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1	Evaluation of air oxidized PAPC: a multi laboratory study by LC-MS/MS.
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21	\mathbf{Q}
22	Keywords
23	Oxidized phospholipids, PAPC, reverse phase chromatography, tandem mass spectrometry,
24	multi-laboratory study
25	
26	Abbreviation
27	CID: collision-induced dissociation; CV: coefficient of variation; DAMP: damage associated
28	molecular pattern; DMPC: 1,2-dimyristoyl- <i>sn</i> -glycero-3-phosphocholine; HCD: high-energy
29	collisional dissociation; HOOAPC: 1-palmitoyl-2-(5-hydroxy-8-oxo-octenoyl)-sn-glycero-3-
30	phosphorylcholine; LC-MS/MS: Liquid chromatography coupled on-line to tandem mass
31	spectrometry; LPC: lysophosphatidylcholine; LPP: lipid peroxidation product; oxLDL:
32	oxidized LDL; oxPL: oxidized phospholipid; PAPC: 1-palmitoyl-2-arachidonoyl-sn-glycero-

33 3-phosphocholine; PC: phosphatidylcholine; PEIPC: 1-palmitoyl-2-epoxyisoprostaneE2-*sn*34 glycero-3-phosphocholine; PGPC: 1-palmitoyl-2-glutaryl-*sn*-glycero-3-phosphocholine;
35 POVPC: 1-palmitoyl-2-(5-oxovaleroyl)-*sn*-glycero-3-phosphorylcholine; PL: phospholipid;
36 TS: truncation score; XIC: Extracted ion chromatograms.

37

38 Abstract

39 Oxidized LDL (oxLDL) has been shown to play a crucial role in the onset and development of cardiovascular disorders. The study of oxLDL, as an initiator of 40 inflammatory cascades, led to the discovery of a variety of oxidized phospholipids (oxPLs) 41 responsible for pro-inflammatory actions. Oxidized 1-palmitoyl-2-arachidonoyl-sn-glycero-3-42 phosphocholine (PAPC) is frequently used by the scientific community as a representative 43 oxPL mixture to study the biological effects of oxidized lipids, due to the high abundance of 44 PAPC in human tissues and the biological activities of oxidized arachidonic acids derivatives. 45 Most studies focusing on oxPAPC effects rely on in-house prepared mixtures of oxidized 46 species obtained by exposing PAPC to air oxidation. Here, we described a multi-laboratory 47 evaluation of the compounds in oxPAPC by LC-MS/MS, focusing on the identification and 48 relative quantification of the lipid peroxidation products (LPPs) formed. PAPC was air-49 oxidized in four laboratories using the same protocol for 0, 48, and 72 hours. It was possible 50 to identify 55 different LPPs with unique elemental composition and characterize different 51 structural isomeric species within these. The study showed good intra-sample reproducibility 52 and similar qualitative patterns of oxidation, as the most abundant LPPs were essentially the 53 same between the four laboratories. However, there were substantial differences in the extent 54 of oxidation, i.e. the amount of LPPs relative to unmodified PAPC, at specific time points. 55 This shows the importance of characterizing air-oxidized PAPC preparations before using 56 them for testing biological effects of oxidized lipids, and may explain some variability of 57 effects reported in the literature. 58

59 Graphical Abstract

60



63 1. Introduction

The role of oxidized LDL as an initiator of inflammatory cascades that contribute to the 64 pathogenesis of cardiovascular disorders including atherosclerosis, led to the discovery of a 65 variety of oxidized phospholipids (oxPLs) important for the fine tuning of innate and 66 adaptive immune responses [1–3]. OxPLs formed via enzymatic or free-radical driven 67 reactions are closely associated with systemic redox imbalance and low-grade chronic 68 69 inflammation, and thus have been intensively studied from structural, chemical, biophysical, and biological perspectives over the last decades [4–8]. Within the generic classification of 70 oxPL, the oxidation products of 1-palmitoyl-2-arachidonoyl-sn-glycero-3-phosphocholine 71 (PAPC), which have been identified as some of the bioactive components in minimally 72 modified LDL, are the oxPLs most often used to study the biological effects of oxidized 73 lipids [9]. Furthermore, PAPC is among the most abundant PLs in human tissues [10]. 74 Arachidonic acid, present in the *sn*-2 position of PAPC, can form a very diverse range of 75 76 oxidation products, many of which have already been shown to be bioactive in their free fatty acid forms. Indeed, numerous studies have demonstrated the pro- and anti-inflammatory 77 properties of, for example, eicosanoids, which are generated from arachidonic acid via action 78 79 of dedicated enzymes or free radical driven lipid peroxidation [11–13]. However, the action 80 of PC-esterified eicosanoids has been relatively less well studied.

A variety of bioactivities have been attributed to oxidized PAPC (oxPAPC). It has been 81 identified as a major pro-atherogenic factor and pro-inflammatory mediator [14,15], but in 82 contrast anti-inflammatory properties of oxPAPC have also been demonstrated [13,16]. 83 84 OxPAPC has been shown to induce both the disruption as well as enhancement of the endothelial barrier [17,18], and recently a pro-algesic effect of oxPAPC, via stimulation of 85 transient receptor potential channels TRPA1 and TRPV1, was demonstrated [19,20]. 86 OxPAPC was also shown to act as a damage associated molecular pattern (DAMP), 87 88 activating classical and non-classical pattern recognition receptors. Thus, oxPAPC released from injured tissues was shown to be recognized by murine caspase-11 and CD14 leading to 89 inflammasome activation and release of pro-inflammatory cytokines, and this pathway 90 demonstrated differential regulation in macrophages and dendritic cells [21–23]. The variety 91 of biological effects of oxPAPC are well established and their conflicting properties have 92 been discussed recently [24-26]. 93

