PD-1 checkpoint blockade immunotherapy9-12. However, this cancer phenotype represents <5% of advanced stage CRC13,14. The PD-L1 antibody atezolizumab showed poor response in the predominant microsatellite stable form of CRC as a monotherapy, but improved overall response rates in combination with an MEK inhibitor or α-VEGF and standard folic acid, 5-fluorouracil, and oxaliplatin (FOLFOX) chemotherapy15. There is thus an established need for therapies that can improve tumour immunogenicity and induce CD8+ T-cell infiltration to enhance immunotherapy for the broader population of CRC patients.

Most chemotherapy regimens are considered immunologically silent or even tolerogenic. However, a subset of chemotherapeutics have recently been shown to be proinflammatory and capable of inducing immunogenic cell death (ICD), suggesting their potential for combination with checkpoint blockade to afford antitumour immunity. A key component of the FOLFOX regimen, oxaliplatin (OxPt), was identified as an ICD inducer16-19. We have recently shown that OxPt can be combined with a photosensitizer to synergize the ICD of OxPt-based chemotherapy and photodynamic therapy (PDT) and prime the tumour microenvironment for α-PD-L1 therapy20. However, PDT is a localized therapy and light penetration is limited to superficial tumours, severely limiting the clinical translation of PDT-based combination therapies in immunotherapy of CRC.

As reactive oxygen species (ROS) are primarily responsible for cancer cell death by PDT, we posited that ROS-based chemotherapeutics could induce strong ICD using chemicals instead of photons for effective priming of the tumour microenvironment. We report here that the ROS-producing drug, dihydroartemisinin (DHA), can efficiently induce ICD and exhibits synergy with OxPt. DHA is an active metabolite of artesminin derivatives, an antimalarial drug well tolerated by millions of patients21. DHA contains an endoperoxide bridge that reacts with a ferrous iron catalyst to generate free radicals and cause oxidative stress, similar to the effects of PDT. Malignant cells alter iron metabolism to increase uptake and decrease efflux for tumour growth, leading to an increased pool of labile iron and thus tumour-specific activity of DHA. Despite significant potential as an anticancer therapeutic, DHA’s power has not been harnessed for in vivo antitumour treatment due to its instability in aqueous media and low bioavailability. The endoperoxide moiety that endows its antimalarial and anticancer activity reacts non-specifically, leading to premature deactivation in circulation.

Herein, we developed self-assembled nanoscale coordination polymer (NCP) core-shell nanoparticles carrying OxPt in the core and DHA in the shell (OxPt/DHA) for their selective delivery to CRC tumours. In OxPt/DHA core-shell particles, an NCP of Zn and OxPt prodrug was coated with a lipid bilayer containing a cholesterol-DHA conjugate (chol-DHA). OxPt/DHA particles

sequester the drugs from water, reductants, and proteins, enabling spatiotemporal control of drug releases in tumours and reducing systemic drug exposure. OxPt/DHA particles have desired surface properties to minimize uptake by the mononuclear phagocyte system (MPS), allowing for their selective accumulation in tumours after systemic injection. Comprised of two ICD-inducing therapeutics, OxPt/DHA elicits strong antitumour immunity in addition to anticancer efficacy, evidenced by early cell-surface exposure of calreticulin (CRT) and high mobility group box 1 (HMGB-1) protein release. Fragments of dead cancer cells were uptaken by phagocytes, leading to T-cell priming and antitumour vaccination. When supplemented with α-PD-L1, OxPt/DHA treatment completely eradicates CRC tumours, in addition to generating long-term tumour-specific immune memory responses to prevent the formation and growth of new CRC tumours in mouse models.
labelled with cholesterol-conjugated pyropheophytin a (chol-pyro, Supplementary Methods 3 and Supplementary Figure 8), we showed that particles were taken up rapidly by cells, with ~95% of all cells showing particle fluorescence after 1 h. The fluorescence intensity significantly increased over time, indicating continual nanoparticle uptake (Fig. 2a). Inductively coupled plasma-mass spectrometry (ICP-MS) quantification of intracellular Pt also revealed time-dependent uptake of both nanoparticles and free OxPt. However, the uptake of nanoparticles was much less than free OxPt (Fig. 2b), possibly due to the surface PEG coating preventing interaction between particles and cells. The lower uptake explains why the particles have slightly higher IC50 than free drugs in vitro. To visualize the intracellular release of drugs, fluorescent nanoparticles were synthesized by doping xylenol orange (blue) into the core and coating chol-pyro (red) and FITC-DOPE (green) on the shell, which allows simultaneous visualization of OxPt in the core, chol-DHA in the lipid shell, and the lipid layer, respectively (Supplementary Methods 4). In the first 10 min, all three dyes were primarily found on the cell surface, viewed as cyan (green and blue merged) and/or white (green, blue and red merged) in the merged image, indicating that the particles were mainly bound to the cell surface. Over time, the xylenol orange and chol-pyro nonspecifically distributed inside the cells as magenta (blue and red merged) fluorescence with varying intensity, suggesting independent release of the drugs. FITC-DOPE mainly localized to the plasma membrane (Fig. 2c), possibly due to the lipid bilayer fusion with endosome membranes to expose the NCP core and the trafficking of FITC-DOPE to the plasma membrane with the recycled endosomes. These data demonstrate that upon cellular uptake, the core–shell structure is disrupted to expose the OxPt NCP core and chol-DHA to high concentrations of intracellular reducing agents, such as glutathione (GSH, 5 mM) and ascorbate (100 μM), which further reduce and/or hydrolyse the prodrugs into parent drugs to exert efficacy.

