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Intraperitoneal Oil Application Causes Local Inflammation with Depletion of Resident Peritoneal Macrophages — Source link

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2	resident peritoneal macrophages
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26 Abstract

27 Oil is frequently used as a solvent to inject lipophilic substances into the peritoneum of laboratory animals. Although mineral oil causes chronic peritoneal inflammation, little is known 28 whether other oils are better suited. Here we show that olive, peanut, corn or mineral oil causes 29 xanthogranulomatous inflammation with depletion of resident peritoneal macrophages. 30 31 However, there were striking differences in the severity of the inflammatory response. Peanut and mineral oil caused severe chronic inflammation with persistent neutrophil and monocyte 32 recruitment, expansion of the vasculature and fibrosis. Corn and olive oil provoked no or only 33 mild signs of chronic inflammation. Mechanistically, the vegetal oils were taken up by 34 35 macrophages leading to foam cell formation and induction of cell death. Olive oil triggered caspase-3 cleavage and apoptosis, which facilitates the resolution of inflammation. Peanut oil 36 and, to a lesser degree, corn oil triggered caspase-1 activation and macrophage pyroptosis. 37 38 which impairs the resolution of inflammation. As such, intraperitoneal oil administration can 39 interfere with the outcome of subsequent experiments. As a proof-of-principle, intraperitoneal peanut oil injection was compared to its oral delivery in a thioglycolate-induced peritonitis 40 model. The chronic peritoneal inflammation due to peanut oil injection impeded the proper 41 recruitment of macrophages and the resolution of inflammation in this peritonitis model. In 42 43 summary, the data indicate that it is advisable to deliver lipophilic substances like tamoxifen by oral gavage instead of intraperitoneal injection. 44

45

46 Introduction

Oil is frequently used as solvent in animal research. For instance, inducible gene recombination using the Cre-ERT2 -loxP system requires administration of tamoxifen which is usually dissolved in olive, peanut, corn or mineral oil. The oil solution is administered orally or by intraperitoneal injection (*i.p.*) (1, 2). Also, in a liver fibrosis model carbon tetrachloride (CCl₄) is delivered by *i.p.* injection in oil, inhalation or oral gavage (3). Interestingly, *i.p.* injection generates stronger liver fibrosis when compared with the other two administration methods (4),

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raising the question whether CCl₄ or its solvent act locally within the peritoneum. Indeed, *i.p.*injection of mineral oil causes chronic inflammation (5-9). Also, subcutaneous injection of olive
oil can cause lipogranuloma, a granulomatous inflammatory soft tissue reaction (10).

Therefore, it can be assumed that any experimental immune cell analysis within the peritoneal cavity would be strongly affected by oil. It is surprising how little is known about the peritoneal immune cell reaction towards oil and comparative studies of different oils are missing to our knowledge.

Peritoneal inflammation can be divided into the initiation and resolution phase. 60 Pathogens trigger infiltration of neutrophils, which phagocytose pathogens, clear apoptotic 61 cells and recruit monocytes from the blood stream into the peritoneal fluid. Recruited 62 monocytes eliminate dying neutrophils and differentiate into monocyte-derived macrophages 63 64 (11). This is important, as the number of resident peritoneal macrophages, which are derived from embryonic progenitors and have self-renewal capacity (12), get strongly decreased as a 65 result of the so-called "macrophage disappearance reaction" (13). As such, resident peritoneal 66 CD11b⁺ macrophages, expressing high F4/80 levels (F4/80^{hi}) get replaced by monocyte-67 68 derived CD11b⁺ macrophages, expressing low F4/80 levels (F4/80^{low}) on the membrane (12, 14, 15). Subsequently, monocyte-derived macrophages increase surface expression of F4/80 69 from a low to an intermediate level (F4/80^{int}) to initiate the resolution phase (16). 70

The switch from inflammatory to resolving macrophages is triggered by phagocytosis of apoptotic cells. Deficiency in this phagocytic process leads to chronic inflammation (17). For instance in atherosclerotic plaques, macrophages take up excessive amounts of lipids and become foam cells, which cannot initiate the resolution phase, perpetuating further neutrophil and monocytes infiltration (18).

The aim of this study was to analyze how the most commonly used oils in animal research affect the myeloid cells within the peritoneum and whether this would diminish their capability to resolve the peritoneal inflammation.

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80 Materials and methods

81 Animal models

The study was approved by institutional and regional animal research committees. All animal procedures were in accordance with institutional guidelines and performed according to the guidelines of the local institution and the local government. Female C57BL/6 mice were grouphoused under specific pathogen-free barrier conditions.

Administration of peanut (P2144, Sigma-Aldrich, St. Louis, USA), corn (C8267, Sigma-Aldrich,

87 St. Louis, USA), olive (88631, Carl Roth, Germany), mineral oil (HP50.2, Carl Roth, Germany)

or 0,9% sterile NaCl (Braun, Germany) in 8 to 12-week-old randomized mice was performed

by daily *i.p.* injection of 100 µl for 5 consecutive days or by oral gavage of peanut oil once with

 100μ l. After three weeks mice were euthanized. For peritoneal lavage, 5 ml of cold PBS

91 (Gibco/Thermo Fisher Scientific, NY, USA) was injected *i.p.* after a careful massage to mobilize

cells, peritoneal fluid was collected. Cells were isolated by centrifugation (5 min, 200 g) and

93 suspended in 1 ml of PBS.

8 to 12-week-old randomized mice were euthanized and administrated with peanut, olive, corn
and mineral oil. After 5 minutes the peritoneal lavage was collected.

Three weeks after oil treatment, mice were *i.p.* injected with thioglycolate (2 mg in 1 ml H₂O; B2551, Sigma Aldrich, St. Louis, USA). After 24 or 72 hours mice were sacrificed and peritoneal lavage collected. All groups were randomized.

99

100 Immunofluorescence and tissue histology

Histological analysis was performed on formalin-fixed paraffin-embedded sections (3 µm).
Sections were deparaffinized and rehydrated. For hematoxylin-eosin (H&E) and Sirius Red
(Dianova, Germany) staining, sections were processed according to standard protocols. For
myeloid cell staining, antigen retrieval at pH 6 with citrate buffer and the primary antibody rabbit
anti-mouse CD11b (1:200) (ab133357, Abcam, Cambridge, MA, USA) and antigen retrieval

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106 with 1:20 proteinase K/TE buffer and rat anti-mouse F4/80 (1:100) (T-2006, Dianova, Germany) incubated at 4°C overnight. After washing, sections were incubated with secondary 107 108 antibodies coupled with HRP (1:200) (DAKO, Agilent Technologies, Santa Clara, CA, USA) for 109 one hour at room temperature. For immunofluorescence staining, antigen retrieval at pH 9 was performed using citrate buffer and sections were incubated with the primary antibody rabbit 110 anti-mouse CD31 (1:50) (ab28364, Abcam, Cambridge, MA, USA) at 4°C overnight. After 111 washing, sections were incubated with secondary antibody (1:200) goat anti-rabbit Alexa 112 Fluor-647 (A21245, Life Technologies/Thermo Fisher Scientific, NY, USA) for 1 hour at room 113 temperature. H&E images were obtained with slide scanner (Zeiss Axio Sacn.Z1, Carl Zeiss, 114 Germany). CD11b images were obtained with widefield microscope (Zeiss Axioplan, Carl 115 116 Zeiss, Germany). All images were processed with ZENblue software (Carl Zeiss, Germany). 117 Immunofluorescence was imaged at the confocal (LSM 700, Carl Zeiss, Germany) microscope 118 with ZENblack software (Carl Zeiss, Germany). Sections of seven Z-stacks per omentum and 119 mesentery and three random fluorescence images per slide were taken. Numbers of CD31 120 positive vessels per view field and lipid droplet size from H&E images were counted with 121 ImageJ software (NIH, Bethesda, MD, USA).