One of the main obstacles in elucidating the effects oxPLs on biological systems is the almost complete lack of commercially available and chemically characterized standards for each individual oxPL product. Thus, to study their biological properties, a mixture of

products, usually referred to as "oxPAPC" is often used. The majority of the studies on 97 oxPAPC effects rely on commercial or in-house preparations, usually obtained by exposing a 98 dried film of PAPC to air oxidation for up to 72 h. Experimental details such as room 99 temperature, light exposure, thickness of the phospholipid film and oxPAPC quality control 100 101 are usually not provided. Given the variety of biological functions attributed to oxPAPC in different cellular and *in vivo* models, which may be dependent on variations in sample 102 103 preparation affecting the precise composition of the oxPAPC, we performed a multilaboratory study with the aim of evaluating the reproducibility of PAPC oxidation upon air 104 exposure for 48 and 72 h. Liquid chromatography coupled on-line to tandem mass 105 spectrometry (LC-MS/MS) was used to identify lipid peroxidation products (LPPs) formed 106 upon PAPC oxidation, and the relative quantities of LPPs formed were compared between 107 samples generated independently in four different international laboratories. 108

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110 2. <u>Materials and Methods</u>

111

2.1. Materials

1-palmitoyl-2-arachidonoyl-sn-glycero-3-phosphocholine 1,2-dimyristoyl-sn-112 and glycero-3-phosphocholine were purchased from Avanti Polar Lipids (Avanti Polar Lipids, 113 Inc., Alabama, USA). In laboratory 1 (L1) acetonitrile, isopropanol, water, methanol, 114 ammonium formate (Optima LC-MS grade), chloroform and formic acid (LC-MS grade) 115 were obtained from Fisher Scientific (Loughborough, UK). In laboratory 2 (L2) HPLC-MS 116 grade chloroform, methanol, acetonitrile and isopropanol were purchased from Fisher 117 118 Scientific (Leicestershire, UK). Formic acid was from Honeywell Fluka (Neu Wulmstorf, Germany). Ammonium formate was obtained from Sigma-Aldrich (Sigma-Aldrich, Munich, 119 Germany). The water was of Milli-Q purity (Synergy1, Millipore Corporation, Billerica, 120 MA). In laboratory 3 (L3) acetonitrile, isopropanol, water, methanol, ammonium formate 121 (OptimaTM LC/MS grade) and chloroform (LC/MS grade) were obtained from Fisher 122 Scientific (Schwerte, Germany). Formic acid (LC-MS grade) was purchased from Sigma-123 Aldrich (Sigma-Aldrich, Munich, Germany). In laboratory 4 (L4) acetonitrile, isopropanol, 124 methanol, and formic acid (all ULC-MS grade) were from Biosolve (Valkenswaard, 125 Netherlands). Ammonium formate and chloroform were from Sigma-Aldrich GmbH 126 (Taufkirchen, Germany). 127

128 **2.2. OxPAPC preparation.**

129 OxPAPC was prepared either in one laboratory (L3) or independently in four 130 participating laboratories following the protocol describe below. A mixture of 1-palmitoyl-2-

linoleoyl-*sn*-glycero-3-phosphocholine (PAPC; 50 μ g) and 1-2-dimiristoyl-*sn*-glycero-3phosphocholine (DMPC; 2.2 μ g; a fully saturated PC used as internal standard for relative quantification) in chloroform was dried under a stream of N₂ in a 2 mL amber flat bottom vial to form a thin film at the vial bottom (three independent replicates). The lipid film was exposed to air at room temperature (L1 – 22-25°C, L2 – 16°C, L3 – 28°C, L4 – 25°C) for 0, 48 and 72 hours. After incubation the vial was filled with N₂, tightly sealed and stored at -80°C. Samples from L3 were sent to the other laboratories on freezer blocks at -20°C.

138

2.3. LC-MS/MS analysis.

The experimental replicates (n=3) of sample prepared in L3 (0 and 72 h of oxidation; 139 analysed by L1, L3, and L4), as well as samples generated independently in each of the four 140 participating laboratories and analysed there, were used to assess inter-laboratory variability. 141 In L4, experimental replicate 1 was analysed in triplicate on three consecutive days, to assess 142 intra- and inter-day analytical variability. Comparable chromatographic conditions and the 143 best available tandem MS instrument were used in the 4 laboratories. Details on LC-MS/MS 144 are provided in **Table 1** and Supplementary Methods section. Briefly, UHPLC separations 145 were performed on C18 reverse phase columns using binary solvent systems containing 146 water, acetonitrile, methanol and isopropanol, with ammonium formate (5 mmol/L) and 147 formic acid (0.1%) as additives [27]. MS analyses were performed using data-dependent 148 acquisition (DDA) in positive and negative ion modes (Table 1 and Supplementary 149 150 Methods).

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2.4. Automated identification of lipid peroxidation products (LPPs).