**Fig. 1** Preparation and characterization of OxPt/DHA. a Schematic illustration showing layer-by-layer construction of the hybrid core-shell structure of OxPt/DHA. The OxPt/DHA consists of an OxPt prodrug coordinated to Zn2+ ions in the core and chol-DHA in the lipid shell. Compositions of the three NCPs investigated are also shown. b TEM image of OxPt/DHA. c Number-average diameter of OxPt/DHA characterized by DLS. d Stability test of OxPt/DHA at 37 °C in the presence of BSA (5 mg/mL). OxPt oxaliplatin, DHA dihydroartemisinin, TEM transmission electron microscopy, DLS dynamic light scattering

Triggered release of OxPt and DHA from NCPs. The disulphide linkage of chol-DHA was cleaved by GSH at a physiologically relevant concentration to release DHA (Fig. 3a, Supplementary Figure 9). However, the kinetics of DHA release in the 5 mM GSH solution of phosphate-buffered saline (PBS) was faster than that of GSH reduction alone. Although chol-DHA was stable in aprotic, organic solvents such as THF at 37 °C for >24 h, it rapidly decomposed (>80%) in aqueous or protic solvents (Fig. 3c, Supplementary Figure 9). The addition of acids further accelerated the decomposition of chol-DHA (Supplementary Figure 9).

Based on these observations, we propose a DHA release mechanism via proton-catalysed hydrolysis in aqueous media. The protonation of the carbonate carbonyl oxygen atom in protic solvents can reversibly cleave the C–O bond next to DHA to form a DHA cation stabilized by the nearby oxygen and a carbonate monoester. The carbonate monoester irreversibly releases CO2, providing the driving force for the hydrolysis of
DHA from chol-DHA. While GSH reduction of the disulphide linkage contributes to the release of DHA, hydrolysis of the chol-DHA carbonate linker is predominantly responsible for DHA release (Fig. 3a, Supplementary Figure 10a-b). Importantly, incorporation of chol-DHA into the lipid bilayer shell of OxPt/DHA prevented premature DHA release by limiting exposure to water. Incubation of OxPt/DHA in water at 37 °C for 24 h did not lead to significant loss of chol-DHA. However, disruption of the lipid bilayer of OxPt/DHA by Triton X-100 led to 90% degradation of chol-DHA in water at 37 °C within 5 h (Fig. 3c). The core–shell structure of OxPt/DHA thus protects DHA from exposure to water and reductants during circulation, ensuring the selective delivery of DHA to cancer cells.

The Pt(IV) prodrug OxPt-bp was also reduced via two mechanisms: direct reduction into OxPt or a two-step sequence of hydrolysis to generate Pt(dach)(oxalate)(biscarbamate) (OxPt-bc) followed by reduction to OxPt (Fig. 3b, Supplementary Figure 10c-d, 11). The structure of OxPt-bc was confirmed by single crystal X-ray diffraction (Supplementary Figure 12, Supplementary Table 4). Interestingly, only ascorbate and not
GSH can reduce either OxPt-bp or OxPt-bc to afford OxPt (Fig. 3b, Supplementary Figure 10b). While OxPt does not release from intact OxPt/DHA particles, the disruption of the lipid bilayers upon endocytosis allows access of the robust coordination polymer to ascorbate and acidic aqueous medium, leading to release of both OxPt and OxPt-bc (Fig. 3b, c, Supplementary Figure 10c).

Cell death by classically programmed and immunogenic pathways. The purported antimalarial activity of DHA involves cleavage of the endoperoxide bridge by ferrous iron, generating oxygen- and carbon-based radicals. As cancer cells are highly sensitive to agents that can augment oxidative stress, we investigated whether our combination therapy can synergistically generate ROS, which can directly react with the membrane, DNA, proteins, and organelles or generate secondary products to cause damage. Both OxPt and DHA can individually induce ROS generation in tumour cells, with a significant increase when given in combination (p < 0.001; Fig. 4a, b and Supplementary Figure 13). ROS is a known trigger for dysfunction of mitochondria, which regulates both autophagy and apoptosis. The generated ROS causes release of cytochrome c from mitochondria, as evidenced by the decrease in the colocalization between the mitochondria (red) and the cytochrome c (green) fluorescence (Fig. 4c, d and Supplementary Figure 14), disrupting the membrane potential as a consequence of ROS accumulation. As a result, both OxPt and DHA induced programmed cell death by apoptosis/necrosis (Fig. 4e, f and Supplementary Figure 15). The combination of OxPt and DHA increased both early apoptotic Annexin V+ cells (26.8 ± 1.4% compared to 11.9 ± 1.0% and 14.7 ± 2.5% for OxPt and DHA, respectively) and late apoptotic/necrotic Annexin V+/PI+ cells (36.2 ± 3.0% compared to 15.6 ± 1.5% and 31.6 ± 2.9% for OxPt and DHA, respectively). Treatment with OxPt NCP, Zn/DHA, and OxPt/DHA led to similar trends in the ROS, cytochrome c release, and induction of apoptosis (Fig. 4a–f and Supplementary Figure 13-15).

In addition to mitochondrial dysfunction, ROS can also inhibit cell growth by cell cycle arrest via endoplasmic reticulum (ER) stress. G2/M phase cell cycle arrest was observed in CT26 cells treated by either OxPt or DHA, increasing the percentages of cells in the G2/M phase to 35.6 ± 3.7% (p = 1.1 × 10⁻²) and 34.5 ± 3.9% (p = 1.5 × 10⁻²), respectively, from 20.8 ± 4.4% in PBS. Combining OxPt and DHA further increased the proportion of
cells arrested at the G2/M phase to 44.3 ± 3.7% ($p = 4.4 \times 10^{-2}$ and $3.4 \times 10^{-2}$ compared to OxPt and DHA, respectively). Two sequential, redundant mechanisms for ER stress-regulation of the cell cycle have been suggested: first delay at the G2 checkpoint followed by cell cycle arrest at the G1 checkpoint.22 Interestingly, treatment with OxPt specifically decreased the accumulation of S phase cells, whereas treatment with DHA led to approximately equal reductions of cells in the G0/G1 and S phases. Cells treated with OxPt NCP, Zn/DHA, and OxPt/DHA resulted in similar changes in the cell cycle as free drug treatments (Fig. 4g, h and Supplementary Figure 16).