122

123 Oil Red O staining

124 Peritoneal lavage was plated into one well of a 6-well plate on top of coverslips and incubated for 30 min with Dulbecco's modified Eagle's medium (DMEM) (Gibco/ Thermo Fisher Scientific, 125 NY, USA). Afterwards non-adherent cells were removed by careful washing three times with 126 PBS. J774A.1 cells cultured in DMEM with 10% fetal calf serum (Biochrom, UK) were seeded 127 into 12-well plates on coverslips and treated with 100 µl oil in 1 ml medium for four hours. Cells 128 129 on coverslips were stained with Oil Red O (O0625, Sigma-Aldrich) following the protocol published elsewhere (19) and counterstained with hematoxylin. Images were obtained with 130 widefield microscope (Zeiss Axioplan Carl Zeiss, Germany). 131

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133 Flow cytometry

134 Cells obtained from peritoneal lavage were washed and erythrocytes lysed with ACK lysis buffer (Gibco/Thermo Fisher Scientific, NY, USA). Cells were suspended at approximately 10⁶ 135 cells/ml in PBS with 2% FCS. Cell suspensions were incubated with the different fluorophore-136 coupled primary antibodies for 20 minutes on ice. These antibodies were used: CD45 137 138 (552848), CD11b (552850), CD19 (560375), Ly6G (560600), Ly6C (560594) and F4/80-like (564227) all from BD Biosciences (Bedford, MA, USA), CD3 (100203) and F4/80 (123128) 139 from BioLegend (St. Diego, CA, USA) and Tim4 (12-5866-82, Life Technologies/Thermo 140 Fisher Scientific, NY, USA). Concentration of the different antibodies was determined by 141 142 titration. Flow cytometer results in percentage were extrapolated to the total amount of cells 143 obtained from the previous cell counting.

144

145 Western Blot analysis

Cell lysates were separated by SDS-PAGE and proteins blotted on nitrocellulose membranes. 146 Membranes were blocked with 5% skim milk in TBS with 1% Tween-20. The following primary 147 antibodies were used: CD36 (ab124515), VCP (ab11433) from Abcam (Cambridge, MA, USA), 148 149 ABCG1 (NB400-132SS, Novus Biologicals, CO, USA), Cleaved-Caspase 3 (Asp175; 9664S), Arginase-1 (D4E3M[™]; 93668S) from Cell Signaling (Danvers, MA, USA) and Caspase 1 150 (14F468; sc-56036, Santa Cruz Technologies, Dallas, TX, USA). Primary antibodies were 151 incubated overnight at 4°C and appropriate HRP-conjugated secondary antibodies (DAKO, 152 Agilent Technologies, Santa Clara, CA, USA) for 1 hour at room temperature. 153 Chemiluminescence was detected by Pierce ECL Western Blotting Substrate (Thermo Fisher 154 Scientific, NY, USA) and ChemiDoc imaging system (Biorad, Hercules, CA, USA) and 155 quantified with Image Lab 3.0 software (Biorad, Hercules, CA, USA). 156

157

158 **Quantitative PCR**

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RNA was isolated using the innuPREP RNA Mini kit (Analytik Jena, Germany). cDNA was 159 160 synthesized with the High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems). The cDNA was applied to gPCR using the POWER SYBR Green Master Mix (Applied Biosystems). 161 Fold changes were assessed by $2^{-\Delta\Delta Ct}$ method and normalized with the *CPH* gene. The 162 following primers were used for qPCR: CD36 forward GCAAAACGACTGCAGGTCAA and 163 reverse GGCCATCTCTACCATGCCAA, ABCG1 forward CTTTCCTACTCTGTACCCGAGG 164 and reverse CGGGGCATTCCATTGATAAGG, IL10 forward GCATGGCCCAGAAATCAAGG 165 and reverse GAGAAATCGATGACAGCGCC and CPH forward ATGGTCAACCCCACCGTG 166 and reverse TTCTTGCTGTCTTTGGAACTTTGTC. 167

168

169 Cell death detection

J774A.1 cells were plated at 5x10⁵ cells per well into a 12 well-plate with 100 μl of the different
oils in 1 ml medium and incubated for four hours. Afterwards, supernatant and attached cells
were collected and stained with Annexin V-FITC (640905, BioLegend, St. Diego, CA, USA)
and PI (Cayman Chemical, USA) and incubated for 15 minutes on ice. After washing, cells
were immediately analyzed by flow cytometry.

An apoptosis/necrosis immunofluorescence assay kit (ab176749, Abcam, Cambridge, MA, 175 USA.) was used for detection of necrosis or apoptosis cell death. J774A.1 cells were plated at 176 5x10⁵ cells/well into a 24 well-plate on top of a coverslip. To each well, 50 µl of oil was added 177 in a final volume of 1 ml medium and incubated for two hours. Afterwards, the staining was 178 performed following the manufacturer's protocol. Three fluorescence images of each channel 179 at fixed positions of each triplicate were collected at the wide-field Cell Observer microscope 180 (Carl Zeiss, Germany) with ZENblue software (Carl Zeiss, Germany). FIJI software was 181 employed for the quantification of positive cells of each channel per field. 182

For determination of lactate dehydrogenase activity in the cell supernatant, J774A.1 cells were plated at 5×10^4 cells/well into a 96 well-plate with 100 µl of medium containing 10 µl oil in triplicates and incubated for 2 hours. Oleic acid (O1008, Sigma- Aldrich, St. Louis, USA),

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- diluted in absolute ETOH was added to the medium or mixed with 5 µl peanut oil when
- indicated. Then, levels of LDH were detected using the LDH-Cytotoxicity Assay Kit (Ab65393,
- Abcam, Cambridge, MA, USA) following the manufacturer's protocol.

189

190 Statistical analysis

191 GraphPad Prism 8 (GraphPad Software, San Diego, CA, USA) was used to generate graphs 192 and for statistical analysis. Statistical significance was calculated using one-way or two-way 193 ANOVA as indicated in the figure legends. Data sets are presented as mean \pm SD. *P* < 0.05 194 was considered as significant.