152 LPPtiger software source code version was used for lipid identification ([28]; https://bitbucket.org/SysMedOs/lpptiger). PAPC was oxidized in silico using theoretical 153 154 oxidation level 2. The list of modifications included hydroperoxy, hydroxy, epoxy, and keto groups with the maximum number equal to the number of bis-allylic and allylic sites in the 155 structure. The oxidative cleavage products with aldehyde and carboxylic acid on the terminal 156 carbon were included. LPPtiger predicted 345 unique oxidized fatty acyl chains 157 corresponding to 692 discrete oxPAPC species including lyso species. All raw spectra were 158 converted into mzML format using ProteoWizard MSconvert (Version 3.0.9134 64bit) and 159 the following parameters were used for all files: m/z range 400 to 1000, isotope score filter 160 80, and overall score filter 40. For L1 files 50 ppm on MS level and 200 ppm on MS2 level 161 were used as mass tolerance thresholds. For L2, L3, and L4 mass tolerance of 10 ppm on MS 162 and MS2 levels was applied. Analysis was performed using the integrated batch mode of 163 LPPtiger on a workstation equipped with 2 CPUs (32 cores) and 128 GB of RAM in L4. 164

165 **2.5. Relative LPPs quantification.**

166 15 LPPs detected in all four laboratories and previously reported in the literature as a 167 component of oxPAPC [9,11,13,14,16–18,23,29,30] were chosen for the relative 168 quantification (**Table 2**). Peak area for each lipid was integrated and normalized relative to 169 the DMPC lipid used as non-oxidizable internal standard. Average percent values for each 170 LPP relative to PAPC were calculated using normalized peak areas.

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2.6. Statistical analysis.

172 Coefficient of variation (CV%) was calculated using the following equation CV% = 173 DevSTD/AverageRelativeArea*100 for each LPPs using normalized peak areas. Principal 174 component analysis was performed using normalized peak area of 15 LPPs from L3 prepared 175 samples analyzed by three laboratories (L1, L3 and L4) using EZinfo (version 1.0, MKS 176 Instruments, Crewe, UK). ANOVA (Single Factor) analysis was performed by Analysis 177 ToolPak Excel Add-In for each LPPs using normalized peak areas.

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2.7. Nomenclature.

Lipid nomenclature is based on the LIPID MAPS consortium recommendations [31]. 179 For instance, the shorthand notation PC 36:4 represents a phosphatidylcholine lipid 180 containing 36 carbons and four double bonds. When the fatty acid identities and *sn*-position 181 are known, as in our case, the slash separator is used (e.g., PC 16:0/20:4). Since no unified 182 nomenclature is available for oxidized lipids, the short hand notations provided by LPPtiger 183 tool were used [28]. Short chain oxidized lipids were indicated by the corresponding terminal 184 enclosed in angular brackets (e.g. "<" and ">"), with the truncation site indicated by the 185 carbon atom number (e.g., <COOH@C9> and <CHO@C12). For long chain products our 186 recommendation is to indicate the number of oxygen addition after the fully identified parent 187 lipid (e.g. PC 16:0/20:4 + 10) when the type of addition is not known, or in parenthesis for 188 known functional groups (e.g. PC 16:0/20:4[1xOH@C11]). 189

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- **Table 1.** Summary of LC-MS/MS methods used for the analysis of oxPAPC in four different
- 198 laboratories.

	L1	L2	L3	L4	
Chromatography					
Column	Accucore TM C18	ACE C18	Accucore TM C18	Accucore TM C18	
Column phase	Solid core, C18	Solid core, C18	Solid core, C18	Solid core, C18	
Dimensions	150 x 2.1 mm	150 x 0.5 mm	150 x 2.1 mm	150 x 2.1 mm	
Particle size	2.6 µm	5 µm	2.6 µm	2.6 µm	
Pore size	150 Å	100 Å	150 Å	150 Å	
Gradient	isopropanol: acetonitrile: water	isopropanol: methanol: acetonitrile: water	isopropanol: acetonitrile: water	isopropanol: acetonitrile: water	
Temperature	50 °C	40 °C	50 °C	50 °C	
		Tandem MS			
Instrument	5600 TripleTOF	Q Exactive TM	 ✓ Orbitrap Fusion[™] Lumos[™] Tribrid[™] 	Q Exactive [™] Plus	
MS1	200-1500 m/z	400–1200 m/z	450–1200 m/z	380–1200 m/z	
Resolution	High sensitivity	70,000	120,000	140,000	
DDA top #	5	5	15	10	
Fragmentation	CID	HCD	HCD	HCD	
CE	35V, 10V CES	NCE 20, 23, 25	NCE 20 +/- 10	NCE 18 +/- 6	
Resolution MS2	High sensitivity	17,500	15,000	17,500	

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200 3. <u>Results and Discussion</u>

201

3.1. Identification of LPP species in air oxidized PAPC.

To evaluate the extent of PAPC oxidation and identify the LPPs formed, oxPAPC 202 samples prepared in four different laboratories (L1-L4) were analysed by LC-MS/MS using 203 204 reverse phase separation on C18 columns and data-dependent acquisition on four different MS instruments (Table 1). OxPAPC mixtures were analysed in both positive and negative 205 206 ion modes. Positive ion mode data were used for relative quantification, given the higher ionization efficiency of PC lipids as protonated quasi-molecular ions, whereas the negative 207 ion mode data were used for identification of LPPs based on tandem mass spectra of the 208 formate adduct or deprotonated ions, as the negative ion fragmentation data contains 209 informative fragment anions of the modified acyl chains. 210

Fig. 1 shows representative positive ion mode base peak chromatograms for three 211 different time points of PAPC oxidation obtained from the four laboratories. Representative 212 negative ion mode base peak chromatograms are shown in **Supplementary Fig. S1**. DMPC, 213 which contains two fully saturated C14:0 acyl chains, was spiked into all preparation at equal 214 quantities (PAPC:DMPC, 20:1 molar ratio) and was used as an internal standard for relative 215 quantification. The general pattern of oxidation was very similar between all laboratories. At 216 the 0 h time point two main signals, corresponding to DMPC (m/z 678. 51⁺) and native PAPC 217 $(m/z 782.57^{+})$, are clearly visible, with almost a complete absence of LPPs. After 48 h of air 218 exposure, a large number of signals for analytes with shorter retention times than PAPC can 219 be observed, indicating the presence of species with higher polarities than native PAPC. After 220 221 72 h, most of the signals eluted towards the beginning of chromatographic gradients, generally in the first third, corresponding to chain-shortened oxidation products, accompanied 222 by a corresponding decrease of the signals eluting in the middle third of the gradient, 223 corresponding to long-chain oxidised products, as well a decrease the native PAPC signal 224 225 which appears towards the end of the gradient.