ER stress and ROS production are essential components of the intracellular pathways that govern ICD, which occur in parallel to activate danger signalling pathways that help to traffic damage-associated molecular patterns (DAMPs) to the extracellular space.16,23–26 We confirmed that OxPt is able to induce ICD, and demonstrated that DHA can also effectively induce ICD, as evidenced by calreticulin (CRT) cell-surface exposure (Fig. 5a, b, and Supplementary Figure 17). We further quantified the release of high mobility group box-1 (HMGB-1) from cells treated with both drugs by enzyme-linked immunosorbent assay (ELISA). Incubation with OxPt or DHA led to increased HMGB-1 release from cells, which was further increased by coincubation with both drugs (Fig. 5c).

**Fig. 4** Programmed cell death in colorectal cancer cells by ROS generation. **a, b** ROS generation in cells treated with OxPt/DHA, as indicated by the green fluorescence of 2′,7′-dichlorofluorescein (DCF) that was oxidized from 2′,7′-dichlorodihydrofluorescein diacetate (H2DCFDA) by ROS. **c, d** Release of cytochrome c from mitochondria in cells incubated with OxPt/DHA. Mitochondria (red fluorescence) and cytochrome c (green fluorescence) were stained by MitoTracker Red CMXRos and anti-cytochrome c antibody, respectively. **e, f** Apoptosis induced by OxPt/DHA. After treatment, cells were stained by Alexa Fluor 488-labelled Annexin V and propidium iodide (PI) and analysed by flow cytometry. **g, h** Cell cycle arrest caused by OxPt/DHA. Treated cells were fixed with 70% ethanol overnight, treated with RNase A, stained by PI, and analysed by flow cytometry. Data are expressed as means ± SD, and one of three repetitions with similar results is shown here. *p < 0.05, **p < 0.01, ***p < 0.001 by Student’s two-tailed t test. OxPt oxaliplatin, DHA dihydroartemisinin, ROS reactive oxygen species.
The result was obtained without repetition for (f) (n = 6). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 by Student’s two-tailed t test. CRT calreticulin, OxPt oxaliplatin, DHA dihydroartemisinin, CLSM confocal laser scanning microscopy

demonstrated by staining of the SIINFEKL-H2kb complex on the surfaces of DCs and macrophages (Supplementary Figure 21, 22). This result suggests that both phagocytes are involved in presenting tumour antigens to initiate the adaptive immune response.

To investigate whether OxPt/DHA could prime T cells, dead and/or dying MC38 cells treated with OxPt/DHA were inoculated into the footpads of C57BL/6 mice. Six days after inoculation, the regional popliteal lymph nodes were excised and stimulated with MC38 lysates ex vivo. Both OxPt- and DHA-treated cells were able to prime T cells for IFN-γ production (Fig. 5f), with the combination of OxPt and DHA showing the highest ability to prime T cells. In addition, the T cell priming ability of OxPt/DHA-treated MC38 cell lysates was much stronger than that of the known MC38 antigen KSPWFTTL (Supplementary Figure 23).

Activation of T cells by OxPt and/or DHA treatment led to efficient vaccination specifically against MC38. OxPt- or DHA-treated cells reduced the frequency of tumours developing from live cells to 33 and 17%, respectively, by day 30 (Fig. 5g). In comparison, 100% mice developed tumours with PBS-treated cells. This is consistent with in vitro results showing DHA is a stronger ICD inducer than OxPt, with a greater percentage of CRT⁺ cells and more HMGB-1 secretion. No tumour formation occurred when live MC38 cells were inoculated into mice vaccinated with OxPt/DHA 30 days prior and challenged with untreated, live MC38 cells in one flank and challenging mice with untreated, live tumour cells in the opposite flank 7 days later. Data are expressed as means ± SD. One of three repetitions with similar results is shown here for (a)–(e). The result was obtained without repetition for (f)–(h) (n = 6). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 by Student’s two-tailed t test. CRT calreticulin, OxPt oxaliplatin, DHA dihydroartemisinin, CLSM confocal laser scanning microscopy.

**Fig. 5** Immunostimulatory effects in colorectal cancer cells. **a, b** CRT cell surface expression upon treatment with OxPt/DHA, determined by flow cytometry (a) and CLSM (b). **c** HMGB-1 release from tumour cells treated with OxPt/DHA, detected by ELISA. **d–f** Uptake of treated MC38 cells by bone-marrow-derived dendritic cells (d) and macrophages (e). **f** Priming of T-cell responses triggered by OxPt/DHA. MC38 tumour cells were treated with OxPt/DHA, and injected into the right footpads of C57BL/6 mice to determine the capacity of draining lymph node cells to produce IFN-γ in response to MC38 lysates. **g, h** In vivo anticancer vaccination of OxPt/DHA in immunocompetent C57BL/6 mice (g) and immunodeficient Rag2⁻/⁻ mice (h). The antitumour response was measured by immunizing mice with OxPt/DHA-treated tumour cells in one flank and challenging mice with untreated, live tumour cells in the opposite flank 7 days later. Data are expressed as means ± SD. One of three repetitions with similar results is shown here for (a)–(e). The result was obtained without repetition for (f)–(h) (n = 6). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 by Student’s two-tailed t test. CRT calreticulin, OxPt oxaliplatin, DHA dihydroartemisinin, CLSM confocal laser scanning microscopy.