195

196

197 Results

198 Macroscopic changes upon intraperitoneal oil injection

Peanut, olive, corn and mineral oil were injected into the peritoneum (*i.p.*) of adult mice for five consecutive days. This mimics a typical protocol for delivering tamoxifen to induce gene recombination in transgenic mice expressing Cre^{ERT2} recombinase (2). Analysis was done three weeks later. As controls, untreated mice and mice treated with peanut oil by oral gavage were used (**Figure 1A**).

In contrast to untreated mice or those receiving oil by oral gavage, the *i.p.* injected mice showed macroscopically visible alterations in the peritoneal cavity. Peanut oil was still visible as oil droplets (**Figure 1B**), whereas this was not the case for the other oils. White nodules, in the size of <3 mm, were visible on the surface of liver, diaphragm or colon in mice receiving peanut, olive and mineral oil *i.p.* but not corn oil. The nodules formed due to olive oil treatment were only loosely attached to the organ surfaces, whereas the nodules in mice that were *i.p.* injected with peanut or mineral oil were firmly attached to the liver surface (**Figure 1B**).

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212 Xanthogranulomatous inflammation in the peritoneum upon oil injection

Histological analysis revealed no pathological changes in liver (**Figure 1C**) and spleen (**Supplementary Figure 1A**) of mice that received *i.p.* injection of oil. The nodules that were firmly attached to the liver surface in mice treated with peanut or mineral oil could be classified as xanthogranulomas with foamy macrophages and mixed inflammatory background (**Figure 1C**).

218 Next, we examined the greater omentum. The greater omentum is an organ that filters excessive fluid from the abdominal cavity, senses microorganisms or damaged cells, initiates 219 immune responses and supports repair of damaged organs (20). Omenta of mice treated with 220 oral gavage were indistinguishable from those of untreated mice. However, omenta of mice 221 that received *i.p.* oil injections showed a remarkable change in morphology. The omenta were 222 223 swollen, darker and had enlarged blood vessels, particularly in mice treated with peanut and mineral oil (Figure 2A). Histological analysis revealed that lipid droplet size in adipocytes was 224 reduced in mice treated *i.p.* with any of the different oils. Again, the changes were most 225 pronounced in mice injected with peanut and mineral oil (Figures 2B and 2F). 226

Immunohistochemical analysis of CD31-positive endothelial cells revealed an increase
in vessel density in the omenta after oil injection. It was strongly increased in the case of peanut
and mineral oil but mild in olive and corn oil-treated mice (Figure 2C and 2G). In addition,
higher numbers of CD11b⁺ myeloid cells were present in all four oil-treated mice, but again,
peanut and mineral oil-treated mice had highest infiltration rates (Figure 2D).

So far, the described changes are indicative of peritoneal inflammation upon local oil injection. Prolonged inflammation may impede tissue healing resulting in organ fibrosis. Indeed, Sirius Red staining revealed an increase in collagen deposition, a typical sign of fibrosis, in the greater omentum of *i.p.* oil-injected mice. Such fibrotic changes were in particular observed in mice treated with peanut and mineral oil (**Figures 2E**).

Histopathological scoring of the inflammation grade in the greater omentum by H&E staining was based on the granularity of the tissue, the presence of foamy macrophages or

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other inflammatory cells, multinucleated giant cells, fibrosis or necrosis with a score from 0 (no
inflammation) to 3 (severe chronic inflammation). This showed that *i.p.* injection of all four oils
causes xanthogranulomatous inflammation of the omentum with highest scores for mineral
and peanut oil. (Figure 2H).

243 Similar data were obtained during the analysis of mesentery. The almost transparent 244 membrane became opaque in mice treated *i.p.* with oil. The strongest changes were observed in mice treated with peanut and mineral oil (Figure 3A). Lipid droplets in adipocytes of 245 mesentery from mineral oil-treated mice were much smaller compared to controls. Such 246 changes were also observed but to a lesser extent in mice injected *i.p.* with peanut oil, whereas 247 248 the effects of olive and corn oil were mild (Figures 3B and 3F). The number of blood vessels were increased in peanut and mineral oil-treated mice. We detected increased numbers of 249 250 CD11b⁺ myeloid cells in the mesentery of olive, corn, peanut and to the maximum extent in mineral oil-treated mice (Figure 3D). There was mild fibrosis in the mesentery of mice treated 251 252 with peanut oil and severe fibrosis in mice that had received mineral oil (Figure 3E). Histopathological scoring of the inflammation grade revealed that peanut and mineral oil, but 253 not olive and corn oil, generated xanthogranulomatous inflammation (Figure 3H). 254

In summary, the histopathological analysis revealed that mineral oil and peanut oil induce a strong xanthogranulomatous inflammatory response in the peritoneum. Olive and corn oils also induce inflammation, but to a much lesser degree.

258

259 Intraperitoneal oil injection causes myeloid cell infiltration into the peritoneum

To further analyze the immune response, peritoneal lavage was obtained three weeks after the *i.p.* injection. Flow cytometry revealed that the total cell number in the peritoneal lavage was significantly increased in mice treated with either peanut or mineral oil compared to untreated animals (**Figure 4A**). The vast majority of the cell population were myeloid cells (CD45⁺CD19⁻CD11b⁺). Peanut and mineral oil increased the total number of myeloid cells,

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whereas olive and corn oil had no significant effect (Figure 4B and Supplementary Figure
266 2A).

We also observed that oil injection led to a decrease in the number of B (CD45⁺CD19⁺CD3⁻) and T lymphocytes (CD45⁺CD19⁻CD3⁺) (**Supplementary Figure 2B**). This was expected as during peritoneal inflammation lymphocytes migrate from the peritoneal fluid into the greater omentum (21, 22).

271

272 Peanut and mineral oil increases neutrophil and monocyte recruitment

A more detailed evaluation of the myeloid population revealed that peanut and mineral oil injection strongly increased the presence of neutrophils (CD45⁺CD11b⁺Ly6G⁺Ly6C^{int}) and recruited monocytes (CD45⁺CD11b⁺Ly6G⁻Ly6C⁺) in the peritoneal fluid. Neutrophils and infiltrated monocytes were almost absent in peritoneal lavage derived from untreated mice or mice treated with olive oil, corn oil or oral gavage (**Figures 4C, D and E**).

Importantly, the injection itself did not cause such alterations. Injection of 0.9% NaCl
did not lead to macroscopic or histological changes, nor to significant changes in total number
of cells in peritoneal fluid or changes within the myeloid cell compartment (Supplementary
Figure 3A-G).