226

Fig. 1. Positive ion mode base peak chromatograms of PAPC lipid oxidized by air exposure
for 0, 48 and 72 h and analysed using reverse phase chromatography coupled on-line to
tandem mass spectrometry detection in L1 (A), L2 (B), L3 (C) and L4 (D).

230

LPPtiger, a new software for the identification of oxidized phospholipids from data-231 dependent LC-MS/MS datasets, was used for the automated identification of LPPs in the 232 oxPAPC preparations [28]. LPPtiger performs identification based on the fragment ions 233 observed in CID or HCD experiments in the negative ion mode. The presence of specific 234 anions corresponding to native and oxidized acyl chains in PLs allows identification of the 235 oxidation type, whereas fragment ions and specific neutral losses provides identification of 236 PL class. Identification of positional isomers of oxPL is not supported by the current version 237 of the software, but the MS/MS based identification of oxidized fatty acyl chains together 238 239 with isotopic scores correction provide a solid basis for the identification of molecular species. 240

Overall 55 different LPPs with unique elemental compositions were identified, providing 143 potential LPPs after considering functional group isomers (**Supplementary**

Table S1). The 55 identified LPPs covered a number of different types of oxidative products, 243 including the lysophosphatidylcholines (LPC) LPC(16:0) and LPC(20:4), as well as 40 short 244 chain and 15 long chain LPPs. 24 LPPs were identified by all three laboratories that 245 performed DDA analysis (L2 used an inclusion list function and thus was not considered for 246 this comparison), and 45 LPPs were identified by two laboratories. Overall, all of the main 247 LPPs generally regarded as being present in oxPAPC mixtures were successfully identified 248 by LPPtiger at the MS/MS level, including the truncated species 1-palmitoyl-2-(5-249 oxovaleroyl)-sn-glycero-3-phosphorylcholine (POVPC; m/z 594.38⁺) and 1-palmitoyl-2-250 glutaryl-sn-glycero-3-phosphocholine (PGPC; m/z 610.37⁺), long chain products with 251 hydroperoxy, hydroxy, and keto groups, and 1-palmitoyl-2-epoxyisoprostaneE2-sn-glycero-252 3-phosphocholine (PEIPC; m/z 828.54⁺). 253

The different numbers of LPPs identified by LPPtiger from the MS/MS data from the 254 four laboratories most probably relates to the differences in MS methods and instruments 255 used in the study. Thus, the highest number of LPPs was identified by L3 utilizing a Tribid 256 Orbitrap Fusion Lumos, which is capable of acquiring a high number of tandem mass spectra 257 (15 MS/MS) per one survey scan, followed by L4 and L1 in which top 10 and top 5 DDA 258 methods were applied using a Q Exactive[™] Plus and a TripleToF 5600 respectively. L2 259 performed targeted acquisition using inclusion lists for 63 LPPs, of which 12 were 260 successfully identified. 261

262

3.2. LC-MS/MS based identification of isomeric LPPs.

Reverse phase chromatographic separation allowed the resolution of multiple isomeric 263 LPP species. The C18 phase proved to be a good stationary phase for the separation of the 264 relatively polar analytes formed by the oxidation of PLs, which are characterized by a mid-265 range polarity compared to the other lipid classes. Reverse phase separation strongly depends 266 on the hydrophobicity of the analyte, usually described by the partition coefficients logP and 267 268 logD [32], along with to a lesser extent some other physico-chemical properties, such as dipole-dipole interactions, proton acceptor/donor interactions and analyte polarizability. In 269 comparison with native PAPC, which has a logP 8.4, introduction of a single hydroxy group 270 results a shift of the logP to 7.2, whereas presence of three hydroxy groups or the isoprostane 271 ring structures (e.g. PGF2 α) further shifts the logP values further to 5.1 and 4.5, respectively. 272 The use of a C18 stationary phase with an isopropanol-acetonitrile-water elution gradient 273 ensures optimal binding of these mid-polarity lipid analytes while still providing efficient 274 separation of lipid species. The combination of LC separation with tandem mass 275

276 spectrometry allows the acquisition of separate tandem mass spectra for the separated 277 isomeric LPPs, and CID or HCD fragment spectra acquired in the negative ion mode 278 provides the information necessary to identify the type and often the position of the 279 modification.

Fig. 2 illustrates three examples of LC-MS/MS based identification of short chain
oxPAPC species.