**References**

27. Recognizing the immunogenicity of OxPt and DHA, we investigated OxPt/DHA in combination with α-PD-L1 checkpoint blockade. Syngeneic tumour models of CT26 and MC38 were established on immunocompetent BALB/c and C57BL/6 mice, respectively, and tumours were allowed to grow for approximately 12 days, reaching 80–120 mm³ before treatment. As minimal toxicity by body weight was observed after 50 doses of Zn/DHA at 5 mg DHA/kg, four weekly doses of OxPt/DHA at 60 mg OxPt/kg, or one dose of OxPt/DHA at 80 mg
Fig. 6 Enhanced anti-PD-L1 immunotherapy on colorectal cancers. a Experimental design for the treatment and challenge of CT26 tumour-bearing mice. Tumours were allowed to grow for 12 days before treatment to form more immunosuppressive tumours. Then, tumour-bearing mice were intraperitoneally injected with OxPt/DHA combined with α-PD-L1 every 3 days for 12 total doses. Three months after all tumours had disappeared, mice were challenged with CT26 cells, followed by rechallenge with 4T1 1 month later. b Growth curves of CT26 tumours after treatment with OxPt/DHA combined with α-PD-L1 and challenge with CT26 cells (red arrow). c Experimental design for surgery control of CT26 tumour-bearing mice. Percentage tumour-free mice (d, f) and tumour growth curve (e, g) after challenge with CT26 cells (d, e) or rechallenge with unrelated 4T1 tumour cells (f, g) in naïve mice or OxPt/DHA and α-PD-L1-treated mice. h Growth curves of MC38 tumour on C57BL/6 after treatment with OxPt/DHA (8 mg/kg OxPt) combined with α-PD-L1. i, j Therapeutic effect of OxPt/DHA plus α-PD-L1 on C57BL/6 (i) at the dose of 16 mg/kg OxPt and Rag2−/− mice (j) at the dose of 8 mg/kg OxPt. Data are each pooled from two independent experiments and expressed as means ± SD (n = 6 except for (j); n = 5 for (j)). OxPt oxaliplatin, DHA dihydroartemisinin.

OxPt/kg (Supplementary Figure 26), we chose a dose of 8 mg OxPt/kg, 2.8 mg DHA/kg, and/or 75 μg α-PD-L1/mouse. This dose was at least ten times lower than the maximum tolerated dose (MTD) and allowed for frequent low-dose metronomic dosing as opposed to conventional infrequent doses at or near the MTD28. Mice were i.p. injected once every 3 days for up to 12 total doses (Fig. 6a). With significant OxPt accumulation in tumours 72 h post administration (Supplementary Figure 27), this dosing schedule allowed for a dose-dense chemotherapy schedule with a near continuous presence of chemotherapy in the tumour. This was in line with the Norton–Simon hypothesis that more frequent doses will lead to greater clinical benefit by minimizing the opportunities for cancer regrowth between doses29. Importantly, OxPt/DHA increased the effective doses of OxPt and DHA by ~15 and 1000 times, respectively (Table 1). Nearly all of the free drugs were cleared out of or decomposed in the bloodstream within 1 h of intravenous injection (Supplementary Figure 28, 29). Despite this, free OxPt leads to significant peripheral neuropathy while OxPt/DHA showed no such toxicity (Supplementary Figure 30).
In CT26 tumour-bearing mice, the low-dose α-PD-L1 treatment alone was ineffective at controlling tumour growth. The combination of free OxPt, DHA, and α-PD-L1 proved moderately effective (average tumour volumes of 203.27 ± 81.00 mm³ on day 18 compared to 616.80 ± 46.59 mm³ for PBS) but extremely toxic, as the body weights steadily decreased (Supplementary Figure 31). All mice had to be euthanized after three doses for humanitarian reasons in accordance with our animal protocols. We hypothesized that NCPs can decrease the toxicity of OxPt and DHA by providing a favourable biodistribution profile. In CT26 tumour-bearing mice, <5%ID Pt/g accumulated in key organs such as the liver, spleen, and kidney indicating OxPt/DHA is not significantly uptaken by the MPS. Furthermore, there was <3%ID Pt/g accumulation in the heart and lung, suggesting OxPt/DHA does not aggregate in circulation. The low MPS uptake and slow clearance led to progressive accumulation in tumour to a maximum of 12.3 ± 2.8 %ID/g at 48 h post administration as a result of the enhanced permeability and retention effect compared to 0.56 ± 2.8%ID/g by the free drug (Supplementary Figure 27, 32). OxPt NCP with or without α-PD-L1 was well tolerated and led to similar tumour growth rates, significantly controlling tumour growth and retarding tumour growth for over 1 month. Interestingly, though Zn/DHA did not show substantial anticancer efficacy alone or in combination with α-PD-L1 (Supplementary Figure 33), it significantly enhanced the efficacy of OxPt and delayed tumour growth to ~2 cm³ until day 66. The tumour growth curve of OxPt/DHA plus α-PD-L1 was initially similar to that of OxPt/DHA, but started to deviate around day 18, after which all of the tumours regressed and eventually disappeared on days 40–50. No tumour recurrence was observed for a period of 120 days (Fig. 6b).

The antitumour immune response initiated by OxPt/DHA plus α-PD-L1 resulted in a memory response; OxPt/DHA plus α-PD-L1-treated mice which were tumour-free for at least 120 days were challenged with live CT26 cells on the contralateral flank. No mice grew new tumours over the next month compared to cells implanted in naïve mice (Fig. 6f, g).