282

283 Oil injection leads to a severe reduction of resident peritoneal macrophages

During inflammation, neutrophils are the first cells being recruited to clear apoptotic cells or eliminate pathogens. Afterwards, monocytes reach the inflamed zone to eliminate dying neutrophils and to differentiate into macrophages. The latter is in particular essential when resident macrophages are eradicated. Therefore, we next examined the macrophage population within the peritoneum. The total macrophage (CD45⁺CD11b⁺F4/80⁺) cell number was not significantly changed in peritoneal lavage of mice treated *i.p.* with oil compared to the untreated mice or to those which received oil by oral gavage (**Figure 4F**).

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We further characterized the F4/80 population by analyzing the amount of Tim4 on the cell surface as Tim4 can be employed as a marker to differentiate long-term (F4/80⁺Tim4⁺) from newly recruited (F4/80⁺Tim4⁻) resident macrophages (23). This revealed that all four oils led to a dramatic decrease in long-term resident (F4/80⁺Tim4⁺) macrophages and a replacement by recently recruited (F4/80⁺Tim4⁻) macrophages (**Figure 4G**).

The level of F4/80 on the macrophage cell membrane varies depending on the differentiation stage (15). In this regard, resident macrophages express high levels of F4/80, while newly recruited monocyte-derived macrophages express low to intermediate levels (F4/80^{int}). Peanut oil injection led to a strong decrease in resident F4/80^{hi} macrophage numbers (**Figure 4H**). Mineral oil showed a similar but not significant trend, whereas olive oil and corn oil did not alter the proportion of F4/80^{hi} macrophages.

The full resolution of inflammation is carried out by F4/80^{int} macrophages (16). Analysis of this cell population revealed that only the *i.p.* injection of olive oil led to a significant increase in F4/80^{int} macrophages at this time point (**Figure 4I**).

305 Collectively, the data imply that injection of any oil into the peritoneum triggers an 306 inflammatory response in which resident macrophages get replaced by monocyte-derived 307 ones. However, the resolution of inflammation depends on the type of oil, with olive oil and 308 corn oil (to a lesser extent) showing signs of resolution.

309

310 Oil injection induces foam cell formation

Monocytes and macrophages can take up excessive amounts of lipids (24). Therefore, we examined lipid uptake in macrophages upon oil injection. Peritoneal lavage was performed three weeks after *i.p.* oil injection. Adherent peritoneal macrophages were stained with Oil Red O, which marks lipids and neutral triglycerides. Macrophages derived from the control mice were not stained by Oil Red O while peritoneal macrophages derived from mice injected *i.p.* with peanut oil contained multiple large lipid droplets. Fewer amounts were detected in the

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macrophages from the olive oil and corn oil group, whereas the macrophages of the mineral
oil group contained almost no detectable lipid droplets (Figure 5A).

To further evaluate this, we tested lipid uptake in the J774A.1 macrophage cell line. J774A.1 macrophages took up lipids when in contact with olive, corn and peanut oil, but not mineral oil, suggesting that the effect of this oil is independent of the cellular lipid uptake.

In atherosclerotic plaques, monocyte-derived macrophages endocytose lipids such as oxidized LDL and become foam cells. During this transition, an upregulation of the fatty acid translocase (CD36) expression and downregulation of the cholesterol transporter ABCG1 is evident (25). J774A.1 macrophages showed the same changes in gene expression when cultured for four hours in the presence of vegetal oils (**Figure 5C-F**).

In summary, these results indicate that peanut, olive and corn, but not mineral oil *i.p.*injection leads to foam cell formation.

329

330 Peritoneal macrophage cell death after exposure to different oils

Lipoprotein uptake can cause macrophage cell death (26). Therefore, we evaluated 331 whether peritoneal macrophage cell death is induced by the four different oils. Mice were 332 injected once *i.p.* with 100 µl oil and five minutes later peritoneal cells were harvested and 333 subjected to flow cytometry (Figure 6A). This revealed that compared to untreated mice there 334 was approximately a 50% decrease in CD11b+F4/80+ macrophages in the peritoneum of mice 335 that received any of the oils (Figure 6B). There was a 5-10% decrease in the fraction of live 336 337 cells (Annexin V⁻, PI⁻) and an equivalent increase of necrotic (PI⁺), apoptotic (Annexin V⁺) and 338 Annexin V⁺PI⁺ cells in the peritoneal lavage of mice that received peanut or olive oil (Figure **6C**). The Annexin V^+PI^+ double-positive population can be the result of both, apoptosis and 339 necrosis (27). The increase in cell death was milder in the presence of corn oil, whereas in 340 mineral oil injected mice cell death was not different as compared to untreated mice, 341 342 suggesting that the mechanism for mineral oil-induced injury is different (Figure 6C).

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To further analyze cell death in macrophages, J774A.1 cells were treated with different oils. All three vegetal oils increased cell death. In this case, the increase in Annexin V⁺, PI⁺ double positive cells was present for olive, corn and peanut oil. Moreover, necrotic cell death (Annexin V⁻, PI⁺) was also increased in the presence of peanut and corn oil (**Figure 6D**).

In order to further clarify whether macrophages die by apoptosis or necrosis we incubated J774A.1 cells with different oils to determine apoptotic and necrotic cell death. There was an increase in apoptotic cells in the case of incubation with olive oil and a mild increase in the presence of corn oil. 7-AAD incorporation (necrosis) was increased upon treatment with peanut and to a lesser extent upon treatment with corn and mineral oil (**Figure 6E-G**).

Another way to detect necrotic cell death is measuring lactate dehydrogenase (LDH) activity in the cell culture supernatant. Membrane disruption of necrotic cells allows release of cytosolic LDH. Peanut oil caused pronounced release of LDH. There was also LDH release from macrophages treated with olive and corn oil, however to a lesser degree (**Figure 6H**). Interestingly, the LDH release upon treatment with peanut oil could be strongly decreased by supplementing peanut oil with the polyunsaturated oleic acid (**Supplemental Figure 4A**).

We corroborated the different mechanisms of cell death further and observed that only treatment with olive oil induces cleavage of caspase-3 in macrophages, a major effector of apoptosis (**Figure 6J**). On the other hand, peanut oil-treated macrophages showed an increase in active p20 caspase-1, which is a marker for pyroptosis (**Figure 6J-K**), which has similar features as necrosis but is driven by caspase-1 activation (28). In contrast, olive oiltreated macrophages showed increased levels of IL10 and Arginase-1 indicating an antiinflammatory switch towards resolution of inflammation (**Figure 6I-K**).

These results suggest that macrophages in contact with olive oil die by apoptosis, which facilitates the subsequent resolution of the inflammation, whereas peanut oil induces macrophage pyroptosis, which impairs the resolution of inflammation.

368

369 *I.p.* injection of peanut oil impairs the resolution of inflammation in a peritonitis model

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The results presented indicate that *i.p.* oil injection leads to a dramatic change in the peritoneal immune cell composition. Chronic inflammation is induced by peanut oil and this would potentially alter the outcome of experiments executed subsequently. One such example could be a peritonitis experiment in transgenic mice that had been injected *i.p.* before with tamoxifen in peanut oil to induce gene recombination. We decided to test this in experimental peritonitis model, in which mice received peanut oil by *i.p.* injection or by oral gavage as control. Three weeks later, thioglycolate was applied to mimic bacterial peritonitis (**Figure 7A**).