282 Extracted ion chromatograms (XICs) for POVPC (Fig. 2) in negative ([M+HCOOH-H⁺], m/z 638.37) and positive ([M+H⁺], m/z 594.38) ion modes both had two 283 chromatographic peaks with a retention times of 5.6 and 6.1 min. Although the POVPC 284 signal was already present at the 0 h oxidation time point, the intensity of this LPP increased 285 by more than two orders of magnitude over the oxidation time course (e.g. from 5e4 counts at 286 Oh to 8e6 counts at 72 h for the example shown in Fig. 2 of the analysis performed by L3). 287 The positive mode HCD tandem mass spectra showed the presence of the fragment ion 288 characteristic of the PC head group at m/z 184, whereas the negative ion mode tandem mass 289 spectra acquired at 5.1 and 6.1 min provided more detailed information on LPP structure. For 290 example, the fragment ion at m/z 578.35 corresponded to the loss of formate and a methyl 291 group, a characteristic loss for the PC head group. The most informative region was where 292 anions of palmitic acid (m/z 255.23) and the five carbon long modified acyl chain with a C5 293 terminal aldehyde could be detected (m/z 155.04). The differences in MS/MS spectra taken at 294 the different retention times can most likely be attributed to the *sn*-1/*sn*-2 positions of the two 295 fatty acyl chains. Thus, the most intense signal eluting at 6.1 min most probably corresponds 296 297 to 1-palmitoyl-2-(5-oxovaleroyl)-sn-glycero-3-phosphorylcholine whereas the signal at 5.6 min, which has a higher intensity of the m/z 115.04 ion, is probably due to the positional 298 299 isomer 1-(5-oxovaleroyl)-2-palmitoyl-sn-glycero-3-phosphorylcholine [33,34]. Similarly, two chromatographic peaks were observed in the XIC at the mass of PGPC (Supplementary 300 Fig. S2), and the XICs generated from both positive and negative mode data showed up to a 301 three orders of magnitude increase in PGPC signal over the oxidation time course. The 302 fragmentation data showed a different ratio between the anionic acyl chain fragment 303 intensities (m/z 255.23 for palmitic and m/z 113.02 and 145.05 for glutaric acid), and the 304 fragmentation spectra in negative ion mode were characterized by the diagnostic neutral loss 305 of 59 Da from the parent (fragment at m/z 549.28), corresponding to loss of CH₃COOH from 306 a terminal carboxylic acid, and the two fragment ions related to the short chain oxidized FA 307 $(m/z \ 113.02 \ \text{and} \ 145.05)$. These fragmentations with a Δm of 32 Da are due to the 308 rearrangement of the polar head group, as has been described previously [35]. 309

310

311 1-palmitoyl-2-(5-hydroxy-8-oxo-octenoyl)-*sn*-glycero-3-phosphorylcholine

(HOOAPC), a previously reported short chain oxidized lipid in oxPAPC preparations, was 312 also already present at time 0 h, and increased with time by up to two orders of magnitude 313 (Supplementary Fig. S3). Once more, it was possible to identify the presence of two 314 chromatographic peaks showing a different ratio between the intensities of the acyl chain 315 316 anion signals. The loss of formate and methyl groups in negative ion mode and the polar head group PC characteristic fragments in positive ion mode were observed. A diagnostic ion 317 318 confirming the hydroxyl group to be on an eight carbon long truncated acyl chain was present at m/z 153.06, resulting from water loss relative to the short chain oxidized FA (m/z 171.07). 319

Fig. 3 shows three examples of LC-MS/MS based identification of long chain oxPAPC 320 species. In this case, the XICs are much more complex than for the short chain LPPs, 321 indicating the presence of multiple isomeric species. For example, at least 11 different 322 chromatographic peaks, 8 of them showing baseline separation, were detected for a mass 323 corresponding to PAPC with the addition of one oxygen (Fig. 3). Furthermore, the 324 chromatographic profile differed at the three oxidation time points. At 0 h all 11 peaks were 325 present, while after 72 h some had disappeared. The fragmentation spectra in positive ion 326 327 mode showed, as expected, a major fragment ion from the PC head group, but it was also possible to see a fragment ion resulting from the loss of 18 Da, corresponding to the loss of 328 329 H₂O which is diagnostic of an OH group. Negative ion mode tandem mass spectra allowed the identification of many of the modification isomers. Thus, the isomeric PAPC oxidation 330 331 products with a hydroxy (RT 13.1) or an epoxy (RT 13.9) group were identified based on the diagnostic ions at m/z 127.08 and 155.07 respectively, with the hydroxy group identified as 332 333 being at position C7, and the epoxy group at C5-C6 (from the ion at m/z 191.18).

Supplementary Fig. S4 shows the identification of PAPC with the addition of two 334 oxygens, where the XIC shows six main peaks at time points 0 h and 48 h, while at 72 h 335 additional chromatographic peaks with shorter retention times were present. In positive ion 336 mode the earlier eluting peaks showed one and two neutral losses of water molecules (-18 337 and -36 Da), typical of diHETE derivatives, while the later eluting signals were characterized 338 by the neutral losses of 18 and 34 Da, diagnostic of lipid hydroperoxides. Furthermore, with 339 these species for the first time we could identify in negative ion mode a diagnostic fragment 340 ion (m/z 335.22) from the diHETE-PC, corresponding to arachidonic acid carrying two 341 hydroxyl groups, whereas for the hydroperoxide containing oxPAPC the predominant signal 342 was detected at m/z 317.21 (single water loss). Occasionally, it was even possible to detect a 343

signal at m/z 331.23 corresponding to [FA 20:4 + OOH - 2H] and characteristic fragments ions at 203.18 and 129.06 m/z, diagnostic for a hydroperoxide group at position C5.

Supplementary Fig. S5 demonstrates the identification of a long chain oxidized PAPC 346 derivative characterized by the addition of three oxygen atoms and the loss of two hydrogens. 347 The intensity over the oxidation time course increased by one order of magnitude between 0 348 and 48 h and then remained approximately constant. The most studied product with this mass 349 350 is PEIPC. The fragmentation spectrum in negative ion mode is shown in **Supplementary** Fig. 5B. All previously reported [36] diagnostic fragment ions were present (m/z 331.19, 351 313.18, 305.21, 287.21, 269.19), including the characteristic ions for the epoxy group at the 352 position C5-C6 (m/z 233.15, 215.14, 203.14, 147.07, 115.04). In positive ion mode the 353 fragmentation spectra were characterized by the PC-specific fragment ion (184.07 m/z) and 354 diagnostic OH neutral losses (loss of 18 and 36 Da). 355





Fig. 2. Extracted ion chromatograms (A), HCD tandem mass spectra in negative (B), and positive ion modes (C) and proposed structure (D) of POVPC. Color-coding of structurerelated signals in the negative mode MS/MS spectra (e.g. the anion of FA 16:0 in blue, oxFA in orange and green, and LPC fragment derived as NL of one FA in pink) corresponds to the color-coding of fragments assigned to proposed structures illustrated in the lower panel.