We confirmed these results on the more immunosuppressive model of MC38 tumours in C57BL/6 mice, which yielded similar results under the same treatment regimen (Fig. 6h, Supplementary Figure 34). The chemoimmunotherapy of free OxPt, DHA, and α-PD-L1 was initially effective at controlling tumour growth, but eventually the tumours grew aggressively and the mice had to be euthanized due to the tumour burden on day 39. OxPt/DHA showed similar anticancer efficacy, but continued to control the tumours until day 54. Tumours treated with OxPt/DHA plus α-PD-L1 regressed around day 18 and were controlled for an extended period of time, but eventually grew back (Fig. 6h).

By increasing the chemotherapy dose of OxPt/DHA to 16 mg OxPt/kg and 4.6 mg DHA/kg plus 75 μg α-PD-L1/mouse, three out of five treated mice were tumour free and the remaining two mice showed prolonged tumour growth control (Fig. 6i). The contribution of the immune system to OxPt/DHA plus α-PD-L1 antitumour efficacy was immediately obvious as no therapeutic effect was observed when MC38 tumours implanted in immunodeflcient Rag2−/− mice were treated with OxPt/DHA plus α-PD-L1 (Fig. 6j).

### Table 1 OxPt and DHA pharmacokinetic information

| Parameter (Unit) | t1/2 alpha (h) | t1/2 beta (h) | AUC 0-inf (ID%mL⁻¹ h⁻¹) | AUMC (ID%mL⁻¹ h²) | MRT (h) |
|------------------|----------------|---------------|--------------------------|-------------------|--------|
| Free OxPt        | 0.38 ± 0.02    | 14.67 ± 3.04  | 5.77 ± 2.65              | 122.5 ± 77.5      | 20.1 ± 4.4 |
| Free DHA        | (0.22 ± 0.04)  | (8.09 ± 11.2) | (33.6 ± 5.8)            | (300.7 ± 79.8)    |
| OxPt/DHA         | 0.27 ± 0.30(0.39 ± 0.38) | 20.70 ± 4.50(7.01 ± 1.13) | 80.9 ± 11.2(33.6 ± 5.8) | 2431.5 ± 809.7(330.2 ± 79.8) | 29.51 ± 6.27(9.79 ± 1.41) |

Data are expressed as means ± SD

OxPt oxaliplatin, DHA dihydroartemisinin

*This was fitted to a one-compartment model

**The numbers in parentheses refer to DHA values

Engaging the innate and adaptive immune systems. Immunogenic therapies are known to induce an innate immune response, including rapid infiltration of phagocytic DCs and macrophages. As the activation of DCs may constitute the first of several steps in the immune development process, we first investigated the intratumoural levels of antigen-presenting innate immune cells. We observed increased tumour infiltration of CD11c⁺ and F4/80⁺ cells 2 days after the first treatment with OxPt/DHA plus α-PD-L1 (Fig. 7a–c). The main function of DCs is to process antigen materials and present them to T cells to promote immunity, while macrophages can engulf and digest cellular debris and secrete proinflammatory cytokines to recruit other immune cells in addition to presenting antigens to DCs. The enhanced infiltration of DCs and macrophages in tumours might have resulted from the improved immunogenicity caused by OxPt/DHA, allowing for efficient antigen capture and presentation. We also investigated the infiltration of DCs and macrophages 12 days after the first treatment. The percentage of DCs slightly, not significantly, increased in tumours treated with OxPt/DHA alone or in combination with α-PD-L1 (Fig. 7e). OxPt/DHA plus α-PD-L1 significantly increased the percentage of M1 macrophages in total macrophages in tumours (Fig. 7f), suggesting the combination treatment enhances polarization to M1 macrophages or recruits more M1 macrophages into tumours to facilitate antigen presentation. By recruiting DCs and macrophages to phagocytose dying and/or dead cancer cells and enhancing the processing/presentation of tumour-associated antigens to T cells, OxPt/DHA may have increased the density of CD8⁺ T cells in tumours to potentiate checkpoint blockade immunotherapy.

Tumours with low densities of CD8⁺ T cells, such as MC38, do not generally respond to PD-1/PD-L1 blockade. Effective combination therapy can increase the intratumoural infiltration of CD8⁺ T cells to significantly increase the response rate. Immunofluorescence analysis performed 12 days after the first treatment with OxPt/DHA plus α-PD-L1 showed significant
Immunoprimarily CD8+ T cells. The number of antigen-specific leucocyte or CD4+ cells increased in the density of innate immune cells. The percentages of dendritic cells (d), total macrophages (e) and M1 macrophages (f) by flow cytometry of cell-surface staining 12 days after the first treatment. Data are each pooled from three independent experiments for (d) and (e). Data were obtained without repetition for (f). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001 by Student's two-tailed t test.

Discussion
As a well-tolerated antimalarial drug, DHA is a strong candidate for anticancer therapy. Higher levels of iron in tumours catalytically decomposes the reactive endoperoxide to generate ROS. However, the low bioavailability and rapid decomposition of DHA in sera present two major obstacles to using DHA in cancer therapy. By synthesizing a cleavable prodrug and encapsulating it into the lipid bilayer shell of an NCP, DHA can be shielded from the water in circulation and released only after particle dissociation upon cellular uptake. Intracellular release of DHA from chol-
DHA can occur via disulphide cleavage and hydrolysis. The OxPt-bp coordination polymer can also release free OxPt directly by ascorbate reduction or via hydrolysis to OxPt-bc followed by reduction (Fig. 3a, b). The dual release pathways for both DHA and OxPt may be beneficial in avoiding resistance mechanisms. Specifically, the OxPt-bc prodrug is resistant to deactivation by GSH and thiol-containing proteins, allowing for prolonged drug exposure when reduced to parent OxPt by ascorbate in cytosols and even nuclei of cancer cells.