It is known that thioglycolate initially induces a massive neutrophil and monocyte 377 378 infiltration, followed by differentiation into macrophages that resolve inflammation by clearance 379 of apoptotic cells (16). Consistently, mice that had received peanut oil by oral gavage had approximately 50% increase in myeloid cell numbers 24 and 72 hours after thioglycolate 380 injection. However, mice pretreated *i.p.* with oil, already had high numbers of CD45⁺CD11b⁺ 381 382 myeloid cells in peritoneal fluid at baseline and this was not further increased upon 383 thioglycolate administration (Figure 7C). In mice treated by oral gavage, the number of monocytes and neutrophils in peritoneal fluid increased strongly upon thioglycolate injection 384 and subsequently returned below baseline. This suggests that the first inflammatory response 385 by these cells had already been cleared. However, in mice that had been *i.p.* injected with 386 387 peanut oil there was a higher proportion of monocytes and neutrophils already under basal conditions, which was maintained after thioglycolate administration (Figure 7B and D-E). 388

389 Mice treated orally showed the expected increase in CD45⁺CD11b⁺F4/80⁺ 390 macrophages 72 hours after thioglycolate injection. However, mice that had been injected *i.p.* 391 with peanut oil had only few macrophages present in the peritoneum (approximately 12% of 392 all myeloid cells) and these increased only marginally (Figure 7F). Orally treated mice showed 393 disappearance of F4/80^{hi} macrophages 24 hours after thioglycolate administration and 394 subsequent recovery, which was accompanied by an increase of F4/80^{int} macrophages 395 (Figure 7G-H). This suggests that resident macrophages disappear after thioglycolate administration and get replaced by monocyte-derived macrophages as the inflammation 396 resolves. Yet, in mice that received peanut oil *i.p.* there were only few F4/80^{hi} macrophages at 397

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baseline and there was only a minor increase in F4/80^{int} macrophages (**Figure 7G-H**). The lower presence of F4/80^{int} macrophages, together with the continuous influx of monocytes and neutrophils, suggests that resolution of inflammation cannot take place. As such, *i.p.* peanut oil injection leads to a dramatic change in the myeloid cell composition of the peritoneum that affects the outcome of subsequent experiments.

403

404 Discussion

Animal experimentation requires careful planning and analysis to allow reproducibility 405 and the possibility to translate basic research into successful clinical trials. Oil injection is 406 407 frequently performed in animal research, in particular to deliver tamoxifen for inducible gene 408 recombination (1, 2, 4). Oil is considered to be safe and non-toxic. However, few studies 409 reported peritoneal inflammation after subcutaneous or intraperitoneal oil injection (6, 10, 29). 410 To our knowledge, little is still known about changes in the immune cell composition and related 411 reactions within the peritoneum upon intraperitoneal oil delivery. This work demonstrates that 412 intraperitoneal injection of four different oils causes inflammation, foam cell formation and 413 depletion of resident macrophages. However, the severity of inflammation strongly depends on the type of oil. 414

Within the peritoneum, the omentum plays a major role in recognition and 415 encapsulation of pathogens (30). During this process, it expands, a feature which we observed 416 after injection of the different oils. Interestingly, the applied oils were not completely resorbed 417 even three weeks after injection into the peritoneal cavity. In particular larger amounts of 418 peanut oil were still visible in the peritoneal fluid. The failed clearance can be assumed to 419 420 prolong the phase of acute inflammation (17). Consistently, chronic xanthogranulomatous inflammation and fibrosis were observed, particularly upon peanut and mineral oil treatment. 421 422 Myeloid cell infiltration and fibrosis were more severe in the omentum compared to the mesentery. This is consistent with the fact that the omentum is an immunological niche and 423

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the first organ to react against pathogens, but only when this inflammation becomes chronic itstarts to affect the mesentery (20).

Mechanistically, the type of oil-induced macrophage cell death appears to determine 426 whether inflammation gets resolved. For successful resolution of the inflammation, there is the 427 428 need of efferocytosis, where macrophages engulf apoptotic cells (31). Non-resolving 429 inflammation contributes substantially to the progression of atherosclerotic plagues and other chronic inflammatory diseases (18, 32, 33). Our data indicate that macrophages in contact with 430 olive oil die by apoptosis, which facilitates efferocytosis-mediated resolution of inflammation. 431 Conversely, peanut oil induces pyroptosis of macrophages. Excessive pyroptosis impairs the 432 433 resolution of inflammation (34, 35). As such, peanut oil injection results in chronic peritoneal 434 inflammation, whereas olive oil induces macrophage apoptosis, followed by efferocytosis and initiation of the resolution phase. 435

At the cellular level, all three vegetal oils were taken up by macrophages and caused foam cell formation. This change in the expression pattern has been observed in peritoneal macrophages isolated from obese mice, and blood monocytes from patients suffering from severe atherosclerosis (36-38). In principle, one could even use intraperitoneal peanut oil injection as a fast model to obtain viable foam cells for *in vitro* experiments. Future research will determine the potential of this model.

In conclusion, our study shows that intraperitoneal injection of different oils causes peritoneal inflammation and depletion of resident peritoneal macrophages. Whereas olive oil triggers macrophage apoptosis and resolution of inflammation, peanut oil induces pyroptosis and chronic non-resolved inflammation. This has important consequences for animal experiments. In a proof-of-principle approach, we demonstrated this in a thioglycolate-induced peritonitis model after peanut oil injection. To overcome such limitations, it is advisable to deliver lipophilic substances like tamoxifen by oral gavage instead of intraperitoneal injection.

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455

456 **Conflict of interest**

457 The authors declare that they have no conflict of interest.

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1 Figure legends

Figure 1. Macroscopic changes in mice three weeks after *i.p.* oil injection. A. Schematic illustration of the *i.p.* or oral oil administration protocol. B. Representative images of the peritoneum. Black arrows indicate visible lipid droplets. White arrowheads mark nodules on the surface of organs. Scale bar, 3 mm. C. Representative microscopic images of liver sections stained with H&E. Xanthogranuloma on the liver surface in mice treated with peanut and mineral oil. Scale bar, 50µm.