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Fig. 3. Extracted ion chromatograms (A), HCD tandem mass spectra in negative (B), and positive ion mode (C) and proposed structure (D) of PAPC with the addition of one oxygen. Color-coding of structure-related signals in negative mode MS/MS spectra (e.g. the anion of FA 16:0 in blue, oxFA in orange and green, and LPC fragment derived as NL of one FA in pink) corresponds to the color-coding of fragments assigned to proposed structures illustrated in the lower panel.

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3.3. Relative quantification of LPPs in air oxidized PAPC

15 LPPs, including lyso PCs (LPCs), short and long chain oxidation products, were
selected for relative quantification (**Table 2**). Before evaluating the reproducibility of PAPC
air oxidation in the four different laboratories, intra- and inter-laboratory analytical variability
was examined.

374

Table 2. Elemental composition and proposed structures for the 15 LPPs in the oxPAPC mixture used for quantification. For each LPP the exact mass and m/z values for corresponding formate adduct (or m/z deprotonated ions in case of short chain LPPs with carboxylic acid terminal groups, marked with *) and protonated ions are provided.

Formula_ Neutral	Proposed structures	Exact mass	neg m/z [M +HCOO ⁻] ⁻	pos m/z [M+H⁺]⁺
C ₂₄ H ₅₀ NO ₇ P	LPC(16:0)	495.3325	540.3307	496.3398
C ₂₈ H ₅₀ NO ₇ P	LPC(20:4)	543.3325	588.3307	544.3398
C ₂₉ H ₅₆ NO ₉ P	PC(16:0/5:0 <cho@c5>)</cho@c5>	593.3692	638.3669	594.3670
C ₂₉ H ₅₆ NO ₁₀ P	PC(16:0/5:0 <cooh@c5>)</cooh@c5>	609.3636	608.3564*	610.3715
C ₃₂ H ₆₀ NO ₁₀ P	PC(16:0/8:1[10H] <cho@c8>)</cho@c8>	649.3949	694.3931	650.4022
C ₃₄ H ₆₂ NO ₉ P	PC(16:0/10:2 <cho@c10>)</cho@c10>	659.4157	704.4139	660.4235
C ₃₇ H ₆₆ NO ₉ P	PC(16:0/13:3 <cho@c13>)</cho@c13>	699.4470	744.4452	700.4548
C ₄₄ H ₇₈ NO ₉ P	PC(16:0/20:4) +10 – 2H	795.5409	840.5391	796.5482
C ₄₄ H ₈₀ NO ₉ P	PC(16:0/20:4) +10	797.5565	842.5547	798.5638
C ₄₄ H ₇₆ NO ₁₀ P	PC(16:0/20:4) +20 – 4H	809.5201	854.5183	810.5274
C ₄₄ H ₈₀ NO ₁₀ P	PC(16:0/20:4) +20	813.5514	858.5496	814.5587
C ₄₄ H ₇₈ NO ₁₁ P	PC(16:0/20:4) +30 – 2H	827.5307	872.5289	828.5380
C ₄₄ H ₈₀ NO ₁₁ P	PC(16:0/20:4) +20	829.5464	874.5446	830.5536
C ₄₄ H ₈₀ NO ₁₂ P	PC(16:0/20:4) +40	845.5413	890.5395	846.5486
C ₄₄ H ₇₈ NO ₁₃ P	PC(16:0/20:4) +50 – 2H	859.5205	904.5187	860.5284

379

The LC-MS method used for relative LPP quantification had already been validated in 380 different laboratories for the analysis of complex lipidomes [27,37]. To provide an estimation 381 of the likely intra-lab accuracy of the quantification results for the LPPs, L4 evaluated intra-382 sample as well as analytical variation (CV%) by analyzing one experimental replicate of the 383 oxPAPC preparation for each oxidation time point on three consecutive days. Normalized 384 peak areas were calculated for both positive and negative ion modes (Supplementary Table 385 386 S2). The coefficient of variation was under 20% for all LPPs at all time points, except for 387 LPCs at time 0 h, which showed higher inter-day variability (up to 34%), probably as the storage of the oxPAPC sample over 3 days at -80 °C does not prevent the oxidation or 388 389 hydrolysis process completely.

To assess inter-laboratory analytical variance, oxPAPC samples generated in L3 (0 and 390 72h oxidation) were analyzed in three different laboratories (L1, L3 and L4). Normalized 391 peak areas were calculated and used to perform principle component analysis (PCA; 392 Supplementary Table S3). As expected, the analyses for the 0 and 72 h time points are well 393 394 separated on the PCA plot, whereas the analytical replicates are tightly clustered. The 0 h data is also well clustered, but there is clear separation at 72h for each laboratory. The first 395 two principle components on the PCA scores plot (Supplementary Fig. S6) explain 85 % of 396 the variation. The first principle component demonstrates separation based on the time of 397 PAPC air exposure, whereas second principle component (explaining 18 % of the variation) 398

399 corresponds to separation based on inter-laboratory instrument variations; samples analysed 400 by L1 are grouped together at the bottom of the PCA scores plot, while L3 and L4 analysed 401 samples cluster more closely together. The minor variations explained by second principle 402 component (18%) are probably due to the type of mass analyzers used in the study (L1 - Q-403 TOF mass spectrometer; L3 and L4 - quadrupole-Orbitrap MS).