Chemotherapies are often beleaguered by toxicities arising from accumulation in healthy tissues, which can be alleviated with nanoparticle delivery. OxPt/DHA are optimally sized pegylated nanoparticles that are large enough to avoid renal filtration (~10 nm) but small enough to penetrate through the leaky tumour vasculatures in tumours. The use of ~20 mol% PEG in this formulation helps to reduce plasma protein binding, thus minimizing MPS uptake after systemic injection. OxPt/DHA showed low uptake in the liver and other major organs associated with clearance, and significantly increased both the single and repeated dose MTDs of OxPt. Mice were dosed near the free drug MTD, but only one tenth the single dose MTD of the NCP formulation, allowing for dose-dense metronomic chemotherapy. This significantly reduced the most common dose-limiting toxicity of peripheral neuropathy while maintaining strong anticancer efficacy and immunity, suggesting that OxPt/DHA may be a strong clinical candidate.

Nanoparticle-supported chemoimmunotherapy has been increasingly studied to alter the tumour microenvironment in recent years. Though many chemotherapeutic agents mediate their cytotoxic effects by inducing immunologically silent or tolerogenic apoptosis, certain chemotherapies kill cancer cells via immunogenic apoptosis. This changes the cell-surface composition and releases DAMPs, of which there are three hallmarks: (i) the preapoptotic exposure of CRT on the cell surface, (ii) release of ATP during the blebbing phase of apoptosis, and (iii) post-apoptotic exodus of the chromatin-binding protein HMGB-1. While OxPt is a known ICD inducer, we have directly shown that treatment of CRC cells with DHA causes translocation of CRT to the cell surface and HMGB-1 release. This facilitates the recruitment of antigen-presenting cells (APCs) into the tumour

Fig. 8 OxPt/DHA promotes tumour-specific T cell response. a Immunofluorescence analysis showing the infiltration of CD8+ T cells 12 days after the first treatment. b The density of CD8+ T cells in the tumour sections, analysed from the confocal images of immunofluorescence staining (n = 3). c CD8+ T cells in tumours by flow cytometry of cell-surface staining 12 days after the first treatment. d KSPWFTTL antigen-specific IFN-γ producing T cells detected by ELISPOT assay 12 days after the first treatment. e The percentage of effector memory T cells in total CD8+ T cells in spleens after treatment with OxPt/DHA plus α-PD-L1. Data are each pooled from two independent experiments for (c)–(e). *p < 0.05, **p < 0.01, ***p < 0.001 and ****p < 0.0001 by Student’s two-tailed t test. OxPt oxaliplatin, DHA dihydroartemisinin, ELISPOT Enzyme-Linked ImmunoSpot.
bed (stimulated by ATP), the engulfment of dying tumour cells and their debris by APCs (stimulated by CRT)23,24, and optimal antigen presentation to T cells (stimulated by HMGB-1)35,56. DHA-treated CRC cells were uptaken more by phagocytes and led to strong T cell priming and antitumour vaccination. Altogether, these processes result in a potent IFN-γ-mediated immune response, which eventually can lead to tumour rejection57.

OxPt/DHA combined with α-PD-L1 is highly effective in treating well-developed tumours of CT26, likely due to innately high T cell and low suppressor cell infiltration58,59. This treatment led to eradication of 6/6 tumours and tumour-specific immune responses resulting in vaccination against subsequent cell challenge. In contrast, the MC38 tumour model is highly immunosuppressive, with myeloid-derived suppressor cells constituting >50% of the CD45+ immune cells in tumours58. A higher dose of OxPt/DHA could approximately recapitulate the effects observed in CT26, with tumour eradication in 3/5 mice and long-term tumour control in the other two. Treatment with OxPt/DHA led to ICD in tumours followed by infiltration of and engulfment by phagocytes. These APCs can present tumour-specific antigens to naive T cells in the lymph nodes and eventual T cell infiltration into the tumours as part of the adaptive immune response. Anti-PD-L1 ameliorates immune suppressive mechanisms of tumour cells causing immune evasion and T-cell anergy and/or exhaustion (Fig. 9)60.

In summary, we present an approach to initiate and stimulate anticancer and immunostimulatory properties of ROS-producing DHA and OxPt. By inducing CRT expression and HMGB-1 release, OxPt/DHA directly converted treated tumours into an in situ vaccine, recruiting antigen-presenting DCs and macrophages, facilitating cancer cell phagocytosis, enhancing antigen processing and presentation, and finally increasing intratumoural infiltration of CD8+ T cells to significantly potentiate checkpoint blockade immunotherapy. The animals treated by metronomic dosing of OxPt/DHA and α-PD-L1 were tumour-free for at least 3 months and immunized against live tumour cell challenge by generating long-term antitumour immunity. The scalable and tunable nature of NCP synthesis should allow their rapid optimization to lead to potential clinical candidates for combination therapy with immune checkpoint inhibitors.

Methods

Materials, cell lines, and animals. All starting materials were purchased from Sigma-Aldrich and Fisher (USA), unless otherwise noted, and used without further purification. 1,2-dioleoyl-sn-glycero-3-phosphate (DOPA), 1,2-dioleyl-sn-glycero-3-phosphocholine (DOPC), cholesterol, and 1,2-distearoyl-sn-glycero-3-phosphoethanolamine-N-[amino(polyethylene glycol)2000] (DSPE-PEG2k) were purchased from Avanti Polar Lipids (USA).