8

9 Figure 2. Xanthogranulomatous inflammation in the omentum upon oil injection.

10 A. Representative images of omenta from untreated mice and such that were treated with oil 11 by *i.p.* injection or oral gavage. Analysis was performed three weeks after treatment. **B.** 12 Representative confocal images of mesentery sections stained with H&E. Scale bar, 50 µm. 13 C. Immunofluorescence microscopy to detect CD31⁺ endothelial cells (white). Scale bar 20 14 μm. **D**. CD11b⁺ myeloid cells. Scale bar, 100 μm. **E** Sirius Red staining to detect fibrosis. Scale 15 bar, 100 µm. F. Lipid droplet size quantification from H&E images. Untreated micen=3, peanut 16 oil n=4, olive oil n=3, corn oil n=5, mineral oil n=5 and oral gavage n=3. Bar graphs show 17 mean ± SD, *, p<0.05 (one way ANOVA). G. Quantification of microvessel density. Olive oil n=4 and all other groups n=5. Bar graphs show mean \pm SD, *, p<0.05 (one way ANOVA). **H**. 18 19 Xanthogranulomatous inflammation score. 0: no inflammation to 3: very strong inflammation. All data represent n=5 mice, bar graphs show mean \pm SD, *, p<0.05 (one way ANOVA). 20

21

Figure 3. Xanthogranulomatous inflammation in the mesentery upon peanut and mineral oil injection. A. Representative images of mesentery from untreated mice and such that were treated with oil by *i.p.* injection or oral gavage. Analysis was performed three weeks after treatment. B. Representative confocal images of mesentery sections stained with H&E. Scale bar, 50 µm. C. Immunofluorescence microscopy to detect CD31⁺ endothelial cells (white). Scale bar, 20 µm D. CD11b⁺ myeloid cells. Scale bar, 100 µm E Sirius Red staining to

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detect fibrosis. Scale bar, 100 µm F. Lipid droplet size quantification from H&E images. n=3
mice. Bar graph shows mean ± SD, *, p<0.05 (one way ANOVA). G. Quantification of
microvessel density. n=5 mice. Bar graph shows mean ± SD, *, p<0.05 (one way ANOVA). H.
Xanthogranulomatous inflammation score. 0: no inflammation to 3: very strong inflammation.
Olive oil n=4 and all other groups n=5. Bar graph shows mean ± SD, *, p<0.05 (one way
ANOVA).

34

Figure 4. Resident peritoneal macrophage depletion and persistent monocyte and 35 neutrophil infiltration upon peanut and mineral oil injection. Peritoneal lavage (PL) was 36 obtained three weeks after treatment with oil. A. Total cell number in peritoneal lavage B. 37 Myeloid cells (CD45⁺CD11b⁺) С. Neutrophils 38 in peritoneal lavage. 39 (CD45⁺CD11b⁺Lv6G⁺Lv6C^{int}) in peritoneal lavage. **D**. Monocytes (CD45⁺CD11b⁺Lv6G⁻Lv6C⁺) in peritoneal lavage. F. Macrophages (CD45+CD11b+F4/80+) in peritoneal lavage. E. 40 Representative blots of flow cytometry analysis of monocytes, neutrophils and macrophages. 41 G. Percentage of CD45⁺CD11b⁺F4/80⁺Tim4⁺ and CD45⁺CD11b⁺F4/80⁺Tim4⁻ macrophages. H. 42 Percentage of CD45⁺CD11b⁺F4/80^{hi} macrophages. I. Percentage of CD45⁺CD11b⁺F4/80^{int} 43 macrophages. All data represent n=4 mice for untreated, peanut, corn and mineral groups and 44 45 n=5 for olive and oral gavage groups. Bar graphs show mean \pm SD, *, p<0.05 (one way 46 ANOVA).

47

Figure 5. Macrophage foam cell formation upon contact with vegetal oil. A. Isolation of
peritoneal macrophages from untreated mice and three weeks after peanut, olive, corn,
mineral oil intraperitoneal injection. Cells from peritoneal lavage were plated for 30 minutes
and stained with Oil Red O. Scale bar, 5 μm. B. J774A.1 macrophages after four hours in
contact with the different oils. Representative images of Oil Red O staining. Scale bar, 50 μm.
C. Representative Western blot of CD36 expression in J774A.1 macrophages after four hours
in contact with the different oils. D. Representative Western blot of ABCG1 expression in

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J774A.1 macrophages after four hours in contact with the different oils. **E.** Quantification of CD36 and ABCG1 mRNA expression levels expression in J774A.1 macrophages after four hours in contact with the different oils. Fold change in comparison to untreated cells. All data from n=3 biological replicates, bar graphs represent mean \pm SD, *, p<0.05 (one way ANOVA).

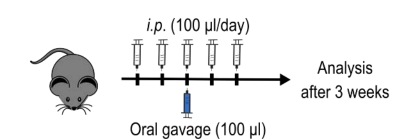
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60 Figure 6. Macrophage cell death upon exposure to vegetal oil. A. Schematic illustration of i.p. oil administration. Peritoneal lavage was obtained five minutes later. B. Percentage of 61 macrophages (CD45⁺CD11b⁺F4/80⁺). Untreated mice n=3, peanut oil n=6 mice and all other 62 groups n=5 mice, mean ± SD, *, p<0.05 (one way ANOVA). C. Percentage of live (Annexin V⁻ 63 PI⁻), apoptotic (Annexin V⁺PI⁻), necrotic (Annexin V⁻PI⁺) or double positive (Annexin V⁺PI⁺) cells 64 65 from total number of cells in peritoneal lavage. Untreated mice n=6, peanut and olive oil n=9 66 mice, corn and mineral oil n=10 mice, bar graphs represent mean \pm SD, *, p<0.05 (2-way ANOVA). D. J774A.1 macrophage cell line untreated or incubated with different oils for four 67 hours and analysis of the percentage of live (Annexin V⁻PI⁻), apoptotic (Annexin V⁺PI⁻), necrotic 68 (Annexin V⁻PI⁺) or double positive (Annexin V⁺PI⁺) cells. n=3 biological replicates, bar graphs 69 70 represent mean ± SD, *, p<0.05 (2-way ANOVA). E. J774A.1 macrophages untreated or 71 incubated with different oils for two hours. Quantification positive cells per field in comparison to untreated, blue (alive cells), green (apoptotic) and red (necrotic). n=3 biological replicates, 72 73 mean ± SD, *, p<0.05 (one-way ANOVA). H. LDH activity in the supernatant upon treatment 74 of J774A.1 macrophages for two hours. n=3 biological replicates, mean \pm SD, *, p<0.05 (oneway ANOVA). I. Normalized mRNA expression levels of IL10. n=4 biological replicates, 75 mean ± SD, *, p<0.05 (one-way ANOVA). J. Representative Western blot cleaved caspase-3. 76 arginase-1 and active p20 caspase-1 of J774A.1 macrophages untreated or incubated for four 77 78 hours with different oils. K. Quantification of Western blots. n=3 biological replicates, 79 mean ± SD, *, p<0.05 (one-way ANOVA).