Furthermore, to compare the impact of sample preparation at different locations with 404 the variation in analysis of a single, centrally prepared sample analyzed in different 405 laboratories, we performed PCA of the normalized peak areas of the 72h samples generated 406 407 in L3 and analysed in L1 (labeled L1a) and L4 (labeled L4a) and the samples generated 408 independently by the four laboratories and analysed in those laboratories (L1, L2, L3, and L4) (Supplementary Tables S3 and S5, Fig. 4). It is clear that the data for the sample 409 centrally generated in L3 but analysed at different locations cluster closely together and show 410 minimal variation, while the samples generated in different locations showed much higher 411 412 distribution. This demonstrates that the main variance was in the sample preparation, and that the analysis of the same sample at different laboratories, using three different instruments 413 (L1:5600 TripleTOF, L3: Orbitrap Fusion[™] Lumos[™] Tribrid[™] and L4: Q Exactive[™] 414 Plus), gave statistically very similar results. The reproducibility of analysis across the four 415 laboratories is supported by the 0 h data in **Supplementary Fig. S6**, which is also tightly 416 417 clustered.



Fig. 4: PCA scores plot for normalized LPPs peak area in sample generated at 72h in L3 and
analysed in three different laboratories (L1a, L3 and L4a), and samples prepared
independently in the four laboratories (L1, L2, L3, L4). Violet squares – L1, blue squares –
L1a, blue stars – L2, pink circles – L3, yellow triangle – L4, orange triangle – L4a.

418

Combining the results of the relative quantification of 15 LPPs from oxPAPC preparations produced and analyzed independently in four different laboratories (**Fig. 5**, **Table 3** and **Supplementary Table S4**), six LPPs were among the most abundant signals in all labs and could be routinely identified and quantified, and therefore were chosen to serve as representatives of the oxPAPC mixture at the 48 and 72 h time points.



428

Fig. 5. Relative quantities (expressed as % of unmodified PAPC after the normalization to internal standard) of 15 representative LPPs from PAPC exposed to
air oxidation for 0, 48 and 72 h. Graphs are shown for the time course at individual laboratories: L1 (A), L2 (B), L3 (C), and L4 (D) at 0 h, 48 h and 72 h.
Identifications for each of the LPP masses can be found in Supplementary Table S1.

Table 3. The most abundant LPPs in air oxidized PAPC preparations for each laboratory (L1L4) quantified relative to unmodified PAPC after normalization of corresponding peak areas
to internal standard.

LPP, give as m/z values of protonated ions	L1	L2	L3	L4
594.38+	24 %		232 %	211 %
610.37+		39 %	99 %	195 %
650.40^{+}	13 %	25 %		103 %
814.56+	11 %			
828.54^{+}	18 %	28 %	97 %	149 %
830.55+		15 %	42 %	
846.55+	18 %	55 %	98 %	157 %

435

These correspond to three short $(m/z 594.38^+, 610.37^+, 650.40^+)$, and three long chain 436 (m/z 828.55⁺, 830.55⁺, 846.55⁺) LPPs. Based on the results of the MS/MS identifications, the 437 short chain LPPs can be assigned to POVPC, PGPC and HOOAPC. For the long chain LPPs 438 each m/z value, and thus elemental composition, can correspond to several isomeric species 439 (Table 2 and Supplementary Table S1). For instance, the LPP with elemental composition 440 $C_{44}H_{78}NO_{11}P$ (*m/z* 828.55⁺), usually referred to in oxPAPC mixtures as epoxyisoprostane E2 441 containing PC (PEIPC), could also be represented by the following combinations of the 442 functional groups on a long chain LPP derived from AA: [OOH and keto], [2OH and keto], 443 [2 epoxy and keto], and [epoxy, OH and keto]. The LPPs with elemental compositions 444 $C_{44}H_{80}NO_{11}P$ (e.g. 3OH or OH + OOH) and $C_{44}H_{80}NO_{12}P$ (e.g. 4OH or 2OOH) are clearly 445 represented by a mixture of isomeric structures, as some of these were at least partially 446 447 separated by chromatography in this study. However, for the relative quantification reported here the total peak area for each m/z were combined to report overall abundance of all of the 448 449 LPP isomers in order to be comparable with previously reported values for oxPAPC mixtures, which have commonly been analyzed by direct infusion methods where the 450 isomeric species have not been differentiated. 451

Although the most abundant LPPs were essentially the same between all four participating laboratories, the abundances of oxidized lipids relative to unmodified PAPC were quite different (**Fig. 6**). Thus, after 48 h air exposure the oxPAPC prepared in L3 showed the highest oxidation levels, with POVPC being the most abundant LPP formed (245

456 % relative to unmodified PAPC). Among the long chain LPPs, the signals at m/z 846.55⁺ 457 (128 %) and 830.54⁺ (96 %) were the highest after 48 h of air exposure. After 72 h exposure 458 to the air oxidation, the highest LPPs quantities were detected in L4, with POVPC (211 %), 459 PGPC (195 %) and long chain LPPs at m/z 846.55⁺ (149 %) and 830.54⁺ (157 %) being the 460 most abundant.



461

Fig. 6. Relative quantities (expressed as % of unmodified PAPC after the normalization to internal standard) of 15 LPPs in PAPC exposed to the air oxidation comparing directly the relative quantities identified in L1, L2, L3, and L4 at 48 (A) and 72 h (B).