Murine colon adenocarcinoma cell CT26 and MC38 cells, mouse mammary carcinoma cell 4T1, mouse Lewis lung carcinoma cell LL/2 were all obtained from the American Type Culture Collection (ATCC, Rockville, MD). CT26 cells were cultured in RPMI 1640, MC38, 4T1 and LL/2 were grown in Dulbecco’s modified Eagle’s medium, respectively, supplemented with 10% FBS, 100 U/mL penicillin G sodium and 100 μg/mL streptomycin sulphate in a humidified atmosphere containing 5% CO2 at 37 °C. Mycoplasma was tested before use by MycoAlert detection kit (Lonza Nottingham, Ltd.). BALB/c female mice (6 weeks, 18–22 g), C57BL/6 female mice (6 weeks, 18–22 g), Rag2−/− female mice (6 weeks, 18–22 g), and CD/SD female rats (6 weeks, 160–200 g) were provided by Harlan-Envigo Laboratories, Inc (USA). We have
In vitro cytotoxicity. CT26 cells or MC38 cells were seeded in 96-well plates at a density of 2 × 10⁴ cells per well and allowed to adhere for 24 h. Cells were then treated with different concentrations of free OxPt, free DHA, or OxPt plus DHA (OxPt/DHA) for 24 h, with or without DHA. The medium was collected for detection of HMGB-1 secretion by ICP-MS and LC-MS, respectively. The livers, lungs, kidneys, and spleens were harvested for the determination of microarrays. The ability of OxPt/DHA-treated MC38 cells to promote the secretion of microarrays was determined by ELISA (Chondrex, Redmond, WA).

Antigen presentation. Bone-marrow-derived dendritic cells and macrophages were cocultured with free drugs or nanoparticles at a dose of 10 µM OxPt or/and 5 µM DHA for 24 h. The medium was collected for detection of HMGB-1 secretion by ELISA according to the manufacturer’s instructions (Chondrex, Redmond, WA).

Phagocytosis assay. Bone-marrow-derived dendritic cells and macrophages were isolated according to previously published protocols. Briefly, the hind legs of mice were cut away just above the hip and below the ankle. The muscle was cut away and the bones were soaked in 70% ethanol for 30 s. In a sterile environment, the ends of both bone were cut away and an insulin syringe filled with RPMI complete media was inserted into the exposed end of the bone and used to flush out the bone marrow into a cell culture dish filled with RPMI complete media. This process was repeated until the bone appeared white and translucent. For DC activation, bone-marrow-derived monocytes were cultured with GM-CSF (20 ng/mL) and IL-4 (10 ng/mL) for 6 days, then nonadherent cells in the culture supernatant were harvested and the expression of CD11c, CD11b, F4/80 and Gr-1 was analysed by flow cytometry to determine the purity of DC before further use. For macrophage differentiation, bone-marrow-derived monocytes were cultured with SF (20 ng/mL) for 6 days, then adherent cells were harvested and the expression of CD11c, CD11b and F4/80 was analysed by flow cytometry to determine the purity of macrophage before further use. TdTamato-transfected MC38 tumour cells were first incubated with OxPt/DHA at 10 µM OxPt and 5 µM DHA for 24 h, then treated tumour cells were cocultured with F4/80-labelled (red) macrophages or F4/80-labelled (blue) macrophages for another 1 h at 37 °C. Cells were then collected, washed twice with cold PBS, resuspended in PBS, analysed by flow cytometry, and calculated as the percentage of TdTamato+ macrophages or CD11c+ DCs.

Antigen presentation. Bone-marrow-derived dendritic cells and macrophages were cocultured with free drugs or nanoparticles at a dose of 10 µM OxPt or/and 50 µM DHA for 24 h, and then injected into the footpads of C57Bl/6 mice. Six days later, popliteal lymph node cells were collected by homogenizing and filtering the organ through a sterile cell strainer. The cell suspension was cultured in culture medium in the presence of MLR medium. Cells were cultured for 24 h and analysed for Pt concentration by ICP-MS. Another 25 µL plasma was added to 5 µL 20% Triton X-100 to disrupt the lipid bilayer of the nanoparticles, chol-DHA was then extracted from plasma by adding 100 µL ethyl acetate, followed by centrifugation at 6700 g for 10 min. The chDHA content was quantified by LC-MS.

Antimurine vaccination. A 1 × 10⁶ MC38 cells treated with 100 µM OxPt or/and 50 µM DHA for 24 h were subcutaneously inoculated into the lower flank of 6-week-old female C57Bl/6 mice or Rag-2−/− mice. Seven days later, 2 × 10⁵ living MC38 cells were inoculated into the contralateral flank. Mice were then monitored for the appearance of tumours for 30 days.

In vivo pharmacokinetics and biodistribution analysis. SD/CD rats were intravenously (i.v.) injected with OxPt/DHA at an OxPt dose of 6 mg/kg (2.14 mg/kg DHA). The blood was collected at 5 min, 30 min, 1 h, 3 h, 5 h, 8 h, 24 h, and 48 h post injection and immediately centrifuged at 604 × g for 10 min to harvest plasma samples. Twenty-five microliters plasma was digested with concentrated nitric acid for 24 h and analysed for Pt concentration by ICP-MS. Another 25 µL plasma was added to 5 µL 20% Triton X-100 to disrupt the lipid bilayer of the nanoparticles, chol-DHA was then extracted from plasma by adding 100 µL ethyl acetate, followed by centrifugation at 6700 g for 10 min. The chDHA content was quantified by LC-MS.

RABL/c mice were subcutaneously injected in the right flank with 1 × 10⁶ CT26 cells. When the tumours reached ~100 mm³, mice were intraperitoneally (i.p.) administrated OxPt/DHA at an OxPt dose of 8 mg/kg (2.86 mg/kg DHA). The blood was collected at 1, 3, 5, 8, 24, 48, and 72 h post injection and immediately centrifuged at 604 × g for 10 min to harvest plasma samples. The content of Pt and chDHA was quantified by ICP-MS and LC-MS, respectively. The livers, lungs, spleens, kidneys, bladders, and tumours were also harvested, digested with concentrated nitric acid for 24 h, and analysed for Pt concentration by ICP-MS.