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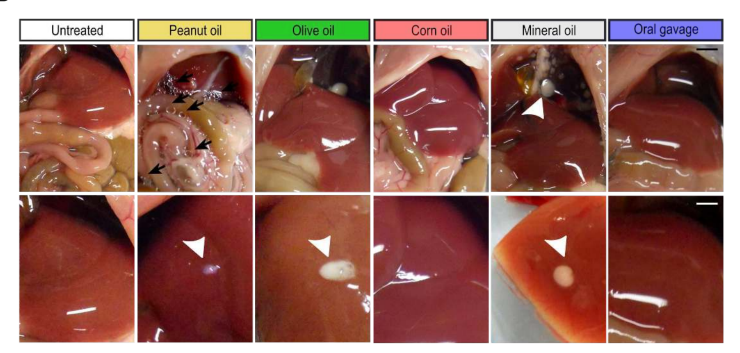
Figure 7. Intraperitoneal injection of peanut oil impairs the resolution of inflammation in

a thioglycolate-induced peritonitis model. A. Schematic illustration of *i.p.* or oral oil 82 administration followed by thioglicolate *i.p.* injection three weeks after treatment with oil. B. 83 84 Representative blots of flow cytometry analysis of monocytes, neutrophils. C. Percentage of myeloid cells (CD45⁺CD11b⁺) in peritoneal lavage. **D**. Percentage of neutrophils 85 (CD45⁺CD11b⁺Ly6G⁺Ly6C^{int}) in peritoneal lavage. **E**. Percentage of monocytes in peritoneal 86 lavage. F. Percentage of macrophages (CD45+CD11b+F4/80+) in peritoneal lavage. G. 87 Percentage of macrophages CD45⁺CD11b⁺F4/80^{hi} in peritoneal lavage. I. Percentage of 88 macrophages CD45⁺CD11b⁺F4/80ⁱⁿ in peritoneal lavage. n=3 mice for oral gavage followed 89 by thioglycolate and analysis at 24 hours. All other groups n=4 mice. 90



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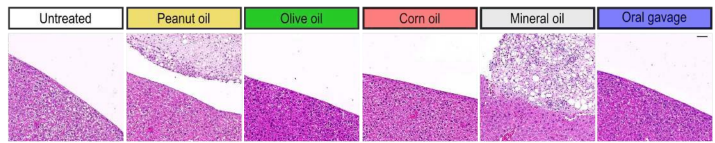
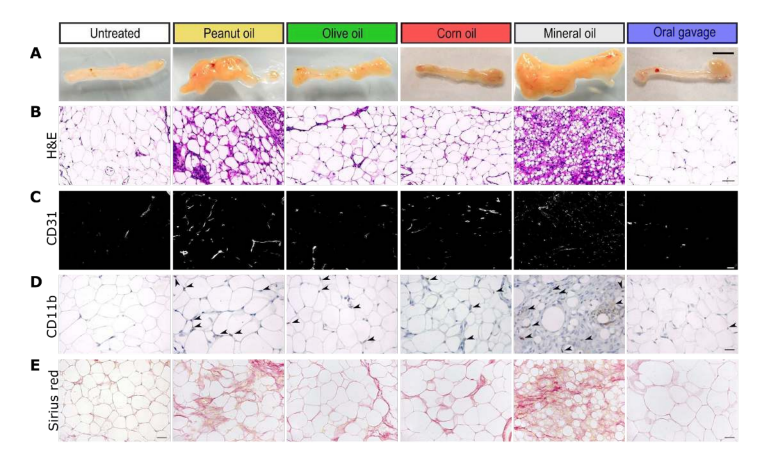
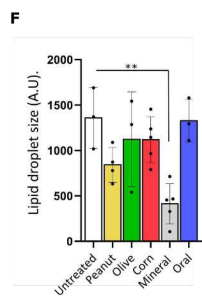
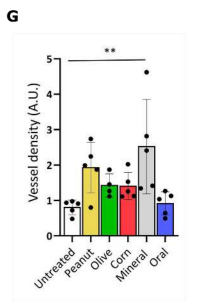


Figure 1







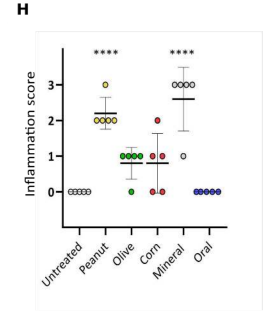
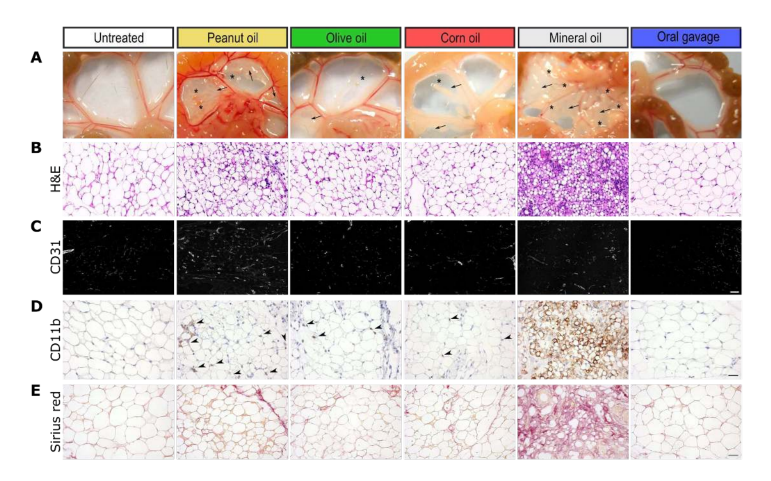
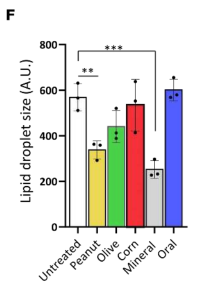
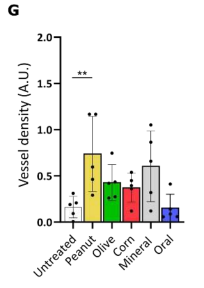