Thus, overall relative abundances for the same LPPs produced in four laboratories 465 differed markedly across the time course, and by up to 13.5-fold (e.g. signal at m/z 828.55⁺ in 466 L1 and L4) even after 72 h of PAPC oxidation. The most variation between laboratories was 467 seen at the 48 h time point and by 72 h the profiles in L2, 3 and 4 showed relatively similar 468 overall abundance of oxidized products, although still with significant variation in individual 469 species. In L1 the overall level of oxidation was still low even after 72 h. ANOVA analysis 470 471 (Table S5) confirmed the statistical significance of the differences, with p-values below 0.05 for all quantified LPPs except PGPC at the 72 h time point. 472

One likely source of the difference in the oxidation rates could be small differences in 473 the laboratory temperatures during PAPC air exposure, as average room temperatures in L1 474 were 22-25 °C, L2 16 °C, L3 28 °C, and L4 25 °C; this could result in changes in the rate of 475 the initiation of the free radical oxidation, which is then amplified by the radical chain 476 reaction. The laboratory temperature at which air oxidation of oxPAPC is performed is 477 usually not reported in publications and might be a significant confounder in reported results. 478 However, this is clearly not the only reason, as the L1 had the lowest level of oxidation 479 throughout, but was not the coldest laboratory. Most importantly from the point of view of 480 experiments that are dependent on bioactivity, there is clearly significant variability in the 481 482 rate of reaction and the distribution of oxidized products produced.

483

Table 4. "Truncation score" (TS) for the oxPAPC preparations rom the four participating labs, calculated as the ratio between the summed abundances of three of the most abundant short chain LPPs (signal at m/z 594.38⁺, 610.37⁺, 650.40⁺) to three of the most abundant long chain LPPs (signals at m/z 828.54⁺, 830.55⁺, 846.55⁺).

	()	oxPA	PC TS
2		48 h	72 h
	L1	0.48	1.00
	L2	0.28	0.68
7	L3	1.21	1.46
	L4	0.66	1.41

488

In the literature, as a very general classification of bioactive compounds in air oxidized
 PAPC preparations, short chain LPPs are usually associated with pro-inflammatory effects
 whereas long chain LPPs are often associated with anti-inflammatory or protective functions

492 [17,24,25]. As is clear from the data presented above, despite the oxPAPC samples being 493 generated in the four different laboratories using the same protocol, which required minimal 494 sample manipulation (air exposure of the PAPC dried film), and each preparation giving rise 495 to approximately the same set of abundant LPPs, there was a high variability in the total 496 oxidation levels and in the extent of generation of individual LPPs. For this reason, a MS 497 based evaluation for all in-house produced oxPAPC mixtures would be beneficial to clarify 498 their exact composition before testing the biological effects.

Furthermore, a useful marker to help to characterize oxPAPC samples could be the ratio 499 of short to long chain oxidation products, which we suggest could be called the "truncation 500 score" (TS). This can be calculated for each oxPAPC preparation using the simple formula 501 TS = \sum % of short chain LPPs/ \sum % of long chain LPPs. The TS calculated using, for 502 example, the three most abundant species from each classification (Table 3), would be 503 indicative of the ratio between short and long chain LPPs for each preparation, which is likely 504 to be one of the most important characteristics for bioactivity. A TS value equal to one would 505 indicate equal contribution of short and long chain LPPs, whereas TS > 1 and TS < 1 would 506 correspond to the prevalence of short or long chain LPPs, respectively. The results from this 507 study using the six most abundant species are shown in Table 4. This indicates that while the 508 overall oxidation levels from L2 and L4 at 72 h look similar in the base peak chromatograms 509 in Fig. 1, the preparation from L2 is higher in long chain components, while that for L4 is 510 higher in short chain. Therefore, although TS is a very rough estimation for the prevalence of 511 short over long chain LPPs, it could be used to evaluate the oxidation state balance of 512 513 oxPAPC preparations before using them in biological experiments. It is not necessary to perform full LC-MS/MS characterization to determine the TS, as direct infusion MS data, 514 515 which are usually obtained to validate PAPC oxidation by air exposure for in-house preparations, could be used. 516

517 One should also bear in mind that not it is not only the relative abundance of short 518 chain over the long chain LPPs important, but their absolute concentrations are also highly 519 significant. However, given the wide variety of chemical structures within oxPAPC mixture, 520 and thus differences in ionization efficiency of LPPs with various functional groups, pure 521 synthetic standards for each LPP class would be required to perform absolute quantification.

522 4. <u>Conclusions</u>

523 We have performed a multi-laboratory evaluation of air oxidized PAPC preparations 524 from four different laboratories using LC-MS/MS analysis, focusing on the identification and

relative quantification of the LPPs formed. High intra-sample reproducibility and a very 525 similar overall pattern of oxidation between the four laboratories was demonstrated. 526 However, significant differences in the general extent of lipid oxidation relative to 527 unmodified PAPC were observed. Thus, we propose that an MS based evaluation for all in 528 529 house produced oxPAPC mixtures should be performed before using them to study biological effects of oxidized lipids. Furthermore, we suggest calculating the "truncation score" as a 530 531 rough estimation of the balance between short vs long chain products of arachidonic acid oxidation. 532

533

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547

548 **Conflict of interests**

549 The authors declare no conflict of interests.

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Highlights

- Multi-laboratory evaluation of air oxidized PAPC preparations by LC-MS/MS.
- Identification and relative quantification of lipid peroxidation products (LPPs).
- oxPAPC preparations should be characterized by MS prior testing biological effects of oxidized lipids.
- "Truncation score" is proposed as a rough estimation of oxPAPC oxidation status.