For surface detection of CRT, CT26 cells were seeded on 10 mm² glass coverslips placed in six-well plates at a density of 2 × 10⁵ cells per well. After treatment, cells were washed with PBS three times, incubated with Alexa Fluor 488-CRT antibody (diluted 1:100) for 2 h, stained with DAPI, and observed under CLSM using 405 nm and 488 nm lasers for visualizing nuclei and CRT expression on the cell membrane, respectively.
In vivo toxicity on mice. Balb/c mice received i.p. doses of Zn/DHA at 5 mg/kg DHA every 3 days for a total of 50 doses. C57Bl/6 mice received a weekly i.p. dose of OxPt/DHA at 60 mg/kg OxPt (21.5 mg/kg DHA) for a total of four doses or a single i.p. dose at 80 mg/kg OxPt (28.6 mg/kg DHA). The activity level and body weights of the mice were monitored for toxicity.

SD/CD rats received weekly i.v. doses of OxPt/DHA or free OxPt at 8 mg/kg OxPt (2.86 mg/kg DHA) for a total of three doses to measure peripheral neuropathy. A hind paw was subjected to a constant heat source through a 3/8” glass pane and tested for the time to withdrawal to measure peripheral neuropathy.

In vivo anticancer efficacy. 1 x 10⁶ cells CT26 or MC38 cells were subcutaneously injected into the right flank region of 6-week BALB/c, C57Bl/6 wild-type or Rag2−/− mice, respectively. Twelve days after tumour inoculation, mice were i.p. dosed with 8 mg/kg OxPt, 2.86 mg/kg DHA, and/or 75 µg PD-L1 antibody once every 3 days for up to 12 doses. Tumour growth was monitored by measurement with a digital caliper, where tumour volumes were calculated as follows: width² x length/2. Tumour-free BALB/c mice were challenged with 5 x 10⁶ CT26 cells on the contralateral flank 3 months after the tumours disappeared. The mice were monitored for 1 month and then rechallenged with 5 x 10⁶ unrelated 4T1 cells.

Immunofluorescence assay. Tumours were collected 2 days or 12 days after the first treatment, and frozen tissue sections of 5 µm thickness were prepared using a cryostat. The sections were fixed in acetone for 10 min at −20°C, blocked with 2% BSA for 1 h, and incubated with individual primary antibodies against CD11c (eBioscience), F4/80 (eBioscience), CD3e (Santa Cruz) or CD8 (Thermo Scientific) overnight at 4°C, followed by incubation with dye-conjugated secondary antibodies for 1 h at room temperature. After staining with DAPI for another 10 min, the sections were washed twice with PBS and observed under CLSM.

Flow cytometry assay for immune response. Tumours were harvested on 12 days after the first treatment, treated with 1 mg/ml collagenase I (Gibco®, USA) for 1 h, and ground with the rubber end of a syringe. Cells were filtered through nylon mesh filters and washed with PBS. The single-cell suspension was incubated with anti-CD16/32 (clone 93; eBiosciences) to reduce nonspecific binding to FcRs. Cells were further stained with the following fluorescence-conjugated antibodies: CD45 (30-F11), CD3e (145-2C11), CD8a (53-6.7), CD11b (M1/70), CD11c (N418), F4/80 (RM4), MHC II (AF6-120.1), CD86 (PO3), CD206 (C068C2), CD44 (IM7), CD62L (MEL-14), and Zombie NIR (all from eBioscience). A Millipore Multiscreen HTS-IP plate was coated with anti-CD16/32 (clone 93; eBiosciences) to reduce nonspecific binding to FcRs. Cells were incubated with primary antibodies for 1 h at room temperature. After staining with DAPI for another 10 min, the sections were washed twice with PBS and observed under CLSM.

ELISPOT assay. Tumour-specific immune responses to IFN-γ was measured in vitro by an ELISPOT assay (Mouse IFN-γ ELISPOT Ready-SET-Go!, Cat. No. 88-7384-88; eBioscience). A Millipore Multiscreen HTS-IP plate was coated overnight at 4°C with anti-Mouse IFN-γ capture antibody (diluted 1:250). Single-cell suspensions of splenocytes were obtained from MC38 tumour-carrying mice on 12 days after the first treatment and seeded onto the antibody-coated plate at a concentration of 2 x 10⁶ cells/well. Cells were incubated with or without KSPWF TTL stimulation (10 µg/mL; in purity 495%; PEPTIDE 2.0) for 48 h at 37°C in 5% CO₂. After 12 days, 100-µL volume of IFN-γ antigenic peptide mixture was added for each 48-well plate (100 µg each per well). After 24 h of incubation, 200 µL/well of AEC substrate solution (Sigma, Cat. AEC101) was added for incubation with Avidin-HRP for 2 h at room temperature. 3-aminio-9-ethylcarbazole (AEC) substrate solution (Sigma, Cat. AEC101) was added for cytokine spot detection.

Reporting summary. Further information on experimental design is available in the Nature Research Reporting Summary linked to this article.

Data availability
The authors declare that all the data supporting the findings of this study are available within the article and its Supplementary Information files or from the corresponding author upon reasonable request. The crystal structure reported is deposited at the Cambridge Crystallographic Data Centre (CCDC) under deposition number CCDC 1875999. The crystallographic file can be obtained free of charge from the Cambridge Crystallographic Data Centre via http://www.ccdc.cam.ac.uk/data_request/cif.

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