Figure 2







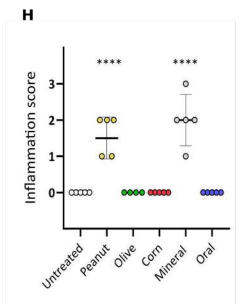


Figure 3

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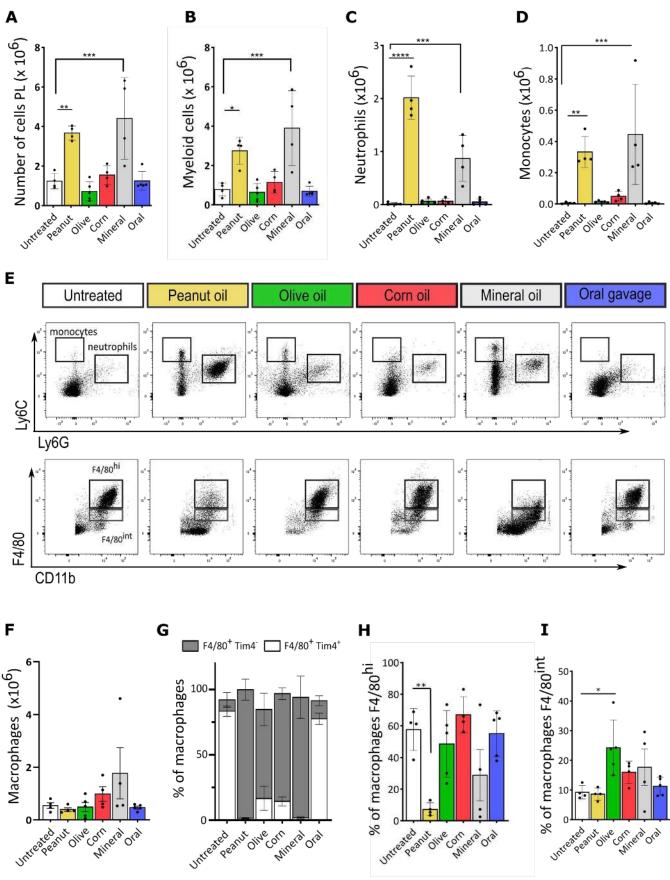
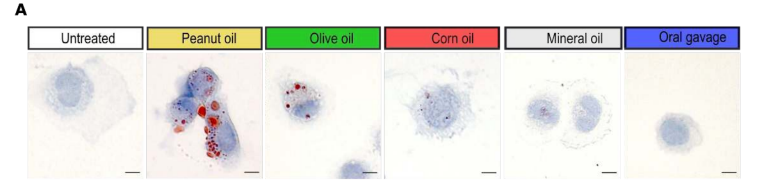
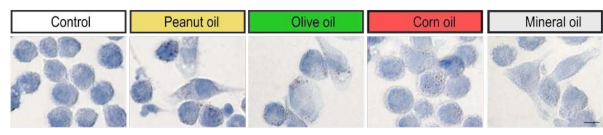
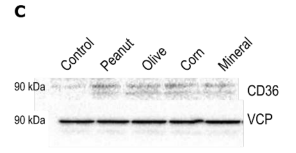


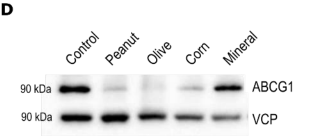
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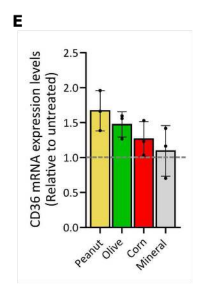


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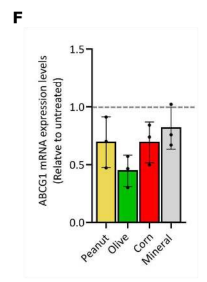
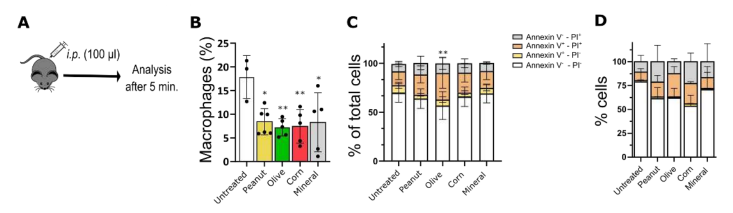
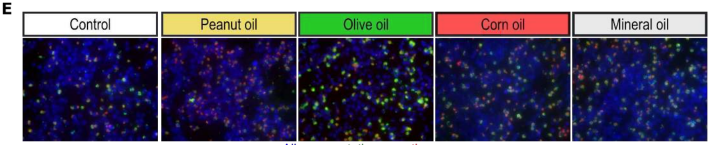


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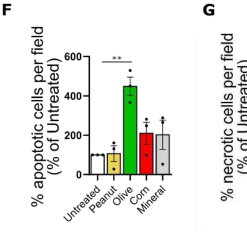


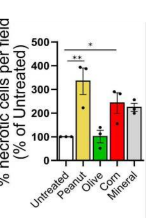


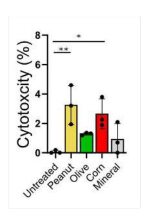
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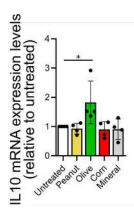
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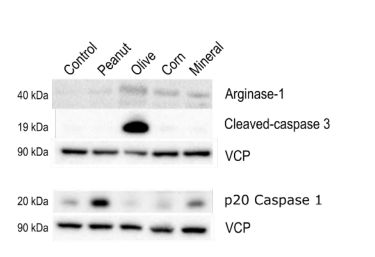


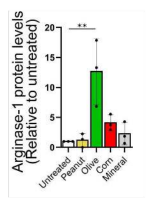




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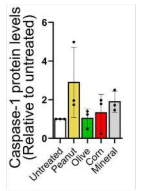


Figure 6.

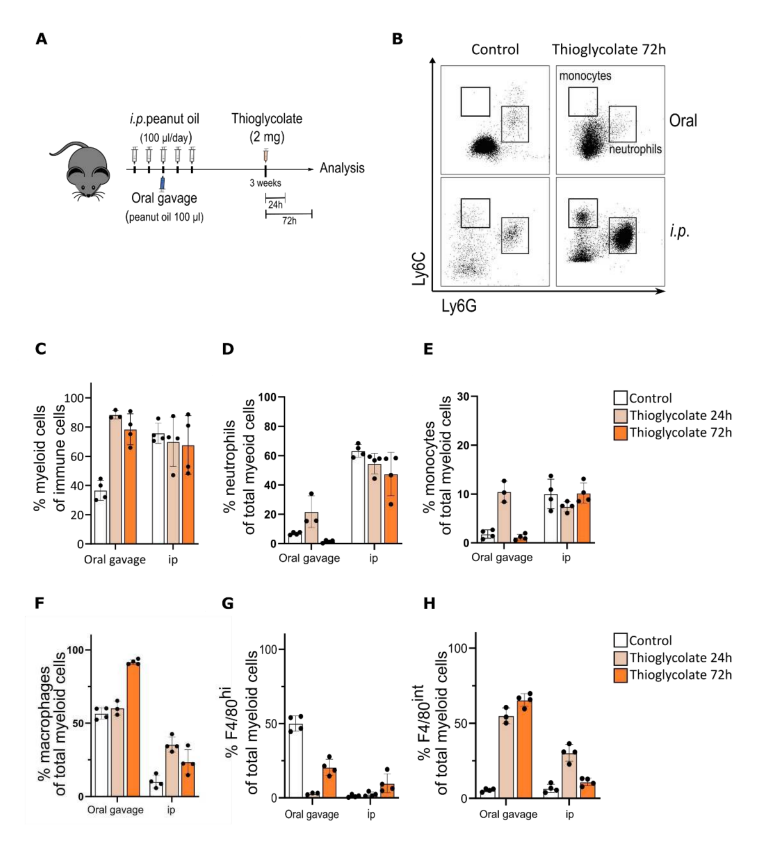


Figure 7.